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## Autecology and Genetic Diversity of *Cletocamptus* Copepods in Coastal Hypersaline Lagoons of Puerto Rico

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**ABSTRACT**—Copepods of the genus *Cletocamptus* are common inhabitants of coastal saline lagoons of the greater Caribbean. They are abundant in southern Puerto Rico, where past studies indicate the presence of one species, *Cletocamptus dominicanus*; but genetic and ecological studies have yet to be conducted. For a year, we surveyed the *Cletocamptus* populations of three lagoons (Candelaria, Fraternidad, and Playa Sucia) in southwestern Puerto Rico. Densities fluctuated within and between lagoons, suggesting patchy distribution. Their average density per lagoon ranged from 0 to 608 copepods/cm<sup>2</sup> of sediment. Salinity, temperature, and pH were not correlated with copepod densities, suggesting tolerance to environmental fluctuations. Specimens exhibited morphological variability, motivating us to assess the standing genetic variability and the potential of cryptic species by employing two molecular markers. The mitochondrial COI and the nuclear ribosomal 28S genes were sequenced from 130 and 118 specimens, respectively, collected from Candelaria, Fraternidad, and Playa Sucia, as well as from an additional four lagoons (Petrona, Encarnación, Providencia, and Flamenco). Phylogenetic analyses using both genes indicate the presence of at least four lineages. Based on COI sequences, corrected sequence divergence between the lineages ranged from 13% to 43%. Given the large sequence divergence among the *Cletocamptus* lineages, we used single-locus species delimitation tools (GMYC, bPTP, mPTP, ASAP, and ABGD) to assess for the presence of additional species beyond the previously reported *C. dominicanus*. Most delimitation tests grouped taxa into five putative species. *Cletocamptus* spp. are often the numerically dominant benthic metazoans in these important coastal habitats and warrant taxonomic, genomic, and physiological studies to understand their divergence and adaptations to this unique environment.

Copepods are the most abundant metazoans in the marine environment and are highly diverse, with ~14,000 described species (Turner 2004; WoRMS Editorial Board 2024). They fulfill essential roles in marine and freshwater ecosystems by promoting bioturbation by reworking sediments, driving food web dynamics by linking primary producers to larger metazoans, and having important roles in the global carbon and nitrogen cycles (Hicks and Coull 1983; Frangoulis et al. 2004; Schratzberger and Ingels 2018). They are ubiquitous throughout the planet, and are found in nearly all habitats with water, including extreme environments like the deep sea, glaciers, and hypersaline waters (Kikuchi 1994; Anufriieva 2015; Rosli et al. 2018).

Harpacticoida alone has ~3,600 described species (Suárez-Morales 2015), and estimates for total diversity could reach 150,000 species (Seifried 2004). Within

the Harpacticoida, members of the genus *Cletocamptus* (Fig. 1) are of interest for their remarkable ability to inhabit a range of salinities, from freshwater (Gómez et al. 2007) to extreme environments like hypersaline lagoons (Suárez-Morales et al. 2013; Anufriieva 2015; Gómez et al. 2017; Yakovenko et al. 2024). These euryhaline organisms are found in many habitats worldwide, most commonly in estuaries and saline lagoons (Gómez et al. 2004, 2013; Anufriieva 2014). This is observed in the greater Caribbean, where they are considered the most abundant harpacticoid inhabiting coastal saline lagoons (Gómez et al. 2017).

Taxonomic work and species assignment within the genus is difficult due to incomplete or erroneous species descriptions in the early years (e.g., *C. dominicanus*, *C. deitersi*, and *C. albuquerqueensis* had to be redescribed by Gómez et al. (2017)). *Cletocamptus*



FIG. 1. Confocal laser microscopy of *Cletocamptus* specimens found in this study. Specimens stained using Congo red following the protocol of Michels and Büntzow (2010). Each copepod is labeled with its id; specimens 15\_13 and 3\_70 are from Lineage 1, 0\_220 is from Lineage 1c, and 13\_81 is from Lineage 3, as labelled in the COI ML phylogeny.

also has cryptic species (Rocha-Olivares et al. 2001), and some species exhibit high intraspecific morphological variability (Fleeger 1980; Castro-Longoria et al. 2003; Gómez and Yáñez-Rivera 2022). Gómez and Yáñez-Rivera (2022) have done the most comprehensive morphological phylogenetic work in revising the genus, with 27 recognized species.

Only a single species, *Cletocamptus dominicanus*, has been reported in the coastal lagoons of Puerto Rico, where they have been found in salinities of up to 87.6 psu (Gómez et al. 2017). Additionally, only two publications, viz. Castro-Longoria et al. (2003) and Rocha-Olivares et al. (2001) focus on the genetic diversity within the genus *Cletocamptus*. This study aims to enhance our understanding of the ecology and systematics of *Cletocamptus* copepods in Puerto Rico by sampling several coastal lagoons and leveraging molecular phylogenetic tools to delineate lineages of *Cletocamptus* from the island.

## MATERIALS AND METHODS

### *Area of study*

The first portion of this work investigates the population dynamics of *Cletocamptus* in three lagoons within the Cabo Rojo National Wildlife Refuge, namely Candelaria (CL), Fraternidad (FL), and Playa Sucia lagoons (SL) (Fig. 2I) (Permits DNER 2020-IC-059 and USFW 2020-04). These lagoons within the Cabo Rojo National Wildlife Reserve are an important stop-over in the Caribbean for migrating shorebirds (Collazo et al. 1995), including the stilt sandpiper (*Calidris himantopus*) and the black-necked stilt (*Himantopus mexicanus*) (Parks et al. 2016). The lagoons are heavily impacted, as they are thought to have been used for salt extraction since 700 AD. Their topography has been altered to make evaporation basins, where the water level is managed for salt production rather than left to fluctuate naturally.



FIG. 2. Area of study; I. Inset map includes lagoons of the Cabo Rojo National Wildlife Refuge used for the demographic portion of the study. Lagoon sites are represented by the first letter of their names (C = Candelaria, F = Fraternidad, S = [Playa] Sucia) along with letters (A, B, and C) indicating specific sampling areas; II. The island of Puerto Rico and the lagoons included in the genetic analyses.

