Untangling the effects of size, habitat and invertebrate biodiversity on parasite prevalence in the Caribbean spiny lobster

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Abstract

The spiny lobster *Panulirus argus* is an important benthic mesopredator and a major fishing resource across the Wider Caribbean region. This species is host to the pathogenic virus PaV1 and metacercariae of *Cymatocarpus solearis*, a digenean trematode whose first intermediate host remains unknown. Previous studies found that the probability of infection with PaV1 was higher in juvenile lobsters and in densely vegetated habitats (suggesting that marine vegetation can be an environmental reservoir for PaV1), whereas the probability of infection with *C. solearis* was higher for larger lobsters and in poorly vegetated habitats. To increase insight into the role of habitat and body size in the ecology of lobster diseases, the presence of both *C. solearis* and PaV1 in *P. argus* was investigated across three contrasting zones in Bahía de la Ascensión, Mexico (19°35′27″N, 87°38′06″W): reef, lagoon and shallow habitat. Additionally, habitat complexity, cover of benthic components, and macroinvertebrate biodiversity were characterized in each zone. Consistent with previous studies, probability of infection with PaV1 (both at a clinical and infective level) decreased with increasing lobster size and was highest in the seagrass-rich lagoon, supporting the idea that marine vegetation could be an environmental reservoir for PaV1. In contrast, the probability of infection with *C. solearis* increased significantly with lobster size but did not vary with zone, suggesting no relationship with benthic substrate type. However, based on results of macroinvertebrate diversity, the gastropods *Cerithium litteratum* and *Tegula fasciata* are put forward as potential candidates for the first intermediate hosts of *C. solearis*.

Introduction

In the changing climate, marine diseases are thought to be on the rise (Ward and Lafferty 2004; Lafferty and Hofmann 2016). Anthropogenic factors associated with climate change, pollution and invasive species have all been research targets for understanding marine disease dynamics (Harvell et al. 2002, 2004). However, ecosystem health as a whole can be encompassed by a range of categories, from anthropogenic disturbances to ecological factors such as species richness and diversity as well as disease (Lafferty and Gerber 2002; McCallum et al. 2005; Aburto-Oropeza et al. 2015).

In the Wider Caribbean region, the spiny lobster *Panulirus argus* is an important benthic mesopredator and a major fishing resource, with annual landings of over 27,000 tons in the past 10 years (FishStatJ 2019). This species, however, can be infected with PaV1 (*Panulirus argus* virus 1), the first known naturally occurring virus of a lobster, as well as a range of macroparasites, bacteria, and fungi (see Shields 2011 for review). *Panulirus argus* virus 1 was first discovered as the causative agent of an emergent disease affecting *P. argus* in Florida Bay, USA in 1999 by Shields and Behringer (2004). This species, however, can be infected with PaV1 (*Panulirus argus* virus 1), the first known naturally occurring virus of a lobster, as well as a range of macroparasites, bacteria, and fungi (see Shields 2011 for review). *Panulirus argus* virus 1 was first discovered as the causative agent of an emergent disease affecting *P. argus* in Florida Bay, USA in 1999 by Shields and Behringer (2004). It has since been found throughout the Caribbean (Moss et al. 2013). *Panulirus argus* virus 1 has been mainly reported in juvenile lobsters (Behringer et al. 2006) and is highly pathogenic. Laboratory trials have shown that PaV1 is mainly transmitted through contact...
between healthy and diseased lobsters, and to a lesser extent through ingestion of diseased tissue and through water (Butler et al. 2008).

*Panulirus argus* has also been found to host metacercariae of the digenean trematode, *Cymatocarpus solearlis* (Gómez del Prado et al. 2003). The cysts of the metacercariae are large, measuring ~1 mm in diameter on average (0.6–1.5 mm, Gómez del Prado et al. 2003; Cruz-Quintana 2012). However, trematode life cycles are typically very complex involving several intermediate hosts in addition to the definitive hosts (Cribb et al. 2003). Although the first intermediate host of this parasite is unknown, it is known that *P. argus* is a second intermediate host, with the definitive host being marine turtles. Little is known about the effects of this parasite upon the physiology or behavior of *P. argus* (Briones-Fourzán et al. 2016).

Host–parasite interactions are typically embedded in complex communities that may alter parasite transmission (Thieltges et al. 2008; Orlofske et al. 2012), but research into reservoirs of PaV1 and effects upon the ecosystem as a whole has also been limited (e.g., Butler et al. 2008). However, studies have noted higher PaV1 prevalence in densely vegetated areas compared to poorly vegetated areas, suggesting that marine vegetation may be acting as an environmental reservoir for the disease (Briones-Fourzán et al. 2012; Huchin-Mian et al. 2013). Both seagrass meadows and macroalgae have been shown to attenuate flow velocity of water (e.g., Scoffin 1970; Peterson et al. 2004), therefore, it could also be hypothesized that densely vegetated areas may cause pooling of virions from infected animals. In contrast to the disease ecology of PaV1, it was found that the prevalence of the parasite *C. solearlis* was higher in poorly vegetated habitats relative to densely vegetated habitats (Briones-Fourzán et al. 2016). Also, the probability of infection decreased with increasing lobster size for PaV1, but increased with increasing lobster size for *C. solearlis*, and coinfection with both parasites was almost nil. Previous studies have indicated that the biodiversity of species present as well as the habitat type and complexity can contribute to disease risk (Keasing et al. 2006; Johnson and Thieltges 2010) and it is well known that vegetation and food sources may act as vectors or reservoirs for disease (Small and Pagenkopp 2011).

