Infection prevalence, intensity, and tissue damage caused by the parasitic flatworm, *Bdelloura candida*, in the American horseshoe crab (*Limulus polyphemus*)

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Abstract

Parasite infection dynamics can have profound implications on a host’s fitness; yet, there is a dearth of information on parasites in the American horseshoe crab (*Limulus polyphemus*) (Linnaeus 1758), a species that has experienced population declines in recent decades. Therefore, we aimed to quantify the prevalence, intensity, and gill surface area coverage of the ectoparasitic flatworm (cocoon and adult stages), *Bdelloura candida* in adult (*n* = 29), sub-adult (*n* = 7) and juvenile (*n* = 32) horseshoe crabs collected from Moriches Bay, NY (40.7810° N, 72.7171° W) in 2019 and 2020. Subsamples of horseshoe crab gill tissue (10%) were collected from live specimen, then *B. candida* cocoons were enumerated across the gill subsamples using microscopy while the extent of tissue damage was quantified with histology. *B. candida* was present in all adult and sub-adult crabs (100%), whereas juveniles exhibited 6.2% prevalence. Cocoon intensities per sample ranged from 28 to 805 cocoons, with 4.0–94.0% of gill lamellae harboring cocoons. In infected individuals, the total cocoon surface area coverage on gill tissues ranged from 0.06–14.51%, with higher cocoon intensities observed in the ventral-most gill quartiles relative to the dorsal-most gill regions. Sex was strongly supported as a primary driver behind *B. candida* infection intensities with adult females harboring higher intensities. Among infected gill lamellae, cocoon intensity was lower in mitochondrial-rich regions relative to mitochondrial-poor regions. These results provide novel insight into *B. candida* infection dynamics across horseshoe crab demographics, but further research is necessary to quantify the physiological impacts of the infection on *L. polyphemus*.

Introduction

The American horseshoe crab (*Limulus polyphemus*) is an iconic marine arthropod species that has persisted ~445 million years (Rudkin et al. 2008) and plays critical ecological roles in coastal marine systems from Maine, USA to the Yucatan peninsula in Mexico (Botton 2009; Smith et al. 2017). These ecological roles include bioturbation (sediment irrigation), structural habitat for epibionts, predation on marine bivalves and benthic macrofauna (Botton and Ropes 1987; Botton et al. 2003), and these animals provide a critical food source for migratory shorebirds, such as the endangered red knot (*Calidris canutus rufa*) (Tsipoura and Burger 1999; Smith et al. 2017). In addition to their numerous ecological roles, horseshoe crabs are an important human health and economic resource. The blood of horseshoe crabs serves to extract a unique compound, *Limulus* amebocyte lysate, (referred to as LAL) that is used to detect the presence of endotoxins from gram-negative bacteria in human medical supplies, and this practice has resulted in a threefold increase in biomedical harvest since the 1980s (Eyler et al. 2015; Smith et al. 2017). Moreover, roughly 1–2 million crabs are exploited annually for bait in the whelk and eel fishery throughout the US East Coast (ASMFC 2019). Exploitation from these industries has been perceived to be the primary contributor towards their recent coastwide declines, and thus, led to their “Vulnerable” conservation status in the US (Smith et al. 2016). Despite the current management interest, the impacts of biotic stressors (e.g. parasite infections) on horseshoe crab fitness and population dynamics have rarely been addressed, as management strategies have primarily focused on direct
anthropogenic exploitation. In particular, there is a dearth of information in regard to the role host-parasite relationships play on horseshoe crab fitness.

Parasites are ubiquitous in nature with most wild animals harboring at least a mild infection, with the prevalence and intensity frequently increasing with size, age, and density of the host (Zelmer et al. 1998). As established stressors, parasites are capable of influencing numerous aspects of their host’s biology including survival, fecundity, population cycles, and behavior (Lehman 1993; Hudson et al. 1998; Tompkins and Bergon 1999; Ebert et al. 2000; Poulin 2010) with cascading impacts on the entire ecosystem. Although variable between species, mild parasite intensities are typically well tolerated by a host (e.g. pinworm in humans) (Lehman 1993; Stjernman et al. 2008); however, intense infections can exacerbate this relationship, leading to adverse outcomes for a host. For instance, infections by roundworm (*Ascaris lumbricoides*) in the human intestine are considered to be mildly symptomatic, but the mere presence of several worms can cause blockages leading to indirect intestinal damage (Despommier et al. 2020). Similar observations have been made in aquatic animals, as many Monogenea (commonly observed ectoparasites of fish and amphibians) have been observed to damage their host not through resource exploitation, but via their attachment organs that penetrate the epithelial layer leading to excess mucus production and inflammation (Whittington 2005). Moreover, parasites infecting gill tissue have been demonstrated to adversely impact survival of aquatic organisms exposed to oxygen-poor conditions (Molnar 1994), with oxygen carrying capacity reduced up to 50% in crustaceans and fish (Taylor et al. 1996). Although several internal parasitic symbionts have been recorded in horseshoe crabs (Leibovitz and Lewbart 2003), the basic information on external parasites (i.e. prevalence, intensity and associated physiological impacts) such as the flatworm, *Bdelloura candida*, are not well established in the horseshoe crab literature.