The second portion of this work investigates the genetic diversity of *Cletocamptus* copepods in shallow coastal lagoons with elevated salinities at the time of sampling, including the aforementioned Cabo Rojo lagoons (> 41 psu) and four additional sites, namely, Providencia (45 psu), Encarnación (75 psu), Petrona (61 psu), and Flamenco lagoons (40 psu) (Fig. 2II). All are located on the southern coast of Puerto Rico, except for Flamenco Lagoon, which is located on Culebra Island, about 25 km East of the eastern coast of Puerto Rico.

### Demographics

Samples were collected from Candelaria, Fraternidad, and Playa Sucia lagoons (Fig. 2I) at a randomly selected date each month, from January 2020 to February 2021, with a gap in April–June 2020 due to the COVID-19 crisis (total of 10 sampling dates). Each lagoon was sampled in three areas (denominated A, B, and C), considering accessibility to the lagoon and intending to sample different areas (Fig. 2I).

Three sediment samples were collected with a sediment corer (5 cm<sup>2</sup> area, 10 cm<sup>3</sup> volume) from three areas of each lagoon, i.e., nine sediment cores were collected from each lagoon at each sampling time. For easier accessibility, only the lagoon borders were sampled, where water depths ranged from 8 to 15 cm. Each core was stored in a 50 ml centrifuge tube, then sifted with a 125 µm sieve and preserved in 95% EtOH with rose bengal to facilitate visualization of meiofauna. Each

core was then divided into eight subsamples using a modified Jensen meiofauna sample splitter (Jensen 1982). Two of the subsamples from the meiofauna splitter were inspected with a Nikon SMZ800 dissection microscope. Individuals of *Cletocamptus* were counted and categorized as female, gravid female, male, or copepodid. Nauplii were seldom seen due to the sieve size (125 µm) used and are not included in this study. Three abiotic parameters (temperature, pH, and salinity) were measured in all lagoon areas at each sampling time. Temperature and salinity were measured in situ with a field thermometer and refractometer; pH was measured with a SevenEasy Metler Toledo pH meter in the laboratory using water samples from each lagoon area.

The data from the two subsamples were extrapolated to estimate the abundance of the whole sample by multiplying the counted copepod abundances by four since there were eight subsamples in the meiofauna splitter. This was used to generate an Euclidian distance resemblance matrix. The matrix was used for a PERMANOVA using PRIMER v. 7 (Anderson et al. 2008; Clarke and Gorley 2015), to test whether the factors *date*, *lagoon*, and *area* (nested in lagoon) affected the variance of *Cletocamptus*' abundance. A biotic and environmental matching analysis (BIOENV) within PRIMER was also conducted to correlate environmental variables (salinity, pH, and temperature) with the abundance of *Cletocamptus*. BIOENV is a multivariate correlation analysis weighted by season (rainy vs. dry).

After examining collected copepods under a dis-

TABLE 1. Primer pairs used in this study.

Primer Pairs (5'–3')	Gene	Product Size	Annealing Temperature	Author(s)
dgLCO1490: ggtcaacaaatcataaagayatygg dgHCO2198: taaacttcagggtgaccaaaraayca	COI	614 bp	47° C	(Meyer 2003)
LCO1490: ggtcaacaaatcataaagatattgg Cop-CO1-2189R: gggtagacaaaaaatcaraa	COI	580 bp	45° C	(Folmer et al. 1994) (Bucklin et al. 2010)
mlCO1intF: ggwacwggwtgaacwgtwtayccycc jgHCO2198: taiacytciggrtgicraaraayca	COI	313 bp	54° C	(Leray et al. 2013) (Geller et al. 2013)
28S-F1a: gcggaggaaaagaaactaac 28S-R1a: gcatagtttcaccatctttcggg	28S	800 bp	51° C	(Ortman 2008)

secting microscope, we noticed high morphological variability, which suggested the possibility of more than one *Cletocamptus* species. This observation motivated us to use DNA sequencing to test whether several genetic lineages/species of *Cletocamptus* are present in the coastal lagoons of Puerto Rico.

#### **Molecular work: DNA extraction, amplification, and sequencing**

Copepods were collected from the sediment of the three Cabo Rojo Lagoons, three additional coastal lagoons of southern Puerto Rico and one lagoon from Culebra Island (Fig. 2) (Permits DNER 2020-IC-059 and USFW 2020-04). Sediment was obtained by scraping the lagoon bottom with a bottle, the sediment was then sifted in situ with a 125 µm sieve and preserved with 95% EtOH. To extract genomic DNA, individual copepods from each location were transferred from EtOH storage to dH<sub>2</sub>O for about 10 minutes at room temperature to eliminate the majority of EtOH, a possible amplification inhibitor. Each sample was labeled with its location code (0: Cabo Rojo National Wildlife Refuge (i.e., Fraternidad, Candelaria, and Playa Sucia lagoons); 3: Providencia 6: Encarnación; 13: Petrona Point; and 15: Flamenco lagoon on Culebra Island) and the extraction number. Individual copepods were placed in 0.5 ml microcentrifuge tubes with ~20 µl of 5% Chelex 100 or InstaGene Matrix (Bio-Rad Laboratories). Samples were cooled on ice for about 10 minutes, vortexed for 15 seconds, quickly centrifuged to accumulate material at the bottom of the tube, heated to 99° C for 10 minutes, vortexed for 15 seconds, and finally centrifuged at 12,000 rcf for 3 minutes. Afterwards, the Chelex resin and the exoskeleton of the copepod precipitates on the bottom of the tube, and the DNA contain-

ing supernatant is used for PCR. The exoskeleton was retrieved and preserved in formalin for morphological observations.