The current study aims to discover more about PaV1 and the little known parasite *C. solearlis* upon the populations of *P. argus* in a marine protected area in the Caribbean. The roles of habitat and surrounding communities, both as possible vectors or reservoirs, are crucial in understanding the disease ecology of these parasites and also in terms of fisheries due to the extensive use of casitas (artificial shelters) in fisheries for *P. argus* in the Caribbean. Spiny lobsters within high-risk habitats that promote disease acquisition may result in altered population dynamics and further impact upon the lucrative fishery (Stewart 1984; Shields 2011). Investigating potential relationships between disease presence and factors such as habitat complexity, components of the benthic habitat, and species richness and species diversity (i.e., community) would be beneficial to inform scientists, fishers and managers about the current and potential effects of the diseases related to spiny lobster habitat selection.

### Materials and methods

#### Study area

Bahía de la Ascensión (centered at 19°35’27”N, 87°38’06”W) is a large, shallow bay (mean depth: 2.2 m; range <1 to 7 m; Medina-Gómez et al. 2014) located within the Sian Ka’an Biosphere Reserve on the Mexican Caribbean coast of Quintana Roo (Fig. 1). The bay is bordered by extensive wetlands, in particular mangrove forests, with several mangrove keys throughout the bay, most notably at its center. The bay has relatively calm water, because an interrupted coral reef that runs parallel to the mouth of the bay reduces wave surge. Large expanses of the bay are covered by extensive meadows of mixed seagrasses and macroalgae, offering a good settlement habitat for postlarvae of *P. argus* (Briones-Fourzán et al. 2012). The bay is characterized by strong SW–NE salinity gradients due to freshwater supply from combined runoff and sub-marine groundwater discharges in the inner bay, and strong tidal influence in the bay’s central basin favored by the open seaward front (Medina-Gómez et al. 2014). The bay sustains an important local fishery for *P. argus* based on the extensive use of “casitas”, artificial habitats constructed of concrete that mimic large crevice-type shelters sought by *P. argus*, widely used for lobster fishing in the Caribbean (Lozano-Alvarez et al. 1991). The only human settlement along the entire margin of the bay is a small village (Javier Rojo Gómez), which is located on the land point named Punta Allen (Fig. 1). The village is the seat of the local fishing cooperative, which holds the concession to fish for lobsters in the entire bay and adjacent marine areas.

#### Sampling operations

Historically, the fishing cooperative has been supportive of scientific research in the bay; however, they prefer that research activities take place during the lobster closed season (March 1 to June 30) so as not to interfere with fishing activities. Therefore, sampling took place in April–May 2017. Three bay zones were sampled with contrasting types of habitats: the interface between the reef lagoon and the back reef zone along a large reef (hereafter “Reef”, centered at 19°47’29.70”N, 87°26’39.18”W), the middle...
portion of the reef lagoon located between Punta Allen and
the reef (hereafter “Lagoon”, centered at 19°47′34.68″
N, 87°27′10.02″W), and a shallow, inner coastal zone far from
the reef (hereafter “Shallow”, centered at 19°46′46.06″
N, 87°34′59.10″W) (see Fig. 1). Areas were chosen that
included casitas, as these structures harbor lobsters over a
wide size range, from juveniles to adults. Casitas were cho-
sen at random, and once searched, were marked to ensure
they were not re-sampled. In addition to sampling lobsters,
the three sampling zones were each characterized based on
the following criteria: percent cover of benthic community
components, habitat complexity, and diversity of macroin-
vertebrate communities. SCUBA diving was used to conduct
all sampling activities.

Habitat characterization

Within each sampling zone, 7 transects were laid, each
25 m in length and marked every 50 cm. Transects were
laid parallel to the reef and the shore. The point intercept
method (Álvarez-Filip et al. 2011) was used to estimate
the percentage of cover for each of the following benthic
components: sand, seagrasses, erect macroalgae, coralline
algae, sponges, rubble, calcareous pavement, soft corals,
live hard corals, and dead coral. This was done by record-
ing the type of benthic component observed below each
mark (= 50 estimates per transect). Because points are
essentially dimensionless, this method is considered the
least biased and most repeatable for determining cover
(Elzinga et al. 1998).

Next, 3 quadrats, 2 m × 2 m each, were laid at the begin-
ing, middle and end of each transect to assess habitat
complexity using the habitat assessment score (HAS, see
Gratwicke and Speight 2005; Lozano-Álvarez et al. 2017).
HAS provides an overall structural complexity value by
visually evaluating six variables of the local topography
(rugosity, variety of growth forms, height, refuge size cat-
egories, percentage of live cover, and percentage of hard
substratum). Each variable is assigned a score between 1
and 5 (from smallest or lowest to largest or highest), and
the sum of the individual scores is the HAS. Therefore, a
score of 6 would represent the least complex habitats and
a score of 30 would represent the most complex habitats.
The HAS was obtained by adding the scores of all com-
ponents per quadrat. The HAS score, a qualitative metric, has
the advantage that it can be applied in habitats from simple
(e.g., sand) to very complex (e.g., coral reefs) (Gratwicke
and Speight 2005).
Assessment of macroinvertebrate biodiversity

To examine the potential relationship between the invertebrate community and the presence and prevalence levels of PaV1 and/or C. solearis at each zone, conspicuous invertebrates (those visible above the sand or fauna) were identified and quantified within the same 2 m × 2 m quadrats used to assess habitat complexity. All underwater identifications were conducted by two scientific observers thoroughly trained prior to the samplings. Training was achieved by repeatedly studying an extensive guide of local invertebrate species created in our lab from photos and drawings obtained from many different sources (e.g., Stanley 1970; Chace 1972; Abbot 1974; Williams 1984; Abele and Kim 1986; Wagner 1990; Humman 1994; Hendler et al. 1995; García-Cubas et al. 1999; Medina et al. 2006; Ng et al. 2008; Laguarda-Figuera et al. 2009; McLaughlin et al. 2010; de Grave and Fransen 2011; Diez García and Jover Capote 2015; Carmona-Suárez and Poupin 2016), followed by direct identification in the field during preliminary dives, with the results being cross-checked between both divers (Lessios 1996; González-Gómez et al. 2018).