*B. candida* is a 2 cm long triclad flatworm that is an obligate parasite of the American horseshoe crab. Adult worms are primarily located on the walking appendages and book gills, while the cocoons are attached on the inner surface of the gill lamellae by means of an endplate (Sluys 1989). Histological sampling has revealed that cocoons of triclad worms can cause necrosis of surrounding gill tissue (Groff and Leibovitz 1982; Leibovitz and Lewbart 2003). Moreover, few studies have quantified the extent of *B. candida* damage within gill tissue, which could be expansive. For example, Leibovitz and Lewbart (2003) documented over 50 *B. candida* cocoons can be attached to a single gill lamella in adult horseshoe crabs; albeit, their low sample size (< 9 adult horseshoe crab specimens) provided limited inferences about *B. candida* infection dynamics in horseshoe crab populations. Despite the limited knowledge surrounding *B. candida* infection, *B. candida* have been observed throughout the entire range of the American horseshoe crab (Riesgo et al. 2017). Although these studies have provided insight into *B. candida* infection characteristics, information pertaining to *B. candida* prevalence, intensities, and their coverage on horseshoe crab gills remains missing.

In this study, juvenile, sub-adult, and adult American horseshoe crabs were sampled from an estuary along the south shore of Long Island, New York, to better understand *B. candida* infection dynamics. The primary objectives of this work were to: 1) quantify *B. candida* intensity and prevalence across horseshoe
crab demographics (age groups and sex) and 2) investigate intrinsic factors that explain \textit{B. candida} intensities in horseshoe crabs. Additionally, we examined pathology inflicted by cocoons on horseshoe crab gill tissue via histological analysis and quantified the percentage of gill tissue occupied by \textit{B. candida} cocoons. We also assessed the infection intensity across the vertical gill space to determine if infection intensity was random (homogenous) or aggregated (heterogenous) across horseshoe crab gill space (ventral most or dorsal most). Lastly, we enumerated the proportion of \textit{B. candida} cocoons occupying the central- mitochondrial rich area (CMRA) vs. peripheral mitochondrial-poor area (PMPA) of each gill lamellae (Hans et al. 2018) to determine if there was a spatial preference for cocoon presence across these regions that provide different waste and respirational roles (Hans et al. 2018). This study provides novel insight into the infection dynamics of \textit{B. candida} on horseshoe crabs and could serve as the basis of monitoring ectoparasite infection in wild horseshoe crab populations.

**Materials & Methods**

**Sample collection**

On 24 June, 2019, juvenile (instars 8–10; n = 30) and adult (n = 29) horseshoe crabs were randomly collected at Pikes Beach, Moriches Bay, Long Island, NY (40.77°N, -72.71°W; Fig. 1). Juvenile crabs between instars 8–10 (prosomal width range = 41.4mm - 57.9mm) (Sekiguchi 1988; Carmichael et al. 2003) were hand collected haphazardly along a 500 meter transect at the water’s edge (< 0.5m depth), then transported back to the lab where they were euthanized using an overdose of Tricaine-S (MS-222) and frozen at -20°C for long term storage. Adults (prosomal width range = 188.0 mm- 283.0 mm) were collected from the intertidal zone (~ 0.5-1m deep) shortly before high tide and were temporarily placed in totes filled with ambient seawater. Sex, prosomal width, and weight were recorded for every individual. Sex was determined by the presence of modified pedipalps, weight was measured using a Pesola (Schindellegi, Switzerland) 10kg (± 0.3%) spring scale, and prosomal width was measured to the nearest millimeter using Vernalier calipers. The gills of adult crabs were subsampled by removing the upper-right portion of book gills, which constituted approximately 10% of their gills (Fig. 2). Both gill samples and \textit{B. candida} samples were individually stored in 70% ethanol to be counted at a later time. After sample collection, all adult crabs were released immediately back to the water.

Opportunistic samples of \textit{L. polyphemus} were obtained from a trawl survey in September-October 2020 from muddy habitat (~ 2m depth) and intertidal beaches of Moriches Bay (40.79°N, -72.71°W). This sample was comprised of 2 juveniles (n = 2; instars 9 & 10; prosomal width range = 40.5 mm – 57.2 mm) and a sub-adult cohort (n = 7; instars 14–18; prosomal width range = 113.0mm -179.0mm). Gill samples from these crabs were not removed, but instead were carefully examined for the presence of \textit{B. candida} cocoons and adult worms.

To obtain \textit{B. candida} adult intensity in a standardized format, forceps were used to remove \textit{B. candida} worms from the book gills and legs over a timed 5-minute period immediately following the physical measurements of the adult crabs. This served as a proxy for catch-per-unit-effort (CPUE). This time...
approach was implemented for two reasons: 1) every adult flatworm could not be accurately removed from the adults by hand or by brief freshwater rinses and 2) we wanted to minimize the desiccation stress to horseshoe crabs during sampling. All adult worms collected during each 5-minute sampling period were stored in 30ml of 70% ethanol in 50mL Falcon conical centrifuge tubes and were processed later in the lab. Prevalence of \textit{B. candida} was determined by tallying the presence of either adult worms or cocoons on horseshoe crab gill tissues or other appendages. Population prevalence was determined as the percentage of individuals that were infected by \textit{B. candida}.

\textbf{B. candida intensity measurements}

To measure adult flatworm intensity, collected adult flatworms were counted and recounted under a dissecting microscope by three readers. Samples were randomly chosen, and readers were blinded from previous recorded intensities in the same sample to minimize bias. The final intensity count was determined if two or more observers had the same intensity counts. If all counts differed between readers in a sample, a fourth reader counted the sample to finalize intensity based on agreement with another reader. Intensity was defined as the total count of adult \textit{B. candida}.