Genetic variability among the processed specimens was tested using two genes with differing evolutionary drivers (Hellberg 2009). The molecular markers used are a mitochondrial and a nuclear ribosomal gene, i.e., cytochrome *c* oxidase subunit I (COI) and large nuclear subunit (28S) rRNA. COI is the de facto marker in biodiversity studies of metazoans and has been successfully used in population genetics and low-level phylogenetic studies (Rocha-Olivares et al. 2001; Machida et al. 2006; Adamowicz et al. 2010; Garlitska et al. 2012). The 28S gene is more appropriate for higher-level phylogenetic studies (more conserved), but could still provide important insights when closely related cryptic species/lineages are expected to be discovered when used with other molecular markers. Various COI primers (see Table 1) were used due to the difficulty amplifying some specimens. PCR was performed using Bio-Rad T100 Thermal Cyclers. A reaction volume of 20 µl consisted of 10 µl of Kapa Taq ReadyMix 2x (Roche), 0.15–0.45 µl of 10 mMol primer (reverse and forward), 1–3 µl of template DNA, and PCR grade H<sub>2</sub>O to complete the desired volume. If reaction volumes changed, components changed proportionally. Annealing temperature gradients were performed to verify optimal annealing temperatures. PCR cycles included: initial denaturation at 95° C for 3 minutes; 35 cycles of denaturation at 95° C for 30 seconds, annealing for 30 seconds (see Table 1 for annealing temperatures), and extension at 72° C for 1 minute/kb; the last step is a final extension at 72° C for 1 minute/kb. The success of PCR amplifications was verified using gel electrophoresis on a 1% agarose gel run at ~74 V for 30 minutes. Amplicons were sent

to McLab Sequencing Facility (San Francisco, CA) for Sanger sequencing.

### **Sequence processing**

The resulting DNA traces were inspected and edited using CodonCode Aligner v. 10.00.02 or AliView (Larsson 2014). Individual sequences were blasted using the blastn tool on NCBI (Altschul et al. 1990) to test for homology and possible contamination. All contaminated or low-quality sequences were discarded (about 70%). A sequence alignment for each gene was then created using the online version of MAFFT v. 7 (Kato et al. 2019) and trimmed with trimAl (Capella-Gutiérrez et al. 2009). Available *Cletocamptus* COI sequences (AF315014, AF315012, AF315009, AF315011, AF315010; Rocha-Olivares et al. 2001) were also included in the analyses. *Enhydrosoma curticauda* (MH976577), *Stenocaris* sp. (MF077882), and *Laophontodes* sp. (MF077876) were used as outgroups for phylogenetic analysis of COI. *Cletodidae* sp. (MF077804), *Stenocaris* sp. (MF077862), and *Paralaophonte meinerti* (KR048898) were used as outgroups for the phylogenetic analysis of 28S. Except for *Stenocaris* sp., no other harpacticoid species existed with NCBI records for both genes; instead, we used outgroup taxa that at least belonged to the same families. All sequences produced from this study were submitted to GenBank (Appendix 1).

### **Phylogenetic inference**

ModelFinder (Kalyaanamoorthy et al. 2017) was used, within IQ-TREE (Minh et al. 2020) to estimate the best-fit nucleotide substitution model for each dataset according to the Bayesian Information Criterion (BIC). Single gene phylogenies were estimated using maximum likelihood (ML) analysis with 100 non-parametric bootstraps. Multigene phylogenetic analysis was made by concatenating COI and 28S sequences with SequenceMatrix v. 1.9, partitioning the data by gene and constructing the phylogeny with IQ-TREE. The resulting trees were visualized using iTOL (Letunic and Bork 2021). The major clades from the COI ML phylogeny were identified, and then the ‘between group mean distances’ of these clades were calculated using MEGA11 (Tamura et al. 2021). Analyses were performed with 1,000 bootstrap replications, using p-distances and Tamura 3-parameter nucleotide substitution model, allowing for transitions and transversions, alpha

parameter of Gamma distribution equal to 1, heterogeneous patterns among lineages, and pairwise deletion of gaps/missing data.

### **Species delimitation**

Various molecular tests were performed to delimit possible species using the COI dataset, i.e., ASAP (Puillandre et al. 2021), ABGD (Puillandre et al. 2012), mPTP (Kapli et al. 2017), bPTP (Zhang et al. 2013), and GYMC. Online versions of ABGD (<https://bioinfo.mnhn.fr/abi/public/abgd/abgdweb.html>) and ASAP (Puillandre et al. 2021; <https://bioinfo.mnhn.fr/abi/public/asap/asapweb.html>) were used with the JC69 model and default parameters. Online versions of mPTP (<https://mptp.h-its.org/#/tree>; Kapli et al. 2017) and bPTP (<https://species.h-its.org/>; Zhang et al. 2013) were used. The bionj tree from IQ-TREE was used as input for mPTP. A BI consensus tree constructed with MrBayes was used for the bPTP method. MrBayes v. 3.2.7 (Huelsenbeck and Ronquist 2001) was used for BI, analysis was run for  $3 \times 10^6$  generations, sampling every 1,000 chains, with a 25% burn-in. To run a GMYC (Generalized mixed Yule-coalescent) test (Fujisawa and Barraclough 2013), ultrametric trees were generated using BEAST2 with different parameters, i.e., yule model with constant clock, yule with relaxed clock, and coalescent model with constant population and constant clock, all with a rate of evolution set to 1 (Michonneau 2016), with 10,000,000 reps and with the BModelAnalyzer package. Consensus trees were generated using TreeAnnotator with 10% burning. GMYC was implemented using the *splits* v. 1.0.20 (Ezard et al. 2021) package for R v. 4.2.2 (RStudio Team 2020; R Core Team 2021).

## RESULTS AND DISCUSSION

### **Demographics**

*Cletocamptus* copepods were found in three lagoons of the Cabo Rojo National Wildlife Refuge (Candelaria, Fraternidad, and Playa Sucia lagoon; Fig. 2I), and their abundances and demographic composition were measured. Large fluctuations of copepod densities were observed at all lagoons throughout the one-year study (Fig. 3). High standard deviations for the total average density of copepods at each lagoon, 224 (CL), 51 (FL), and 244 (SL), demonstrate high variability. This variability is also seen at the lagoon area level, exemplified by the standard error bars in Fig. 3.