Lobster sampling

The fishing cooperative allowed us to sample lobsters as necessary provided we did not kill them. Therefore, assessment of parasites in lobsters was only visually conducted although, in the case of PaV1, we also extracted hemolymph from lobsters for PCR assays. All lobsters encountered beneath casitas along and around the transects were collected with hand nets. Nets containing lobsters were fastened to the edge of the boat with lobsters remaining in the water to avoid exposure to air. Lobsters were sexed, measured (carapace length, CL, from the inter-orbital notch to the rear end carapace, in millimeter), and carefully examined for clinical signs of infection with PaV1 (milky hemolymph, visible through the translucent membrane between the carapace and abdomen, Candia-Zulbarán et al. 2012) and for the presence of C. solearis cysts (obvious to the naked eye as small spheres of ~1 mm in diameter in the muscles of the abdomen, visible through the translucent membrane between the carapace and the abdomen, Candia-Zulbarán et al. 2012) and for the presence of C. solearis cysts (obvious to the naked eye as small spheres of ~1 mm in diameter in the muscles of the abdomen, visible through the translucent membrane between the carapace and the abdomen, or through the relatively thin cuticle of the ventral side of the abdomen, Gómez del Prado et al. 2003). Although it was not possible to dissect lobsters, we were confident that the cysts belonged to C. solearis, because decapod populations (except for brachyuran crabs) tend to harbor only one species of trematode (Thieltges et al. 2009) and, in previous studies, all metacercariae obtained from cysts in tails of P. argus were identified as C. solearis (Grano-Maldonado and Álvarez-Cadena 2010; Cruz-Quintana 2012). Following measurements, ~200 μl hemolymph was taken from the base of the fifth pair of pereopods using a 30-gauge sterile needle and 1 ml syringe, after disinfection of the puncture site with 70% ethanol. Hemolymph samples were fixed immediately in ice-cold 96% ethanol, transported to the laboratory and frozen at −20 °C. Lobsters were returned to the capture site.

Hemolymph surveillance of PaV1

DNA extraction

DNA was extracted from ~200 μl of hemolymph/ethanol mixture (~25 mg of hemolymph) with the Wizard genomic DNA purification kit (Promega) following a slightly modified manufacturer’s protocol (Online Resource 1). DNA eluted was used as the template for polymerase chain reaction (PCR). Haemolymph DNA extraction was optimized to ensure detection of PaV1 using known, positive controls initially derived from P. argus, confirmed by sequencing. DNA extractions were verified by running 1 μl DNA mixed with 4 μl Promega Green GoTaq® 5× Flexi Buffer on a 1.5% agarose gel.

PCR conditions

All PCRs were carried out using primers synthesized by Sigma and performed on a 3Prime Personal Thermal Cycler (Techne, UK) before being visualized on a 1.5% agarose gel. To test for the presence of PaV1 in lobsters, a PCR was performed using known, specific primers for PaV1 (45aF: TTC CAG CCC AGG TAC GTA TC; and 543aR: AAC AGA TTT TCC AGC AGC GT) that amplify a region of 499 bp (Montgomery-Fullerton et al. 2007). All PCR reactions were carried out in a total volume of 10 μl containing 1 μl extracted DNA (50–200 ng/μl), 0.33 mM of each primer 45aF and 543aR, 2.5 mM MgCl2 (Promega), 0.6× Green GoTaq® 5× Flexi Buffer (Promega), 0.4 mM dNTP mixture (Promega), and 0.75 u GoTaq® Flexi DNA Polymerase (Promega).

Statistical analyses

Habitat characterization

The data of structural complexity (HAS score) were analyzed to verify the normality and homoscedasticity using D’Agostino–Pearson’s normality test. As these assumptions were not met, comparison of values between zones was performed using the Kruskal–Wallis test followed by Dunn’s multiple comparisons test (Dunn 1964), all using a significance level of 95%. For these analyses, GraphPad Prism 7.0 was used.

The percent data on the cover of benthic components were logit transformed (Warton and Hui 2011) and
subjected to a principal component analysis (PCA) using the software PAST v.3.09 (Hammer et al. 2001). The transformed data for each benthic component were then compared among zones with independent one-way analysis of variance (ANOVA) using a general linear model (GLM) approach (Rutherford 2001).

Macroinvertebrate biodiversity

For each sampling zone, the following ecological indices were estimated: species richness (S, number of species), Shannon–Wiener’s diversity (H’), Pielou’s evenness (J’) and Simpson’s dominance (D). S is an informative index as it is the basis of biodiversity estimates, whereas H’, J’ and D are compound indices (i.e., indices that combine richness and abundance of species) that provide a greater ability to discriminate sites than S alone (Morris et al. 2014). Each index was compared among zones with a separate one-way ANOVA using a GLM approach.