To measure the intensity of \textit{B. candida} cocoons, each individual lamella was removed from the book gill subset. \textit{B. candida} cocoons were enumerated under the dissecting microscope for each gill lamella in each gill sample subset. The intensity of cocoons was also enumerated separately on the central mitochondria-rich area (CMRA) and the peripheral mitochondria-poor area (PMPA) of the lamellae. Figure 2 visually illustrates CMRA and PMPA delineations.

\textbf{Gill and cocoon surface area preparations and measurements}

After measuring cocoon intensity, each lamella was placed on a lightbox with a ruler and photographed using a digital Panasonic LUMIX DMC-T380 waterproof camera. Every gill subset was sampled from the ventral to the dorsal side. However, lamella measuring less than 1 cm in diameter (typically the first few in the book gill) at its widest point were not photographed as they were not observed to harbor any infections and their small size contributed little to total respiratory surface area. One gill sample deteriorated during processing and could not be analyzed.

To measure the proportion of gill surface area covered by the cocoons, cocoon area and gill lamella surface area were measured using ImageJ (version 1.8.0) software (Schneider et al. 2012). The local threshold tool was used to automatically detect and measure the surface area of the cocoons against the lamella, which limited human error. In cases where the color threshold inadequately distinguished cocoons or lamella from each other, manual measurements were made in ImageJ. Average cocoon size was determined by randomly sampling 100 cocoon measurements across all individual adult crabs.

\textbf{Histological analysis}
A small gill sample (~2 cm x 1 cm x 0.5 cm) from a horseshoe crab, not used in the other analyses, was removed and placed in a histo-cassette then fixed in 10% buffered formalin, and embedded in paraaffin wax. Heavily infected sections of lamellae were selected to ensure the detection of *B. candida* cocoons. Sections (~5 µm) were mounted on slides and stained with Harris's haematoxylin and Eosin. Multiple slides were cut from the same paraaffin block, and slides were visually inspected for any signs of pathology, such as inflammation, necrosis or encapsulation.

**Statistical analyses**

A generalized linear model (GLM) was employed to determine which intrinsic factors explained the most variance behind cocoon intensity in adult horseshoe crab gill tissues. Only crabs sampled in 2019 were included in the statistical analyses because only prevalence was evaluated in the opportunistically sampled crabs in 2020. The response variable was cocoon intensity, and the explanatory variables were adult flatworm intensities, total gill surface area, and sex in the GLM. Prior to constructing the model, cocoon intensity data were fit to Poisson and negative binomial error structures and each distribution was compared by Akaike's information criterion (AIC) in the *fitdistrplus* R package (Delignette-Muller and Dutang 2015; R version 4.0.2, R Core Development Team 2020) to determine the most appropriate error structure given the data. In all GLMs, multicollinearity between the explanatory variables was assessed by calculating the variance inflation factor (VIF) in the *Performance* R package (Lüdecke et al. 2020). Variables with a VIF greater than 5 were removed from the GLM, as VIFs above 5 are considered to be moderately or strongly correlated with each other and strong correlations between one or more predictor variables can lead to erroneous and biased estimates (Gareth et al. 2013). Given that size is a sexually dimorphic trait in all horseshoe crab populations (Smith et al. 2009; Bopp et al. 2019) and it had the largest VIF in both GLMs, size was not included as an explanatory variable in the global models for cocoon and adult flatworm intensity candidate sets. Outside of size, other explanatory variable VIFs were <3 and were therefore, not removed from the global model prior to fitting and model selection analysis. The fit of all possible GLM model variants was assessed with the dredge function in the *MuMin* R package (Barton and Barton 2015). Inference was drawn from models within each GLM candidate set using the small sample size corrected Akaike Information Criteria (AICc) and AIC weights (Burnham and Anderson 2002). Model variants with ΔAICc < 2 and AIC weights > 0.10 were considered to have moderate support (Burnham and Anderson 2002).

A separate GLM was constructed to examine the relationship between adult flatworm intensity and the explanatory variables total gill surface area, sex, and horseshoe crab size. Similar to the other GLM, negative binomial and Poisson error distributions were fit to adult flatworm intensity data and were assessed with AIC. As aforementioned, the multicollinearity analysis was repeated, eliminating size, and model selection was again carried out using the dredge function.

To determine if intensity was homogenous or aggregated across gill tissue space (ventral to dorsal sections), each individual crab’s gill subsets were split into four quartile groups and the total cocoon intensity for each gill quartile was summed. A Friedman rank-sum test was used to determine if...
unreplicated blocks, while gill quartiles were considered groups in the Friedman model. If intensities differed between gill quartile groups in the Friedman test, multiple pairwise comparisons between the quartile groups were examined using a post-hoc Conover-Iman test (Conover and Iman 1979). A Bonferroni correction was used to control for multiple comparisons with an assumed familywise error rate of $\alpha=0.05$. The Conover-Iman test was conducted in the PMCMR R package (Pohlert 2014). A Spearman rank correlation test was also employed between the quartile group with the highest cocoon intensity (4th quartile) and the total cocoon intensity across gill samples to determine if the most infected gill quartile could serve as a relative proxy of overall cocoon infection intensity.