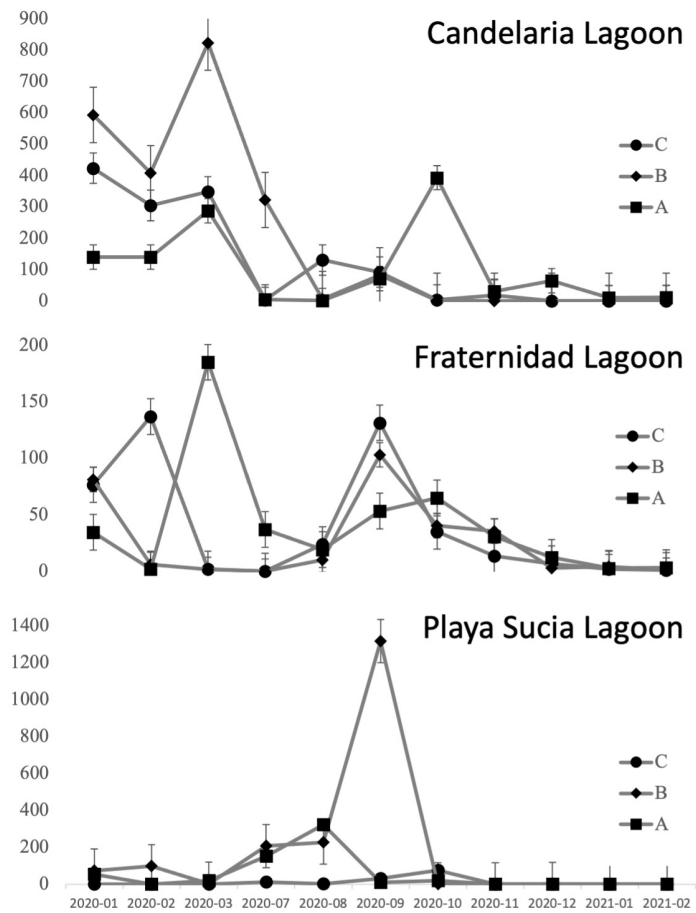


FIG. 3. The average number of *Cletocamptus* spp. per cm<sup>2</sup> in each lagoon area through time.

### *Candelaria Lagoon (CL)*

The highest density of copepods at any lagoon was observed for CL, 608 copepods/cm<sup>2</sup>, in March, 2020 (Fig. 4). CL had the highest overall density (total average) of copepods with 144 copepods/cm<sup>2</sup> throughout all sampling times. CL was the most productive (in terms of *Cletocamptus* copepods per area) out of the three lagoons. CL had an average of 17 gravid females per cm<sup>2</sup> in January 2020 (Fig. 4). CL also had the highest percentage of non-gravid females out of all lagoons (79.8%), but the lowest percentage of gravid females (2.7%), copepodids (8.1%), and males (9.4%) (Table 2).

### *Fraternidad Lagoon (FL)*

The highest average density at FL was found in September, 2020, with 120 individuals/cm<sup>2</sup> (Fig. 4). FL had an overall density (total average) of 35 copepods/cm<sup>2</sup> throughout all sampling times, making it the least populated of the three lagoons. FL had an average of 27 gravid females per cm<sup>2</sup> in September 2020. Frater-

nidad Lagoon had the lowest percentage of non-gravid females at 54.5%, however, it had the highest percentages of gravid females (12.9%), copepodids (18.1%), and males (14.5%) (Table 2).

### *Playa Sucia Lagoon (SL)*

The highest average densities for Playa Sucia Lagoon were observed in September, 2020, the same sampling date as FL, with 565 individuals/cm<sup>2</sup> (Fig. 4). SL had an overall density (total average) of 81 copepods/cm<sup>2</sup>, making it the second most populated of the three. SL had an average of 19 gravid females per cm<sup>2</sup> in July, 2020. If abundance sums for SL are turned into percentages, we can see that all demographic categories fall between the other lagoons. Non-gravid females at 67.7%, gravid females account for only 3.5%, copepodids 16.1%, while males formed 12.8% of all *Cletocamptus* counted for this lagoon (Table 2).

All lagoons have different and fluctuating densities of copepods; non-gravid females are the major demographic category, followed by copepodids, males, and gravid females (Fig. 4). Copepods are patchily distributed, as exemplified by the heterogeneous column heights, and there is a stark decrease in abundance at all lagoons beginning in October, 2020 that persists to the end of sampling in February, 2021 (Fig. 4). During this period, we observed the lowest densities (e.g., 4 (CL), 2 (FL), and 0 (SL) copepods/cm<sup>2</sup>).

Using abundance data, the percentage of non-gravid females is 72.6%. Gravid females account for only 4.3%, copepodids reach a total of 12.0% of all copepods counted, while males only 11.1% (Table 2). Females were the largest demographic representative for all lagoons through time, with few exceptions (Figs. 4, 5). This is nearly a third of all copepods counted throughout this investigation. This pattern is common, as benthic harpacticoids with r-population strategies and large populations have been seen to have larger female-to-male ratios (Dahms and Qian 2004).

Even though abundances approached zero near the end of sampling, total average copepod densities were higher than what is commonly seen in other harpacticoid studies in surface sediment of shallow aquatic ecosystems, where average densities are lower than 30 copepods/cm<sup>2</sup> (Jayabarathi et al. 2015) with harpacticoids dominating (51%).