Differences in community composition between sampling zones were analyzed by non-metric multidimensional scaling (MDS) using the Bray–Curtis similarity measure on a fourth root transformation of the abundance data (Clarke 1993). The significance of the observed differences among zones was further tested with a one-way analysis of similarity (ANOSIM), which provides an R value, typically between 0 and 1. Values close to 0 are indicative of only a little difference, whereas values close to 1 are indicative of a large difference in sample composition (Clarke and Warwick 2001). However, as the stress value of the 2D MDS plot was moderately high (see Results), results from this plot were cross-checked against those from a cluster analysis with group average linkage (Clarke and Warwick 2001; Clarke and Gorley 2015). The significance of the cluster findings was further tested with a similarity profile analysis (SIMPROF), which tests the null hypothesis that the set of samples contains no multivariate structure that can be further examined (Clarke et al. 2008). Finally, a similarity percentage analysis (SIMPER) was performed to identify those species that were more important in creating the observed pattern of similarity (Clarke 1993). PRIMER 6 v6.1.9 (PRIMER-E Ltd) software was used to carry out these analyses.

Lobster size distribution

Size distribution of lobsters was plotted per zone in 5-mm increments. The data did not meet the assumption of normality; therefore, a Kruskal–Wallis test with a significance level of 0.05 was used to compare lobster size among zones.

Disease ecology

Prevalence (the percentage of infected lobsters) was estimated for (a) lobsters clinically infected with PaV1 (i.e., lobsters exhibiting gross signs as described in methodology), (b) lobsters testing positive for PaV1 via PCR, not necessarily exhibiting signs of disease (i.e. those picked up via the highly sensitive PCR method), and (c) lobsters observed harboring at least one cyst of C. solearis, simply referred to as trematodes. In all cases, 95% confidence intervals were computed using Wilson’s score method with continuity correction (Newcombe 1998).

Binomial logistic regression models with logit link functions (Quinn and Keough 2002) were used to determine whether specific predictor variables had a significant effect on the probability of finding lobsters clinically infected with PaV1, lobsters testing positive to PaV1 infection by PCR, and lobsters infected with C. solearis. The predictor variables were size (CL, covariate) and zone (categorical factor with three levels: Reef, Lagoon and Shallow, with Shallow as the reference level). In the models for lobsters testing positive to PaV1 by PCR and lobsters infected with C. solearis, the effect of the presence of the other parasite in the same lobster (i.e., presence of C. solearis for the model examining probability of infection with PaV1 by PCR, and viceversa) was also examined (categorical factor with two levels: presence and absence, with presence as the reference level). It was not possible to test the effect of C. solearis on the probability of clinical infection with PaV1, because lobsters with both C. solearis and clinical signs of PaV1 were not present at all zones. However, Briones-Fourzán et al. (2016) pooled the data from all their sampling sites to test the effect of lobster size and the presence of clinical signs of PaV1 on the probability of finding lobsters with trematodes; therefore, for comparative purposes, this was also done in the present study. All logistic models were run in the software Statistica v.10 (StatSoft, Inc., USA).

Results

Habitat characterization

Habitat complexity

The HAS score showed that the three zones in Bahía de la Ascensión were distinctly separated (Fig. 2). Dunn’s multiple comparisons test revealed that the most significant difference in habitat complexity was between Shallow and Lagoon (p < 0.0001, mean rank diff. =−28.83) followed by Shallow vs. Reef (p =0.0051, mean rank diff. =−17.6). Lagoon vs. Reef, however, was not significantly different (p =0.1348, mean rank diff. =11.24). This indicates
that Shallow was the zone with the least complex habitat, with a lower overall HAS score compared to the other two zones.

**Cover of benthic components**

The first two axes of the PCA of the percent cover of benthic components explained 51.3% and 18.3% of the cumulative variance, respectively. The first axis was positively related with coralline algae and seagrasses (loadings 0.458 and 0.422, respectively) and negatively related with calcareous pavement and sand (−0.498 and −0.426, respectively), whereas the second axis was positively related with rubble (0.648) and negatively related with seagrasses (−0.526) (Fig. 3). Transects on Shallow were distinctly separated from transects on the other two zones along the first axis, whereas transects on Reef and transects on Lagoon differed mostly along the second axis (Fig. 3). The GLMs comparing each benthic component among zones revealed that all components (except rubble) differed significantly with zone (Table 1). In particular, Lagoon was dominated by seagrass (67.7% of total cover) followed at a distance by sand (11.7%), whereas Shallow was dominated by sand (56.9% of total cover) followed by calcareous pavement (17.4%) and macroalgae (16.6%). In contrast, dominance was not so pronounced in Reef, which exhibited a 38% cover by seagrass, followed by 18.6% cover of coralline algae and by several components with coverages between 10 and 13% (sand, dead coral, rubble) (Table 1).

**Assessment of macroinvertebrate community**

Identification of conspicuous invertebrates yielded 63 taxa in total (2 anemones, 6 polychaetes, 25 crustaceans, 7 echinoderms, and 23 mollusks), with 35 taxa in Reef, 38 in Lagoon, and 20 in Shallow (Online Resource 2). Most taxa were identified to species level. Ecological indices $S$, $H'$ and $J'$ were higher in Reef and Lagoon than in Shallow, but $D$ was far greater in Shallow than in the other two zones (Table 2). The 2D MDS ordination plot had a relatively high stress value (0.19); therefore, conclusions derived from this plot were cross-checked against those from the cluster analysis and SIMPROF test (Clarke and Warwick 2001) (Fig. 4a). The resulting plot (Fig. 4b) shows that, at 20% similarity, all quadrats from Reef and Lagoon formed one group, whereas quadrats from Shallow formed two groups. This was mainly due to the high abundance of the yellow mussel *Brachidontes modiolus* in three of the quadrats on Shallow. At 30% similarity, several groups with variable degrees of overlap were formed within each zone, but Shallow remained distinctly separated from the other two zones. This was confirmed by ANOSIM, which yielded an overall $R = 0.704$. $R$ values were higher in pairwise comparisons involving Shallow (vs Reef, $R = 0.747$; vs Lagoon, $R = 0.807$), than between Reef and Lagoon ($R = 0.514$).