To determine if *B. candida* cocoons were spaced randomly or preferentially across mitochondrial regions of the lamellae in infected individuals, we calculated the average proportion of cocoons on CMRA regions vs. PMPA regions. Each of these regions can be assessed visually given their contrasting pigmentation on gill lamellae (Fig. 2C; Hans et al. 2018). Additionally, in order to determine if cocoon placement was proportional to the coverage of CMRA and PMPA regions within gill lamellae, CMRA regions of 100 random lamellae were randomly measured using ImageJ. Moreover, we employed a beta regression model (Ferrari and Cribari-Neto 2004) to determine if total cocoon intensities influenced the proportions of cocoons on the CMRA regions. A beta regression accounts for the fact that proportions are bound between 0 and 1, and error variances are not normally distributed. In the model, cocoon CMRA proportion was the dependent variable, and the total cocoon intensity was the independent variable. Beta regression models were fit using maximum likelihood estimation in the betareg package in R (Cribari-Neto and Zeilas 2010).

**Results**

**Prevalence and intensity**

From the horseshoe crabs collected in June 2019, *B. candida* infections (both adult worms and cocoons) were present in 100% of adult gill samples ($n=29$), while only 1 juvenile (instars 8–11) from this collection date ($n=30$) was observed to harbor any signs of infection (3 cocoons). The opportunistic samples procured in October of 2020, showed a similar pattern with 100% of sub-adults ($n=7$; instars 16–18) being observed to have either adult worms and/or cocoons. Of note, the two juveniles obtained in 2020 (i.e instars 9–11) were the least infected relative to all adults/sub-adults with one having no signs of infection and the other having a single cocoon. From the infected adult crabs, both cocoon intensity and adult worm counts varied considerably, as intensities of cocoons and flatworms ranged from 37 to 805 (mean = 267 ± 41 SE) and 5 to 196 (58 ± 9 SE), respectively. Adult female horseshoe crabs had an average of 306 ± 42 SE cocoons; whereas, males had a lower average cocoon intensity (85 ± 5 SE cocoons). Mean infection count was 3.2 ± 0.50 SE cocoons per lamella; however, total intensity varied across two orders of magnitude, ranging from 0.4 to 12.2 cocoons per lamella. Total number of gill lamellae (>1 cm diameter) also varied between crabs with counts ranging from 53 to 125 with an average of 91 ± 3 SE. Cocoon surface area differed by four orders of magnitude among adult crabs, covering gill subsamples, with 2.3% ±0.6% SE
representing the average (Fig. 3). The proportion of cocoons occupying the CMRA regions ranged from 0.05 to 0.53 ($x \pm 0.21 \pm 0.02$ SE) in infected adult horseshoe crabs (Fig. 4).

**Microscopic analysis**

*B. candida* cocoons were elliptical in shape with an anchor shape protrusion extending from one side, and they were attached to the lamellae epithelium via a cement-like excretion that surrounded the cocoon body and the anchor (Figs. 5C-D). After randomly sampling 100 single cocoons, the average cocoon area was $3.37 \pm 0.07$ SE mm$^2$. Histological observations showed healthy horseshoe crab leaflets consisted of parallel lamellae connected via pillar cells, with space between the pillars believed to be vascular channels (Fig. 5A). The dorsal tips of the leaflets were covered in a thickened proteinaceous matrix (Fig. S1), which also covered the epithelium of the leaflets, albeit not as thick as the blunted dorsal tips (Fig. 5A). Mats of bacteria and algae also frequently covered sections of gill epithelium but were superficial and not observed to cause any inflammation (Fig. 5A). Surface defects that impacted the organization of epithelial tissue and disrupted the typical gill structure were observed in histology sections of infected lamellae with melanization frequently observed on the outer layer of these lesions (Fig. 5E). Granulomas were frequently observed in the vascular lumen of the lamellae causing hemocyte aggregation, encapsulation and inflammation (Figs. 5B, E-F).

**Generalized linear model: cocoon intensity**

The negative binomial distribution was chosen for the cocoon intensity GLM, as the AIC was lowest for the negative binomial distribution relative to the Poisson distribution ($\Delta$AIC 4673.7). Therefore, a negative binomial was used as the family error structure in the candidate model set. The GLM model selection process revealed that three out of 8 model variants had moderate support (cumulative AIC weight $> 0.88$) according to $\Delta$AIC values and AIC weights (Table 1). Adult flatworm intensity was supported in all three models, but the top model had support for gill surface area as well (Table 1). There was also some support for sex influencing cocoon intensities (model 3, Table 1). The evidence ratio of the top model relative to model three was $\sim 1.9$ (Table 1), suggesting that the top model is nearly 2 times more probable than model 3. However, model selection uncertainty was present between the top three models given $\Delta$AIC $< 2$ (Table 1).
Table 1
Candidate model results for generalized linear model (GLM) for the cocoon intensity of adult horseshoe crabs after model fitting with the dredge package in R. A total of 8 possible model combinations were assessed, but only models with moderate support (Akaike weight > 0.10 and ΔAICc < 2) are shown here (see Methods) and are ranked by AICc. The cumulative Akaike weight between these models comprised 0.88. The global model included sex, adult worm intensity, and gill surface area, as explanatory variables with cocoon intensity set as the response variable. NP represents the number of parameters. The negative binomial distribution was the family error structure used in the GLM, as AIC was lower for the negative binomial distribution relative to the Poisson distribution (ΔAIC 4673.7).