### *Statistical inference*

In brackish systems, the abundance of harpacti-

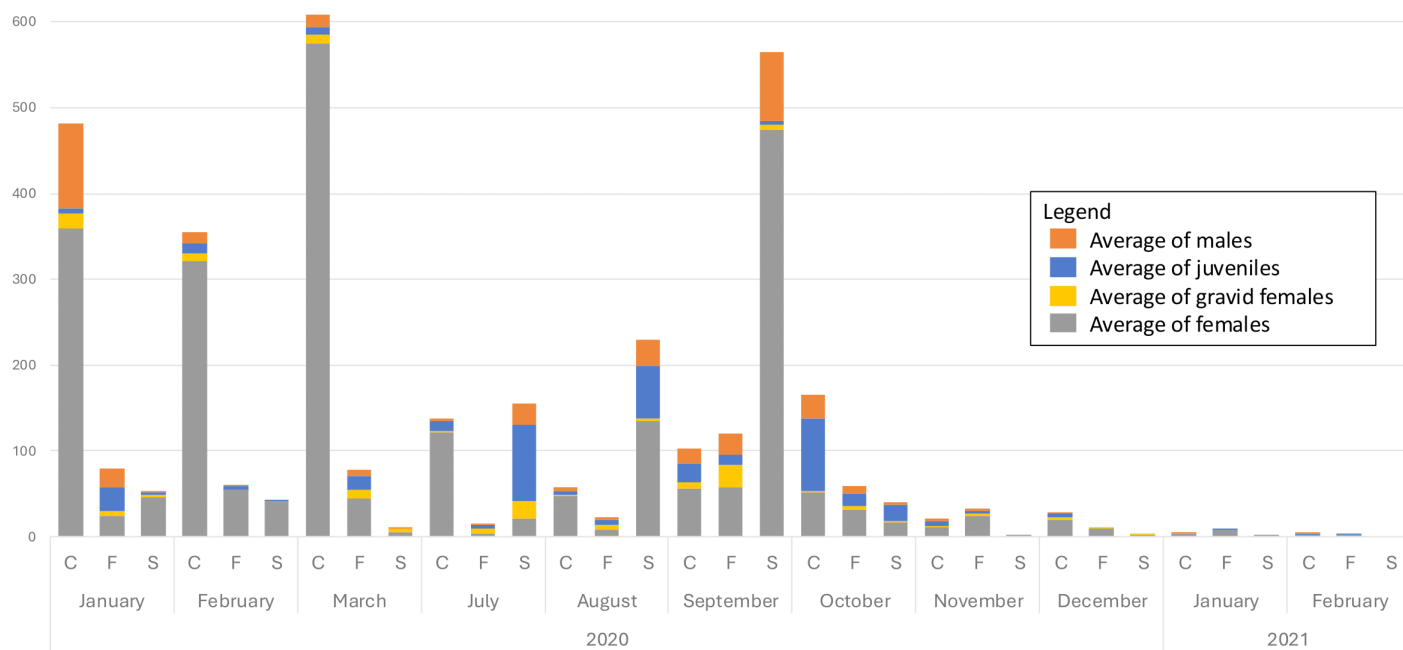


FIG. 4. Density (copepods/cm<sup>2</sup>) of *Cletocamptus* spp. at each lagoon at all sampling times.

coid copepods is positively correlated with increasing salinity. Copepods are less abundant in oligohaline regions, indicating a preference for higher salinity levels (Yamamuro 2000). In tropical estuaries and coastal lagoons, copepod densities are generally lower due to significant salinity fluctuations and temperature stress (Armenteros et al. 2007). Temperature plays a crucial role in the ecology of copepods as high temperatures can increase oxidative and physiological stress, affecting their reproduction and survival (Von Weissenberg et al. 2022). pH also influences copepod populations, as low levels can significantly reduce the egg production rates of certain copepod species. For example, at a pH of 7.6 compared to 7.9, the egg production rate of some copepod species decreased approximately two-fold when combined with increased temperatures (Lee et al. 2020). Furthermore, it is hypothesized that copepods may survive in hypersaline environments by acquiring exosmolytes through their diet, i.e., by eating osmolyte-containing algae (Shadrin and Anufriieva 2013; Anufriieva 2015). Copepod densities in shallow coastal lagoons are affected by interactions of temperature, salinity, and pH, and the ability of copepods to survive in hypersaline environments may be habitat/community dependent.

At the Cabo Rojo lagoons, we measured temperatures ranging from 22 to 42° C, salinities from 27 to 150

psu, and pH from 7.25 to 9.03 (Figs. 6–8; Table 3; and for averages of all lagoons at all sampling times Appendix 2). These parameters reach extreme values, creating a challenging environment for its inhabitants, and should significantly influence the abundance of *Cletocamptus* in these lagoons. However, a BIOENV analysis shows that the interaction of salinity, temperature, and pH did not correlate with the abundance of copepods ( $Rho = 0.043$ ,  $P = 0.87$ ). The same analysis using only single physicochemical variables resulted in even lower correlations. That is to say, the abundance of *Cletocamptus* copepods was not correlated to any of the parameters measured, e.g., events of high salinity or temperature. A PERMANOVA test indicated there is a low probability that the observed variability of copepod densities is explained by the factors (date, lagoon, area) included in this sampling design. What best explained the variability in the abundance of *Cletocamptus* is the date x area (nested in lagoon) interaction ( $p = 0.001$  and Pseudo-F = 13.211), meaning that the variability in the abundance of copepods in each area of each lagoon was independent of sampling time.

A wider sampling grid, i.e., a higher sampling effort, may have detected temporal trends by lessening the effects of patchy distributions. Notwithstanding, the absence of significant statistical results pertaining to the interaction of factors and correlations of physicochem-



TABLE 2. Demographic composition of each lagoon throughout all sampling dates.

Lagoon	Percentage (%) of			
	Females	Gravid Females	Copepodids	Males
Candelaria	79.8	2.7	8.1	9.4
Fraternidad	54.5	12.9	18.1	14.5
Playa Sucia	67.7	3.5	16.1	12.8
Total	72.6	4.3	12.0	11.1

TABLE 3. The highest and lowest parameters for each lagoon where *Cletocamptus* was present. The sampling date is included in parentheses.

Lagoon	pH		Temperature (° C)		Salinity (psu)	
	lowest	highest	lowest	highest	lowest	highest
Candelaria	8.02 (10/2020)	9.03 (01/2020)	27 (11/2020)	42 (07/2020)	27 (11/2020)	124 (02/2020)
Fraternidad	7.25 (02/2020)	8.89 (12/2020)	22 (02/2020)	38 (07/2020)	36 (01/2021)	115 (03/2020)
Playa Sucia	7.89 (08/2020)	8.97 (01/2020)	25 (11/2020)	41 (10/2020)	43 (11/2020)	95 (07/2020)

ical factors with copepod densities in this study can be attributed to temporal cycles in population size (unrelated to seasonal fluctuations), microtopography of the lagoons, sediment particle size (De Troch et al. 2006), and the patchy spatial distribution of benthic copepods. Food availability and dissolved oxygen could also affect the abundance of the small crustaceans (Alcocer et al. 2015). It can also be caused by dormancy (Dahms and Qian 2004), where resting eggs lie dormant during unfavorable periods. Additionally, abundance trends may have been obfuscated by the presence of more than one *Cletocamptus* species in our samples, as we found from our genetic surveys, which inhabit the same ecosystem but may occupy different niches with different population trends.