Results of SIMPER (Online Resource 3) showed a higher average similarity among quadrats within Shallow (42.4%) and within Lagoon (41.4%) than among quadrats within Reef (33.2%). A cumulative similarity of ~90% among quadrats was accounted for by nine species in Lagoon and eight in Reef, but only by four species in Shallow. Average...
nia crassicornis) (Online Resource 3). Modiolus and a polychaete (the spaghetti worm B. Chione cancellata-barred venus and the yellow mussel Pagurus annulipes banded hermit and the brown-included two decapods (P. brevidactylus tricolor). Finally, the five more common species in Shallow were three gastropods (the pine star shell Cribanarius brevidactylus and the blue-legged hermit crab P. Lithopoma phoebium) and two decapods (the nodose clinging crab O. rubicunda).}

Disease ecology

As a whole, across all zones, significantly more lobsters were found infected with PaV1 using PCR (Fig. 6). However, the pattern differed between zones. Significantly more lobsters residing in Lagoon were infected with PaV1, both clinically and detected via PCR, than in Reef and Shallow. Fewer lobsters infected with trematodes occurred in Lagoon than in Reef and Shallow (Fig. 6).

The logistic regression models showed that the probability of finding lobsters with clinical signs of PaV1 infection decreased with increasing size of lobster and was significantly higher in Lagoon than in Shallow (odds ratio, OR: 1.97), but was similar between Reef and Shallow (Table 3A). The probability of finding lobsters testing positive to PaV1 by PCR also decreased with increasing lobster size and was not affected by the presence of C. solearis in lobsters, but was significantly higher in Lagoon (OR: 1.97) and Reef (OR: 1.77) than in Shallow (Table 3B). By contrast, the probability of infection with trematodes increased with increasing lobster size, but was not affected by the presence of PaV1 and did not vary significantly with zone (Table 3C). Upon pooling the data from all sampling zones, the probability of finding lobsters with trematodes increased significantly

| Table 1 | Percent cover (mean ± 95% confidence interval) of benthic constituents on three zones in Bahía de la Ascensión (Reef, Lagoon, Shallow; \( N = 7 \) transects per zone) |
|---|---|---|---|---|
| Benthic component | Reef | Lagoon | Shallow | \( F \) | \( df \) | \( p \) |
| Seagrasses | 38.0 ± 15.6\(^a\) | 67.7 ± 6.0\(^b\) | 6.0 ± 2.8\(^c\) | 28.725 | 2, 18 | <0.001 |
| Erect macroalgae | 5.1 ± 5.9\(^a\) | 9.4 ± 2.7\(^b\) | 16.6 ± 7.4\(^b\) | 6.889 | 2, 18 | 0.006 |
| Coralline algae | 18.6 ± 5.6\(^a\) | 4.6 ± 4.7\(^b\) | 0\(^b\) | 24.602 | 2, 18 | <0.001 |
| Sand | 13.4 ± 10.0\(^a\) | 11.7 ± 4.3\(^a\) | 56.9 ± 10.4\(^b\) | 14.006 | 2, 18 | <0.001 |
| Sponges | 0\(^a\) | 2.3 ± 1.8\(^b\) | 0.3 ± 0.6\(^b\) | 5.179 | 2, 18 | 0.017 |
| Rubble | 10.3 ± 13.7\(^a\) | 3.7 ± 3.5\(^a\) | 2.9 ± 3.6\(^a\) | 0.084 | 2, 18 | 0.920 |
| Live hard coral | 3.1 ± 1.7\(^a\) | 0.6 ± 0.7\(^b\) | 0\(^b\) | 11.784 | 2, 18 | 0.001 |
| Dead coral | 11.4 ± 6.0\(^a\) | 0\(^b\) | 0\(^b\) | 49.351 | 2, 18 | <0.001 |
| Calcareous pavement | 0\(^a\) | 0\(^a\) | 17.4 ± 6.3\(^b\) | 154.103 | 2, 18 | <0.001 |

Mean value (± 95% confidence interval) of species richness (\( S \)), Shannon–Wiener diversity (\( H' \)), evenness (\( J' \)), and dominance (\( D \)) of macroinvertebrates on three sampling zones in Bahía de la Ascensión, México (Reef, Lagoon, Shallow). For each index, different superscripts indicate significantly different means (ANOVA, \( \alpha = 0.05 \)).

Table 2: Ecological indices for macroinvertebrates by quadrat per zone

| Ecological index | Reef | Lagoon | Shallow |
|---|---|---|---|
| \( S \) | 6.80 ± 1.38\(^a\) | 8.52 ± 0.74\(^b\) | 3.86 ± 0.62\(^c\) |
| \( H' \) | 1.65 ± 0.18\(^a\) | 1.85 ± 0.20\(^b\) | 0.91 ± 0.19\(^c\) |
| \( D \) | 0.19 ± 0.04\(^a\) | 0.20 ± 0.08\(^a\) | 0.48 ± 0.10\(^b\) |
| \( J' \) | 0.91 ± 0.03\(^a\) | 0.87 ± 0.08\(^a\) | 0.73 ± 0.11\(^b\) |