| Model Number | Model                                      | NP | Log Likelihood | AICc | ΔAICc | Akaike weight |
|--------------|--------------------------------------------|----|----------------|------|-------|---------------|
| 1            | cocoon intensity ~ adult intensity + gill surface area | 4  | -181.5         | 372.8| 0     | 0.37          |
| 2            | cocoon intensity ~ adult intensity          | 3  | -183.1         | 373.1| 0.3   | 0.32          |
| 3            | cocoon intensity ~ adult intensity + sex    | 5  | -180.7         | 374.0| 1.2   | 0.20          |
| 4            | cocoon intensity ~ adult intensity + sex + gill surface area | 6  | -180.1         | 375.9| 3.1   | 0.08          |
| 5            | cocoon intensity ~ gill surface area        | 3  | -186.6         | 380.2| 7.4   | 0.01          |
| 6            | cocoon intensity ~ sex                      | 4  | -185.4         | 380.4| 7.6   | 0.01          |
| 7            | cocoon intensity ~ sex + gill surface area  | 5  | -184.0         | 380.7| 7.9   | 0.01          |
| 8            | cocoon intensity ~ 1                        | 2  | -189.8         | 384.1| 11.3  | 0.00          |

Generalized linear model: adult worm intensity

The negative binomial distribution was used as the family error structure in the GLM given it was a better fit (AIC = 293.9) compared to the Poisson distribution (AIC = 1166.2). Model selection revealed that 3 out of 4 possible model variants relating adult flatworm intensity to covariates had moderate to strong support (Table 2) with model selection uncertainty present. The top model only included sex and had an evidence ratio of ~ 3 relative to models 2 and 3, indicating this model is nearly 3 times more probable compared to the other models. Model support for gill surface area (models 2 and 3) was weaker compared to sex in terms of affecting adult flatworm intensities (Table 2).
Candidate model results for generalized linear model (GLM) for the adult worm intensity of adult horseshoe crabs after model fitting with the dredge package in R. A total of 4 possible model combinations were assessed, but only models with moderate support (Akaike weight > 0.10) are presented and are ranked by AICc. The cumulative Akaike weight between these models comprised 0.88. The global model included sex and gill surface area as explanatory variables with adult intensity set as the response variable. NP represents number of parameters. The negative binomial distribution was used as the family error structure in the GLM given it was a better fit (AIC = 293.9) compared to the Poisson distribution (AIC = 1166.2).

| Model Number | Model                                | NP | Log Likelihood | AICc | ΔAICc | Akaike weight |
|--------------|--------------------------------------|----|----------------|------|-------|--------------|
| 1            | adult worm intensity ~ sex           | 4  | -140.7         | 291.2| 0     | 0.56         |
| 2            | adult worm intensity ~ gill surface area + sex | 5  | -140.5         | 293.6| 2.4   | 0.17         |
| 3            | adult worm intensity ~ gill surface area | 3  | -143.4         | 293.8| 2.6   | 0.15         |
| 4            | adult worm intensity ~ 1             | 2  | -144.9         | 294.4| 3.2   | 0.12         |

**Spatial patterns of cocoon intensities**

Cocoon infection intensity differed among gill lamellae quartiles (Friedman-test; $F_{3,84} = 76.82, p < 0.001$). Specifically, cocoon intensities were different between all quartile group pairwise comparisons (Table 3). Gill quartile 4 (ventral most) had higher cocoon intensities relative to all other gill quartiles (Fig. 6). Gill quartile 4 exhibited a strong positive correlation (Spearman; $\rho = 0.96, p < 0.001$) with overall infection intensity in the gill subsample (Fig. 7).
Table 3
Post-hoc pairwise Conover-Iman comparison results among the four gill quartile groups following the Friedman test. Quartiles 1 and 4 denote the dorsal-most and ventral-most gill quartiles, respectively. V represents the Conover-Iman test statistic. The p-adjusted value represents the Bonferroni corrected p-value to minimize the inflation of familywise type 1 error. P-adjusted values < 0.05 denote statistical significance.

| Quartile comparison | V    | p adjusted |
|---------------------|------|------------|
| 1 to 2              | 12.54| < 0.01     |
| 1 to 3              | 25.09| < 0.001    |
| 1 to 4              | 29.00| < 0.001    |
| 2 to 3              | 12.54| < 0.01     |
| 2 to 4              | 16.45| < 0.001    |
| 3 to 4              | 3.90 | < 0.01     |

SUPPLEMENTAL

Figure S1. Histological cross-section of the acellular blunted lamella tips

Cocoon intensity on CMRA vs. PMPA

Total cocoon intensity appeared to weakly explain some of the variation behind the proportion of cocoons in the CMRA regions in adult horseshoe crab gill lamellae with cocoons more likely to occupy the CMRA regions with increasing infection intensity (beta regression; pseudo $R^2 = 0.14$, $p = 0.046$, df = 26, coefficient = 0.001). The average proportion of cocoons occupying the CMRA region of gill lamellae was 21.0% ± 2.0% SE. Out of the 100 random lamellae measurements, the average proportion of the CMRA region vs. total gill area was 30.6% ± 0.6% SE.