### Phylogenetic inference

One hundred thirty (130) sequences of COI and 118 sequences of 28S were generated (GenBank Accession Numbers: COI: OQ682279–OQ682408; 28S: OQ819183–OQ819299). After editing and trimming, the final lengths of the COI and 28S sequences were 623 and 865 bp, respectively. ModelFinder suggested K3Pu+F+I+G4 (BIC score 11715.8) for COI and TN+F+G4 for 28S (BIC score 7499.7) as the best-fit substitution models according to the Bayesian Information Criterion. ML gene phylogenies were construct-

ed to test for the presence of one or more lineages of *Cletocamptus*, based on the observed morphological variability. *Cletocamptus* from the Candelaria, Fraternidad, and Playa Sucia lagoons (of Cabo Rojo) and four other coastal saline lagoons (Providencia, Encarnación, Petrona, and Flamenco lagoons) that covered a longitudinal gradient spanning from the southwesternmost part of the main island to the easternmost part of the island archipelago were included (Fig. 2II). Additionally, we included five COI sequences from GenBank generated by Rocha-Olivares et al. (2001). The resulting phylogeny grouped local *Cletocamptus* into five major lineages (Fig. 9). One hundred and eighteen 28S sequences generated from this study were also analyzed phylogenetically (Fig. 10). The 28S ML phylogeny is more conserved and does not recover an identical structure/grouping as COI, yielding four lineages, instead of five. This has been observed in other studies where the phylogenetic structure of the tree changes with different marker genes (González et al. 2020). The generated ML multigene (COI+28S) phylogeny recovers the same major lineages as COI (Fig. 11).

The major lineages recovered by the genetic analysis were used to create groups and estimate their genetic distances. Corrected genetic distances between these groups are high and range from 13 to 43% (Table 4). A 3% and 9% COI divergence is considered a threshold

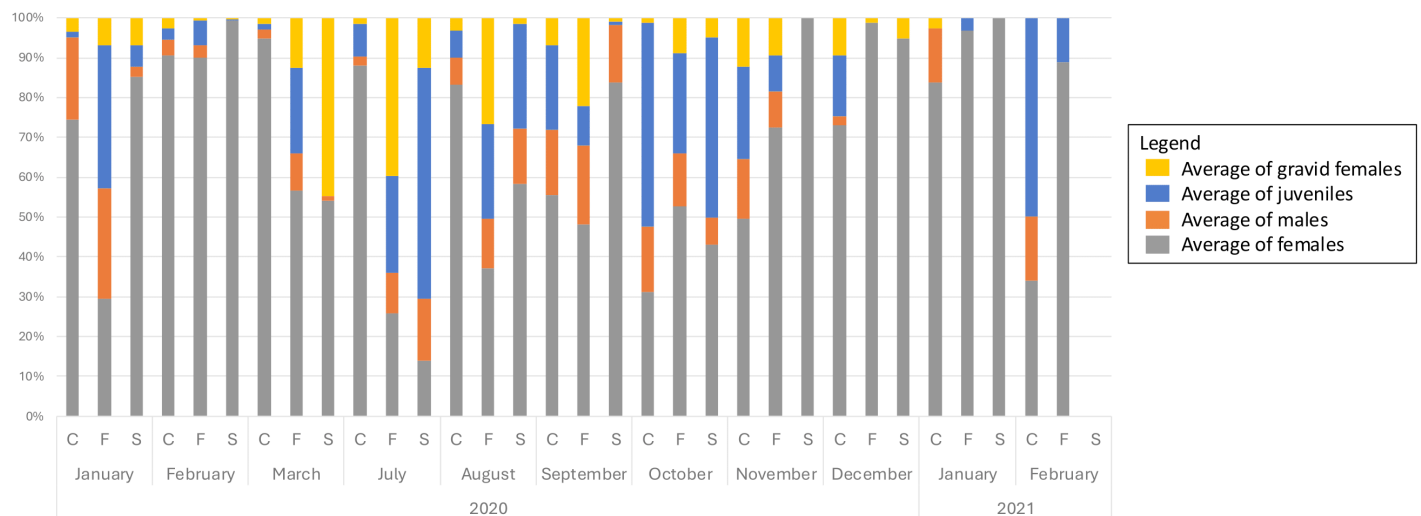


FIG. 5. Percentages of females, gravid females, copepodids, and males of *Cletocamptus* spp. sampled at all lagoons at each sampling time.

to call putative species for Coleoptera and Lepidoptera, respectively (Hebert et al. 2003; Ma et al. 2022). However, genetic distances between harpacticoid copepod species are generally higher than other arthropod taxa. The genetic distances among the *Cletocamptus* lineages concord with those between *Cletocamptus* species (Rocha-Olivares et al. 2001; Table 4) and between species of other copepod genera, such as *Enhydrosoma*, *Nanopus*, and *Tigriopus*, indicating that the major lineages recovered by the COI and multigene phylogenies could be putative species.

The multigene tree with the *Cletocamptus* specimens linked to their geographic coordinates can be seen in Fig. 11. Lineage 3 is composed solely of copepods from Punta Petrona and was recovered by both COI and 28S gene phylogenies; the recovery of all other lineages varied between genes. All lineages except Lineage 3 are found in more than one sampling location. Lineage 4 is the most broadly distributed, having a presence in all sampling locations. The Cabo Rojo lagoons host Lineages 1, 2, and 4 (Table 5). All lagoons host three lineages, except for Punta Petrona, which hosts two. Most copepods from Flamenco lagoon of Culebra comprise Lineage 1, however there are also members of Lineages 4 and 5 represented. The presence of three lineages in Culebra suggests that there are no effective spatial barriers to limit genetic flow between locations of the main island of Puerto Rico and Culebra Island, which is geographically isolated by a ~35 km stretch of ocean. The mechanisms of gene flow for *Cletocamptus* across lagoons and across Caribbean islands are

unknown. However, many migrating waterbirds visit these lagoons, e.g., shorebirds (Collazo et al. 1995), ducks, and egrets; it has been suggested that birds may disperse copepod eggs undigested through their feces, or whole copepods in their feet and plumage (Green and Figuerola 2005; Frisch et al. 2007; Hessen et al. 2019). Dispersal through birds is a promising hypothesis since coastal lagoons have little connection to ocean currents and *Cletocamptus* spp. as benthic harpacticoid species have obligatory benthic nauplii, making dispersal through this route highly improbable.

From an ecological perspective, the potential for spatial barriers to drive genetic differentiation and speciation in aquatic organisms such as copepods has important implications for conservation and management. Understanding the distribution and connectivity of populations can inform conservation efforts by identifying areas of high genetic diversity and evolutionary potential. Furthermore, the ecological context in which organisms live can also affect their evolution and diversity. In the case of *Cletocamptus*, the saline lagoon habitat may play a critical role in driving the genetic differentiation of these copepods. Understanding which factors drive genetic differentiation can help us better understand how species evolve and adapt to changing conditions over time.