Disease ecology

Lobster size distribution

In total, 430 lobsters were collected (132 in Reef, 150 in Lagoon, and 148 in Shallow) with an overall size range of 12.2–131.9 mm CL. The size distribution of lobsters exhibited two modes in Shallow and Lagoon (Fig. 5) and the data did not meet the assumption of normality. There was a significant difference in lobster sizes among zones (Kruskal–Wallis test, \( H_2 = 115, p < 0.0001 \)), with Dunn’s multiple comparisons test revealing that Reef lobsters were significantly larger than those found in Lagoon and in Shallow, but also the lobsters in Lagoon being significantly larger than those in Shallow (mean rank diff. Shallow vs. Lagoon = −37.73, \( p = 0.0264 \); Shallow vs. Reef = −154.5, \( p < 0.0001 \); Lagoon vs. Reef = −116.8, \( p < 0.0001 \)).
Fig. 4 Invertebrate community structure in benthic samples from three sampling zones in Bahía de la Ascensión: Reef (blue dots), Lagoon (green triangles), and Shallow (orange squares). \textbf{a} Hierarchical clustering (group average linkage) and \textbf{b} non-metric multidimensional scaling plots showing groups of quadrats within 20\% (continuous line) and 30\% (broken gray line) similarity contours given by sequence of SIMPROF tests on dendrograms from \textbf{a}. Analyses were done using Bray–Curtis similarity and fourth root transformations. Each symbol denotes a quadrat ($N = 21$ per zone)

Fig. 5 Size frequency distribution of individual lobsters caught in the three chosen zones within Bahía de la Ascensión; Shallow (orange, $N = 148$), Lagoon (green, $N = 150$) and Reef (blue, $N = 132$)
with size but was not affected by the presence of clinical signs of PaV1 (Table 4A) or by the presence of PaV1 via PCR (Table 4B).

Discussion

This investigation aimed to increase insight into the complex relationships between habitat characteristics and parasites of spiny lobsters. An inverse relationship between lobster size and the probability of infection with PaV1 was found, which is consistent with findings of previous studies (e.g., Butler et al. 2008; Behringer et al. 2011; Candia-Zulbarán et al. 2012; Briones-Fourzán et al. 2012; Huchin-Mian et al. 2013). However, even after accounting for the significant effect of size on the probability of infection with PaV1, such probability was highest in Lagoon, where seagrass was far more abundant, supporting the idea first put forward by Briones-Fourzán et al. (2012) that marine vegetation could be an environmental reservoir for PaV1. Seagrass meadows and macroalgal beds trap particles, because they attenuate water flow velocity (Scoffin 1970; Peterson et al. 2004). Thus, they have natural filtration abilities, which can, for example, reduce the abundance of bacteria reaching coral reefs (Lamb et al. 2017). This suggests that marine vegetation can concentrate viral particles, therefore potentially increasing the risk of infection for local lobsters. In contrast, results concerning C. solearis are not consistent with those of the prior study by Briones-Fourzán et al. (2016). These authors found the highest levels of C. solearis prevalence in their “Vigía Chico” zone, whereas the present study found no significant effect of zone on the probability of infection with C. solearis. However, both studies found a positive relationship between lobster size and probability of infection with C. solearis.

Table 3 Estimates for logistic regression analyses testing the effects of size (carapace length, mm) and sampling zone (three levels: Reef, Lagoon, Shallow; reference level: Shallow) on the probability of finding spiny lobsters Panulirus argus: (A) clinically infected with PaV1, (B) testing positive to PaV1 by PCR, and (C) infected with metacercariae of the trematode Cymatocarpus solearis, in Bahía de la Ascensión, Mexico

| Effect                | Level       | Estimate | SE   | Wald   | p       | Odds ratio (95% CI) |
|-----------------------|-------------|----------|------|--------|---------|--------------------|
| (A) PaV1 clinical infection | Intercept   | 0.369    | 0.544| 0.461  | 0.497   | 0.93 (0.91, 0.96)  |
|                       | Size        | −0.070   | 0.013| 28.839 | <0.001  | 0.93 (0.91, 0.96)  |
|                       | Zone Reef   | 0.055    | 0.401| 0.019  | 0.892   | 1.06 (0.48, 2.32)  |
|                       | Zone Lagoon| 0.665    | 0.284| 5.480  | 0.019   | 1.94 (1.11, 3.39)  |
| (B) PaV1 by PCR       | Intercept   | 0.396    | 0.426| 0.863  | 0.353   | 0.98 (0.96, 0.99)  |
|                       | Size        | −0.024   | 0.007| 12.917 | <0.001  | 0.98 (0.96, 0.99)  |
|                       | Trematodes  | 0.126    | 0.221| 0.326  | 0.568   | 1.13 (0.74, 1.75)  |
|                       | Zone Reef   | 0.571    | 0.271| 4.439  | 0.035   | 1.77 (1.04, 3.01)  |
|                       | Zone Lagoon| 0.677    | 0.286| 5.605  | 0.018   | 1.97 (1.12, 3.44)  |
| (C) Trematodes        | Intercept   | −4.784   | 0.658| 52.828 | <0.001  | 0.0001             |
|                       | Size        | 0.047    | 0.010| 20.709 | <0.001  | 1.05 (1.03, 1.07)  |
|                       | PaV1 by PCR | 0.127    | 0.223| 0.325  | 0.569   | 1.14 (0.73, 1.76)  |
|                       | Zone Reef   | 0.312    | 0.286| 1.193  | 0.275   | 1.37 (0.78, 2.39)  |
|                       | Zone Lagoon| −0.237   | 0.291| 0.665  | 0.415   | 0.79 (0.45, 1.39)  |

In B and C, the effect of the other parasite (trematodes in B, PaV1 by PCR in C) was also tested (reference level: present)
Habitat characterization

The habitat characterization revealed substantial differences among the three sampling zones. The Reef zone had the greatest variability in both habitat complexity and cover of benthic components, but seagrass had the greatest cover (38%). These features reflect the location of this zone along the back reef, i.e., the interface between the reef lagoon and the coral reef structure. Results for Lagoon and Shallow were similar to those of Briones-Fourzán et al. (2012) for their larger sampling zones “Punta Allen” and “Vigía Chico”, which encompass Lagoon and Shallow, respectively. The Lagoon zone was vastly dominated by marine vegetation consisting mainly of seagrass, with interspersed macroalgae. In contrast, the Shallow zone consisted mostly of rather featureless, shallow bottoms, either bare rock (“calcareous pavement”) or covered with fine sand, or with small patches of short macroalgae.