Discussion

This study demonstrates that *B. candida* infections are highly prevalent among adult horseshoe crabs, and that the cocoons of this parasite are capable of occupying a considerable amount of gill real estate (>10%). To our knowledge, the only previous studies evaluating *B. candida* infection intensities demonstrated that adult horseshoe crabs had 400–800 (average = 575, n = 4) cocoons across their entire gill area (Pearse 1949) and are capable of having > 50 cocoons within a single gill lamella (Leibovitz and Lewbart 2003). In contrast, the average *B. Candida* intensity was 267 in this study within our subsamples (10% of the total horseshoe crab gill area), and assuming that *B. candida* cocoon intensity patterns in the remaining 90% of the gill are homogenous, our results indicate average cocoon intensities could be much higher (average of 2,670 cocoons in entire gill space). Interestingly, cocoon prevalence and intensity were not uniform across life history stages, as only 6.2% (n = 2) of juvenile crabs (instars < 13; 58mm) had adults were observed to have cocoons and
adult worms. Moreover, all cocoon intensities were orders of magnitude higher on adult crabs than juveniles (when present). Ontogenetic shifts in parasite infrapopulation characteristics, such as the one we observed here, are not uncommon for a species where both prevalence and infection intensities increase with host body size and age. However, the mechanisms leading to the adult-juvenile dichotomy in *B. candida* prevalence observed is intriguing, as juvenile (instar groups 8–10) and adult crabs share the same habitat (spawning beaches) during the spring months possibly exposing the juvenile crabs to the parasite. Moreover, the high (100%) prevalence of *B. candida* in adult crabs suggests that *B. candida* is well adapted at colonizing a susceptible host with transmission unlikely to be limiting to juveniles; therefore, these age-group prevalence trends may presumably be a result of behavioral, foraging, or physiological differences between the stages.

A possible explanation for the ontogenetic differences in infection are due to the decrease in molting frequency observed in horseshoe crabs as they age. Juvenile horseshoe crabs molt several times a year until reaching instar 10 (49.2mm), after which molting occurs annually until a terminal molt is reached upon sexual maturation (Carmichael et al. 2003, Estes et al. 2015). Following the terminal molt, the accumulation of epibionts on horseshoe crabs (slipper snails, barnacles, macroalgae, etc.) are well documented (Walls et al. 2002) with similar dynamics likely to influence the establishment of *B. candida*. For example, the antiparasitic effects of molting have been observed in Antarctic krill (*Euphausia superba*), as recently molted krill had 0% prevalence of ectoparasites as opposed to pre-molt individuals that had a 66% prevalence (Tarling and Cuzin-Roudy 2008). Moreover, this study also found parasite prevalence increased with host age, presumably a result of decreased molting frequency. Further support of molting as a mechanism of parasitic defense was demonstrated in *Daphnia magna*, as molting was found to limit the adhesion of bacteria and subsequent infection (Duneau and Ebert 2012). Similarly, prevalence and intensity of epizootic shell disease in the American lobster were shown to increase in animals with lower molting frequencies (egg-bearing females) and this was suggested to result from the inability of infected lobsters to eliminate pathogenic microbes during molting (Castro et al. 2012).

Although molting seems to be a likely mechanism in *B. candida* regulation, it is difficult to test, as the most heavily infected cohort (adults) rarely molt and are difficult to maintain in laboratory conditions in sufficient numbers to test such a hypothesis.

Outside of molting, other phenomena such as horseshoe crab behavior, ontogenetic differences in size, physiology and resource use may explain prevalence dynamics across age groups. For instance, it is possible that *B. candida* is sexually transmitted and initiates infection during the extensive copulatory process observed in horseshoe crabs in which multiple males may attach to one female for days to months each spring (Brockmann and Penn 1992). However, the observation of 100% prevalence infections among the sub-adults (n = 7, instars 16–18) in this study makes this an unlikely scenario, as this age group exhibits different space-use patterns relative to adults, and they do not engage in mating behavior (Rudloe et al 1981). Differences in gill surface area between instars 8–10 and sub-adults/adults may also be a primary factor behind prevalence disparities, as parasite intensity can correlate with increasing body size such is the case with Salmon louse (*Lepeophtheirus salmonis*) infecting Atlantic salmon (*Salmo salar*) presumably a result of increased surface area.
reducing space competition among conspecific ectoparasites. Additionally, the body size argument may also explain why horseshoe crab sex was a contributing factor in *B. candida* infection rates because adult females are larger than males (Loveland and Botton 1992) and thus, females have more available surface area or “habitat” for *B. candida* to reside. However, host size has not been found to be a limiting factor in ectoparasite infection intensities in some organisms; whereas, host whole body metabolism can be a more important determinant of ectoparasite intensities (Hechinger et al. 2019) because host energy can be more constraining to parasite infection loads.

The difference in infection intensities among age groups may partially reflect contrasting foraging behaviors among life history stages and may make the juvenile cohort (<12 instars) unsuitable for *B. candida* establishment. For example, *B. candida* is believed to indirectly consume food particle remnants from horseshoe crab feeding activities (Jennings 1977). Juvenile crabs (instars < 10) predominantly rely on sedimentary organic matter and meiofauna adjacent to salt-marsh habitats (Botton et al. 2003b); in contrast, older juvenile and adult crabs predominantly forage on larger-bodied prey, such as bivalves and polychaetes (i.e. *Nerites spp.* (Botton and Ropes 1987; Gaines et al. 2002)). Therefore, the nutritional resources on juvenile horseshoe crabs may not be sufficient or optimal for *B. candida's* nutritional requirements. However, the theory of *B. candida* foraging on remnants of horseshoe crab prey items remains controversial due to chemical analysis indicating that *B. candida* may obtain some nutritional energy directly from horseshoe crabs (Lauer and Fried 1977). The application of modern techniques to assess resource-use, such as stable isotope analysis (bulk or compound-specific), could be used to resolve this controversy, as it could identify the nutritional resources adult *B. candida* predominantly relies on.