### *Species delimitation*

Species delimitation tests were conducted using 135 COI sequences (Fig. 9). The ASAP delimitation test creates partitions that range from nine to 22 mo-

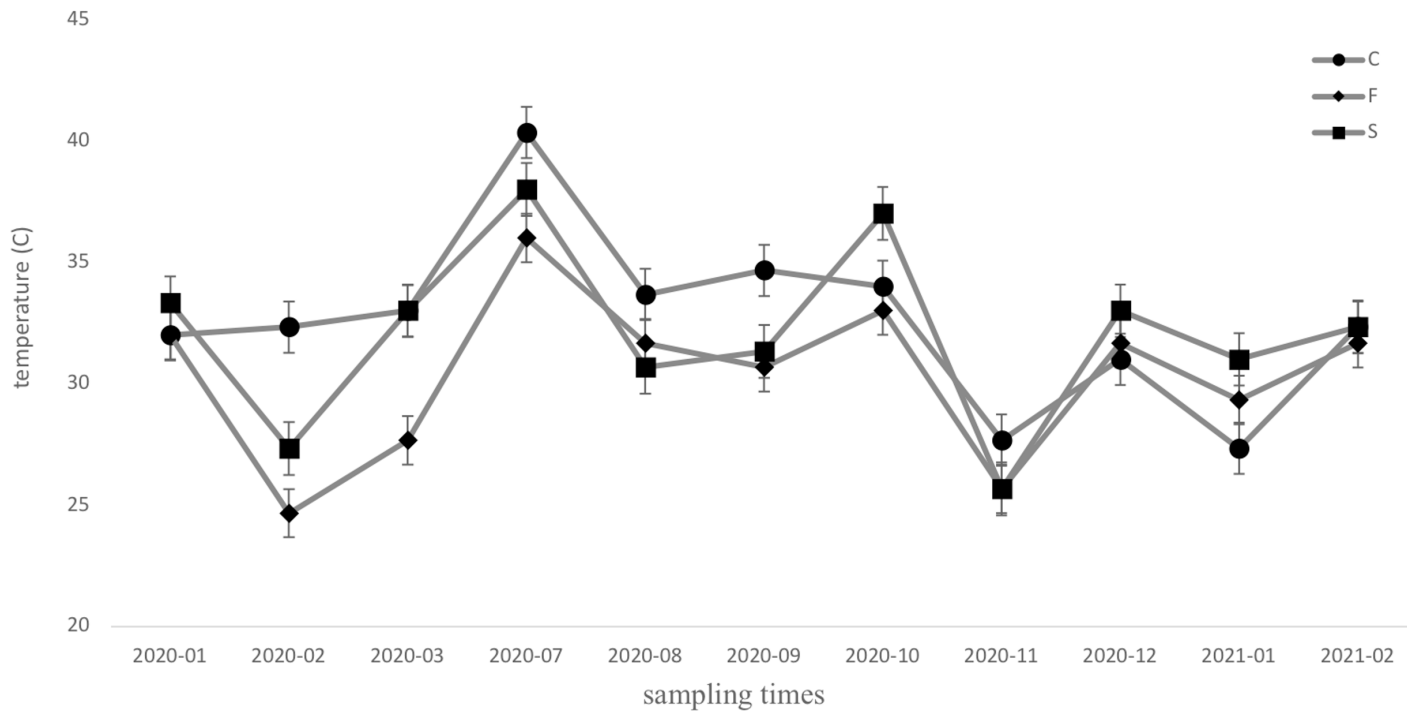


FIG. 6. Average temperatures for Cabo Rojo lagoons at each sampling time; C = Candelaria; F = Fraternidad; S = [Playa] Sucia Lagoon.

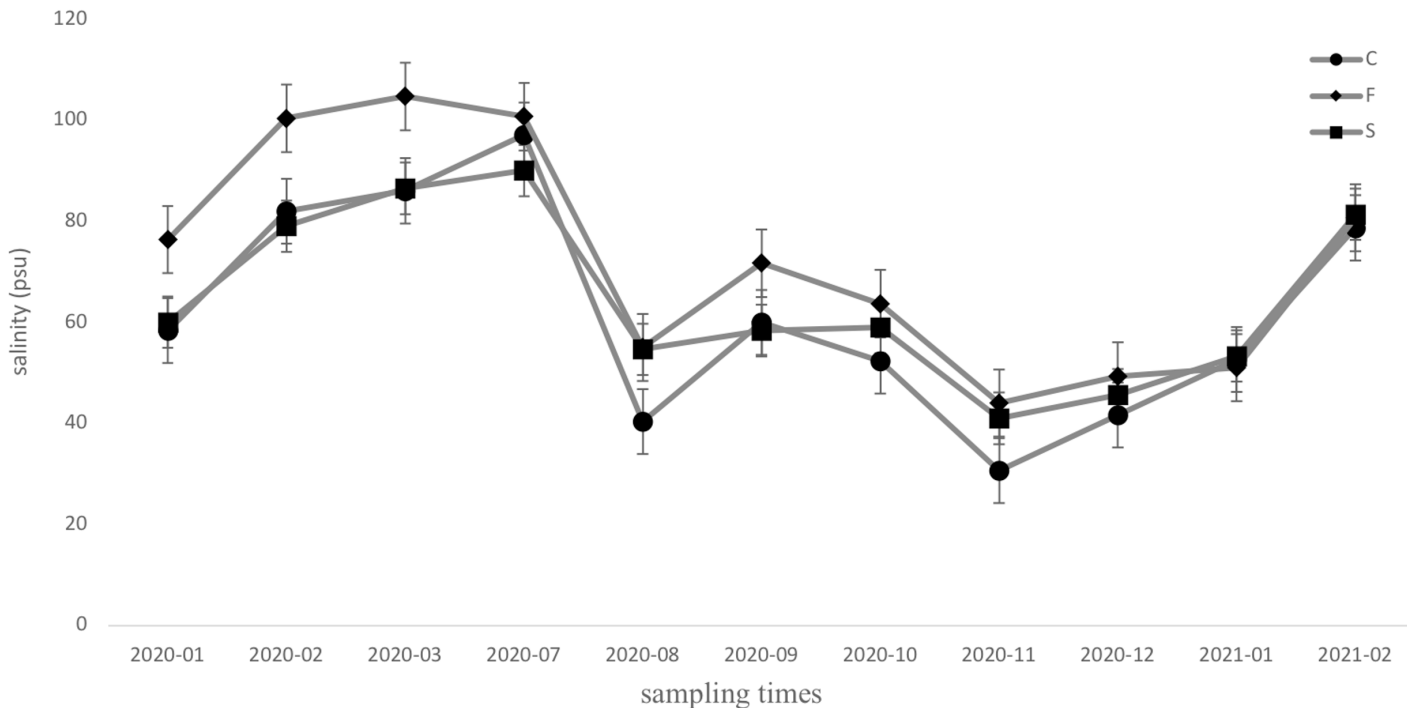


FIG. 7. Average salinity (psu) for Cabo Rojo lagoons at each sampling time; C = Candelaria; F = Fraternidad; S = [Playa] Sucia.