Invertebrate biodiversity

The characterization of the invertebrate community is an important step into the search for potential vectors for PaV1 and first intermediate hosts for C. solearis. Most of the invertebrate species identified in the present study have been recorded in benthic communities of similar coastal systems in the wider Caribbean region (e.g., McNulty et al. 1962; Alcolado 1990), but substantial differences in community composition occurred among the three sampling zones. The invertebrate community differed more substantially in Shallow due to the dominance of two bivalves that were not recorded in either Reef or Lagoon, the mussel *Brachidontes modiolus* and the clam *Chione cancellata*. The former is a highly gregarious species that attaches to hard bottoms (Fields and Moore 1983) and was patchily distributed, whereas the latter is found in sandy and silty bottoms (McNulty et al. 1962). The particular habitat characteristics and invertebrate community composition of the Shallow zone are related to its location in the inner bay area, which is very shallow (< 2 m on average, Briones-Fourzán et al. 2012) and highly exposed to runoff and underground freshwater input (Medina-Gómez et al. 2014). In contrast, the habitat characteristics and invertebrate community composition of the Reef and Lagoon zones, located in the outer bay area, are more typical of coral reef systems, with higher richness and lower dominance values. These zones shared many species but differed in the relative abundance of species. However, the hermit crab *Pagurus brevidactylus* was one of the most abundant decapods in all three zones, whereas *Cerithium litteratum* and *Tegula fasciata* were the two most abundant gastropods in Reef and Lagoon. These species inhabit a wide range of habitats in other Caribbean reef and bay areas (e.g., Alcolado 1990).

Lobster size distribution

In Bahía de la Ascensión, the mean size of *P. argus* has been typically found to be larger in reef habitats and smaller in seagrass habitats of the outer bay area (i.e., between the land points and the reef; e.g., Lozano-Álvarez et al. 1991; Briones-Fourzán et al. 2012, 2016; Candia-Zulbarán et al. 2012; Huchin-Mian et al. 2013). However, the mean size of lobsters in the inner bay area, where the Shallow zone was located, appears to vary widely in time relative to zones in the outer bay area, being larger in some years (e.g., Lozano-Álvarez et al. 1991; Briones-Fourzán et al. 2012, 2016; Candia-Zulbarán et al. 2012; Huchin-Mian et al. 2013) and smaller in other years (Eggleston et al. 1990; Lozano-Álvarez et al. 1991; Eggleston and Lipcius 1992, the present study). Although a potential relationship with disease dynamics cannot be dismissed, the temporal variation in the mean size of lobsters in the inner bay is more likely to be related with movement pulses of lobsters triggered by the broad variation in environmental conditions typical of this shallow area, which is more vulnerable to storm waves and more exposed to underground freshwater input and high
evaporation (Arellano-Méndez et al. 2011; Medina-Gómez et al. 2014). Thus, when salinities decline in the inner bay, large local lobsters tend to migrate towards the outer bay area (Lozano-Álvarez et al. 1991), which is more stable in environmental conditions (Medina-Gómez et al. 2014).

**PaV1**

PaV1 has predilection for juvenile spiny lobsters (Shields and Behringer 2004), thus explaining why the probability of infection with PaV1 (both based on clinical signs and detected via PCR) significantly decreased with increasing size of lobsters. These results are consistent with previous studies concerning PaV1 in Bahía de la Ascensión (Briones-Fourzán et al. 2012; Candia-Zulbarán et al. 2012; Huchin-Mian et al. 2013) and elsewhere (Butler et al. 2008). However, Briones-Fourzán et al. (2012) found a significantly higher probability of infection for lobsters of all sizes in densely vegetated habitats than in poorly vegetated habitats. These authors hypothesized that if PaV1 particles are transmitted through water (e.g., Butler et al. 2008), then their persistence might be influenced by habitat exposure to UV radiation and the degree of shading provided by the local marine vegetation, i.e., that marine vegetation could be an environmental reservoir for PaV1 (Briones-Fourzán et al. 2012). Seagrass meadows in particular can attenuate water flow velocity (Peterson et al. 2004), which could potentially cause the pooling of virions, and in the present study the probability of clinical infection with PaV1 was highest in Lagoon, where the cover of seagrass was far greater. Interestingly, compared to the Shallow zone, the probability of lobsters testing positive to PaV1 by PCR was higher not only in Lagoon, but also in Reef, where the largest lobsters occurred. However, the cover of seagrass was also the highest among all benthic components in Reef due to its location in the interface with the reef lagoon, and it has been previously established that lobsters > 50 mm CL, including fully mature adults, can carry PaV1 in their system without necessarily developing clinical signs (Behringer et al. 2012; Huchin-Mian et al. 2013). Overall, these results lend support to the hypothesis of Briones-Fourzán et al. (2012), although the presence of viable virions in seagrass meadows requires to be tested, e.g., by means of environmental DNA techniques (Taberlet et al. 2018).