Given that this study emphasized one population of horseshoe crab hosts, we cannot state these infection intensity patterns apply to other populations, as other factors such as biogeographic differences in reproduction ratios, environmental conditions, migration patterns, abundance, and size may result in varying *B. candida*. For example, host population density is often positively correlated with ectoparasite intensity as a result of increased probability of direct transmission (e.g. contact, breeding, etc.) among conspecifics within a population after controlling for other covariates (Arneberg et al. 1998). However, controversy surrounds the contribution of population density to increased parasite infection intensities. Bagge et al. (2004) noted that the primary determinant behind infection rate variability for multiple Monogenean species’ in crucian carp (*Carrasius carassius*) was host population size, presumably due to a required infection density threshold for effective transmission, and thus numbers of hosts were the limiting transmission factor. For horseshoe crabs, size-at-maturation and population densities are the largest in Mid-Atlantic populations (Delaware and Chesapeake Bays) and can be 2-400 times greater than their northern counterparts (Shuster 1955; James-Pirri 2005; Smith et al. 2009, 2017 ). In turn, *B. candida* intensity may contrast between host populations due to both disparate population densities, abundances, and body size differences of *L. polyphemus*. Previous studies have demonstrated that ectoparasite intensities increase with host body size in a variety of animal species, including chigger parasites on the Spiny lizard (*Sceloporus clarkii*), (Watkins and Blouin-Demers 2019), multiple species of Woodpeckers in Brook trout (*Salvelinus fontinalis*) (Poulin et
Therefore, assessing *B. candida* population dynamics between horseshoe crab populations with different characteristics (e.g. abundance, density, etc.) may be beneficial for elucidating mechanisms that regulate ectoparasite-host relationships, particularly in such an ancient and stable host species.

Within an individual host or even an organ, parasites are known to aggregate in regions that provide the best niche for them and in turn higher fitness, this was observed in the cocoons of *B. candida* in this study as cocoons of were significantly more prevalent in the dorsal most quartile of gill lamellae (Fig. 6). The ventral most gills were preferentially used for cocoon placement, possibly due to the larger size of these lamellae as larger lamellae size not only provides more habitat, but also allows cocoon location to be away further from edge of the lamellae sheltering the cocoons from excessive flow, a critical concern for ectoparasites (Wootten 1974). Additionally, larger lamellae can pump more water, which may be necessary to meet the oxygen demand of the flatworm cocoons which require oxygen to sclerotize (Huggins and Waite 1993). Unsurprisingly, the realized niche of a parasite is frequently smaller than the potential niche (Sukhdeo and Croll 1981), resulting from constraints on attachment, competition and nutrient acquisition, factors that can lead to hyper specialization within a larger organ. For example, the gills of fish are often segregated between parasites such as Monogenean flatworms or parasitic copepods which will localize to particular gill arches in fish (Arme and Halton 1972; Teemer et al. 2020). Similar results were observed in this study as *B. candida* cocoons were infrequently placed in the CMRA’s, a specialized zone of the lamellae important for nitrogenous waste excretion (Henry et al. 1996; Hans et al. 2018). However, we postulate that cocoon placement is fairly random across the CMRA vs. PMPA sections of horseshoe crab gills because the CMRA region in this study comprised an average area of 0.30 ± 0.60 SE across the total gill surface while the average proportion of cocoons was 0.21 ± 0.02 SE, indicating that the placement of *B. candida* cocoons is nearly proportional to the CMRA. It is important to note that, the CMRA cocoon placement was not entirely avoided in this study and the likelihood of CMRA cocoon placement appeared to slightly increase with cocoon intensity, albeit, the beta regression results indicated a weak relationship between overall cocoon intensity and the proportion of cocoons in the CMRA region. Therefore, in other horseshoe crab populations it is important to determine if the spatial arrangement (random or clustered) of cocoons on gill lamella varies across *B. candida* intensity levels.

The extent of the deleterious impacts imposed by *B. candida* infections remains uncertain; however, horseshoe crab fitness could be affected from the combination of anthropogenic and ambient environmental stressors coupled with *B. candida* infection. This study revealed no more than 15% of gill surface area in any adult was covered with *B. candida* cocoons, however this estimate is likely conservative as our analysis was unable to detect the cocoon cemented regions (Fig. 5C, D). Regardless of this potential underestimation, light infection intensities on gills from ectoparasites may have adverse impacts on horseshoe crab fitness. For example, ectoparasite coverage on gills appears to be directly proportional to reduction on the velocity of oxygen uptake in aquatic organisms (Duthie and Hughes 1987) and may potentially affect horseshoe crab fitness by reducing respiration efficiency, especially in hypoxic conditions. Hypoxic conditions are expected to become more chronic and frequent in coastal marine environments in the coming decades (Diaz and Rosenberg 2008), and the combined stress of *B.
hypoxic conditions (<2.0 $\text{O}_2 \text{mgL}^{-1}$) and parasitic nematode (*Anguillicola crassus*) infections over 4 days, eels with low swim bladder degenerative indexes (0–1) and the highest infection loads exhibited shorter time until death (10–25 hours shorter on average) than their uninfected counterparts (Lefebvre et al. 2007). Additionally, horseshoe crabs face a unique and direct anthropogenic stressor, in the form of blood extraction for biomedical purposes, that may make individuals with intense infections of *B. candida* more susceptible to sublethal effects (i.e. reduced oxygen uptake, increased respiration energy expenditure, etc.) or mortality (Smith et al. 2017; Owings et al. 2019, 2020). The sublethal impacts of biomedical blood extraction on horseshoe crab survivors (~70% survival) are numerous, such as a significant reduction in hemocyanin concentrations following typical blood volume extractions (30% blood volume) and reduced spawning frequency (Owings et al. 2019, 2020), and it can take up to 4 months for amebocytes to fully recover to baseline levels (Novitsky 1984). Hemocyanin is essential for maintaining oxygen transport (Mangum 1980), immune response (Coates et al. 2011), wound repair and molting (Adachi et al. 2005) and can be altered by environmental conditions (Coates et al. 2012). Additionally, these amebocytes are involved in the immunological response to *B. candida* cocoons (Fig. 5), so the impairment of these cells could reduce the crabs’ ability to respond to immunological insults and/or increase baseline level of stress of these animals and potentially increasing susceptibility to stressors. Therefore, understanding the simultaneous impacts of projected intensifying environmental conditions (i.e. temperature, ocean acidification, and hypoxia), biomedical blood harvest on hemocyanin levels, and *B. candida* infections are required, as their combined effect may engender more severe consequences for horseshoe crabs than when these effects are isolated.