To date, *Panulirus argus* is the only known host for PaV1. Butler et al. (2008) investigated the presence of PaV1 by means of histology in individuals of three other species sympatric with *P. argus* (the spotted spiny lobster *Panulirus guttatus*, the channel crab *Maguimithrax spinosissimus*, and the stone crab *Menippe mercenaria*) 90 days after inoculating them with hemolymph of a PaV1-infected lobster, but found no evidence of PaV1 in any of them. In the present study, however, none of these three species were observed in the sampling zones and, to our knowledge, there have been no other attempts to find other potential hosts or vectors for PaV1. The invertebrate biodiversity assessment conducted in the present study could allow screening other species for the presence of PaV1, such as the small and very abundant hermit crabs observed in the reef and lagoon zones, using molecular techniques (e.g., endpoint PCR, Montgomery-Fullerton et al. 2007; qPCR, Clark et al. 2018).

**Cymatocarpus solearis**

In theory, the potential for trematode infection risk for lobsters may be higher in Shallow than in Reef and Lagoon, because the greater diversity of invertebrates in general and gastropods in particular in the two latter zones could result in a “dilution effect” (Civitello et al. 2015), whereby alternate hosts or predators may “intercept” miracidia (e.g., Laracuente et al. 1979) or cercariae (Thieltges et al. 2008; Orlofske et al. 2012). In the study by Briones-Fourzán et al. (2016), the probability of infection with *C. solearis* was significantly higher for lobsters in the site they called “Vigía Chico” (where the Shallow sampling zone is located) than for lobsters in two other more vegetated sites, but in the present study the sampling zone had no significant effect on this probability. However, in the study of Briones-Fourzán et al. (2016), lobsters from Vigía Chico had the largest mean size, whereas in the present study lobsters from Shallow had the smallest mean size. Also, Briones-Fourzán et al. (2016) found that the probability of infection with *C. solearis* significantly increased in the absence of clinical signs of PaV1, whereas no effect of presence of PaV1 on probability of infection by trematodes was found in the present study. These contrasting findings underline the complexity in potential relationships between lobster size, parasite infection, and features of local habitat and communities. On the other hand, as we only used visual inspection in the field, it is possible that we missed some lobsters infected with *C. solearis*, although the cysts of this parasite are large and easily detected by trained scientists. Estimating sensitivity of visual assessment of *C. solearis* prevalence would require comparing results of such assessment to results obtained via a complete dissection of a representative sample of lobsters over a broad size range. This would imply euthanizing the lobsters, which was not possible in the present study.

The characterization of the invertebrate community allowed the identification of some species that may act as first intermediate hosts for *C. solearis* and further explain its distribution. Digeneans use gastropods, bivalves, scaphopods and polychaetes as first intermediate hosts, but gastropods are the most common group (Cribb et al. 2003). All gastropods recorded in our sampling zones belong to superfamilies that have been identified as first intermediate hosts of trematodes in the order Plagiogriphiida (Cribb
lus fauna was rather dominated by the bivalves were overall very rare in the Shallow zone, where the mol-lobatus costatus, C. litteratum. Only C. l. costatus C. litt. act as first intermediate hosts of fasciata, C. litt. sp. Gastropods were overall very rare in the Shallow zone, where the mollusk fauna was rather dominated by the bivalves B. modiolus and C. cancellata. Whilst it is known that the family Brachycoeliidae does not use bivalves as first intermediate hosts (Cribb et al. 2001), other digenean metacercariae have been reported in several bivalve species (e.g., Lassalle et al. 2007). Taking this into account, C. litteratum and possibly T. fasciata are put forward as the most likely potential candidates for the first intermediate host of C. solearis.

Conclusions and future work

The present study builds upon previous studies by adding new information and providing further insight into the parasite/host system dynamics. For example, the findings of our quantitative analysis of cover of benthic components support the notion that marine vegetation could be an environmental reservoir for PaV1, and our logistic regression analyses provide further evidence that lobster size plays a vital role in the probability of infection by both PaV1 and C. solearis. A novel contribution of the present study was the analysis of the invertebrate community in our sampling zones, which allowed the identification of a few species as potential first intermediate host for the latter parasite and as targets for future research into potential vectors for PaV1, but also as a first step into understanding how species diversity in a given community may affect disease dynamics in these two parasite/host systems (Keessing et al. 2006; Johnson and Thieltges 2010). Since the conception of the present study, research surrounding PaV1 has progressed with new techniques which are not only able to detect, but also to quantify PaV1 in P. argus (Clark et al. 2018). In the near future, these techniques could be used to test for the presence of viable PaV1 virions in seagrass and other marine vegetation, as well as in decapod species associated with this type of habitat.

There is also much to discover surrounding C. solearis. First, it would be necessary to test whether C. litteratum, T. fasciata, or any other of the gastropod species herein identified indeed act as first intermediate hosts of C. solearis. This could be done by subjecting the gastropods to standard techniques inducing cercarial shedding (e.g., Morton 2018), which would allow to obtain and describe the cercariae of C. solearis, which remain undescribed. Although to the best of our knowledge C. solearis is the only trematode found in P. argus, the development and incorporation of molecular diagnostics for this trematode are important to determine whether other trematodes also infect P. argus, and to definitively determine the first intermediate host(s) (Théron 2014) as well as the presence of C. solearis in lobsters without the need to euthanize them. The potential effects of C. solearis on P. argus also merit investigation. For example, from a pathological viewpoint, histology could identify some immune reactions visually, whereas from a physiological angle, biochemical assays (e.g., Davies et al. 2015) could shed more light on this issue; however, molecular diagnostics would be needed to observe transcriptomic changes to get a full assessment of immunological effects.

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Compliance with ethical standards

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

Human and animal rights statement This article does not contain any studies with human participants performed by any of the authors, nor were any experiments performed on captive animals. All applicable international, national and/or institutional guidelines for the care and use of animals were followed. All procedures performed in the study involving animals were in accordance with the ethical standards of Mexican law regarding maintaining animal welfare in research.

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