**Conclusions**

This study provides novel insight into *B. candida* infection characteristics across multiple life history stages of *L. polyphemus*, and the results suggest there is a moderate dichotomy of *B. candida* prevalence between instars <12 and sub-adult/adult crabs. This study also demonstrated that cocoon infection intensity and surface area coverage differed substantially in adults, indicating infection is considerably variable. This study only focused on one *L. polyphemus* population and patterns of *B. candida* infection may not be uniform across all horseshoe crab populations. Because parasite abundance can vary due to differences in geographical location, host density, ambient environmental characteristics, and distance between host populations (Poulin et al. 2011), future studies quantifying the *B. candida* intensity and prevalence across other geographically distinct horseshoe crab populations are needed. Additionally, other underlying mechanisms (i.e. horseshoe crab population density, age group composition, etc.) of *B. candida* infection dynamics should be addressed at the population level to understand both host-parasite ecology and the evolutionary underpinnings of *B. candida* infection. Monitoring *B. candida* infection in horseshoe crabs is recommended given that histological analysis indicated *B. candida* may adversely impact gill functioning and could adversely affect host fitness, especially if *B. candida* intensity is higher in populations outside of this study. Based on the findings, overall infection intensity strongly correlates with the 25% ventral-most gill lamellae (Fig. 6), and thus, monitoring programs could expedite intensity enumeration by examining these gill lamellae sections. Considering the vulnerable status of *L.*
polyphemus in the U.S., the impact of B. candida infection on horseshoe crab respiration coupled with environmental stressors on survival should be examined to determine if this host-parasite relationship is contributing to recent population declines.

**Declarations**

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**Data availability:** The datasets and code generated and analyzed during this study are available from the corresponding authors on request.

**Competing interests:** The authors declare no competing interests.

**Ethics approval:** organisms and tissue samples collected for this research were permitted through by the New York State Department of Conservation under the Scientific License to Collect #1145 under the New York State Environmental Conservation Law.

**Informed consent:** No human subjects were used in this study and informed consent was, therefore, not applicable.

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Figures
Figure 1

Map of Moriches Bay in Long Island, NY, USA. The black star represents Pike's beach, the intertidal sampling location for juvenile and adult crabs 2019. The red star denotes the trawl survey location where sub-adult crabs were opportunistically sampled in 2020. (Base map source: NOAA, ESRI)
Figure 2

A) Book gill (10%) removed from an adult horseshoe crab, in which the ventral most lamellae is heavily infected with B. candida cocoons. B) an individual gill lamella with moderate B. candida infection (>25 cocoons). C) The central mitochondria-rich area (CMRA) is located within the mid-section of the lamellae and has a darker pigmentation relative to the more transparent peripheral mitochondria-poor area (PMPA).
Figure 3

Histogram of the percentage of gill surface area covered by cocoons in infected adult horseshoe crabs (n=29). The vertical dashed line represents the average.

Figure 4
Histogram showing the proportion of B. candida cocoons within the central mitochondrial-rich area in the gill lamellae (n=29). Vertical line denotes the mean proportion of cocoons on CMRA (0.21 ± 0.02 SE).

Figure 5

A) Normal gill histology of gills from an area not impacted by B. candida cocoons, nonpathogenic biofilms are present on the acellular layer of the chitinous epithelium stained black (scale bar= 500 μm). B) Granuloma present in lamellae with intense hemocyte infiltration and inflammation (scale bar= 200 μm). C) B. candida attached to a lamellae (scale bar= 1mm). D) B. candida removed from a lamellae with adhesion substance surrounding the cocoons and pointed out with arrows (scale bar= 1mm). E&F) stained cross-section of a wound caused by B. candida, melanization was observed (arrow) with
hemocytes aggregating along the inner surface of the epithelial layer (E scale bar = 100 \mu m; F scale bar = 200 \mu m)

Figure 6

B. candida cocoon intensity for four gill lamellae quartile groups. The first quartile represents the dorsal-most gill lamellae; in contrast, the fourth quartile represents the ventral-most gill lamellae. Letters represent statistically significant differences between gill lamellae quartile groups from Conover post-hoc pairwise comparison tests shown in Table 3. Black dots represent each individual horseshoe crab’s cocoon intensity in each gill lamellae quartile.
Figure 7

Scatter plot of cocoon intensity from the 4th quartile of gill (ventral-most) leaflets (n=27) vs. total gill cocoon intensity. The Spearman rank correlation results were: $\rho = 0.96$ and $p = <0.001$. Standard error is represented by grey shading.

Supplementary Files

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