lecular operational taxonomic units (motus). The best scoring partition, asap score of 2.00 and P value of 1.00e-5, splits all samples into nine motus: local specimens into five and the Rocha-Olivares et al. (2001) into four. All 10 initial partitions of ABGD divide taxa into

10 motus, four recursive partitions split taxa into 14, 15, 25, and 32 motus. The highest scoring partitions suggest 10 motus: five for local specimens and five for the Rocha-Olivares et al. (2001) specimens (P value of 1.00e-01). mPTP splits the data into nine motus: five for

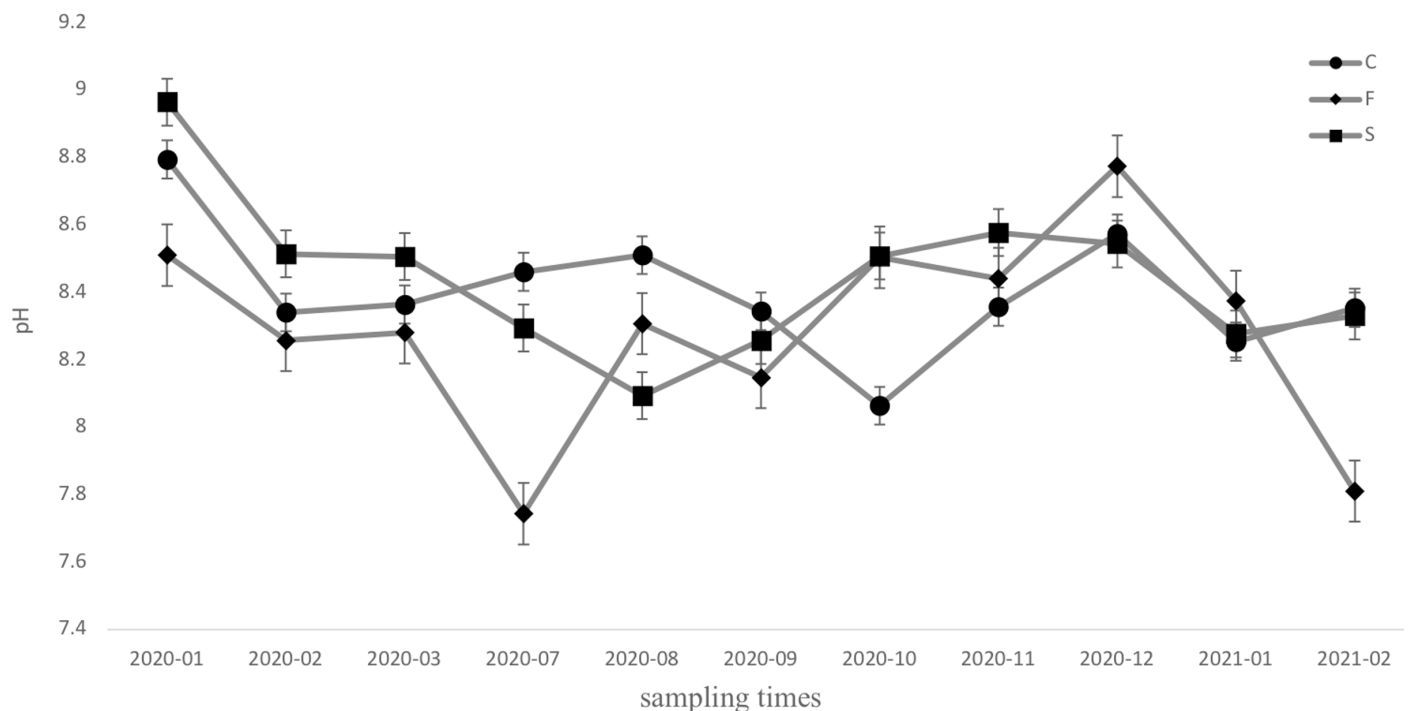


FIG. 8. Average pH for Cabo Rojo lagoons at each sampling time; C = Candelaria; F = Fraternidad; S = [Playa] Sucia.

local specimens and four included from the Rocha-Olivares et al. (2001) study. bPTP splits the data into 11 motus: local specimens into six and Rocha-Olivares et al. (2001) study into five. GMYC splits the data into eight or 11 motus, depending on the model and rate of molecular evolution used to make the BEAST tree. The Yule model with constant and relaxed rates of molecular evolution group local taxa into six putative species and Rocha-Olivares et al. (2001) into five groups. The relaxed clock model splits local taxa and one of the Rocha-Olivares et al. (2001) sequences into five groups, and four of the Rocha-Olivares et al. (2001) samples into three groups, this is the only model that groups one of the Rocha-Olivares et al. (2001) sequences with local *Cletocamptus*.

Species delimitation analysis yields three lineages of *Cletocamptus* in the CRLs, and at least five lineages of *Cletocamptus* in all sampled coastal lagoons. As in other studies, different methods of genetic species delimitation produce different results (Hofmann et al. 2019; Guimarães et al. 2022; Ranasinghe et al. 2022). This is due to model assumption violations and/or to the varying statistical power of each model (Carstens et al. 2013).

Coastal saline lagoons in Puerto Rico harbor several genetic lineages of *Cletocamptus*; some of these are likely undescribed species. The presence of mul-

multiple putative species is congruent with recent studies of *Cletocamptus* (Gómez and Yáñez-Rivera 2022) and Harpacticoida in general (Karanovic and Cooper 2012; Rossel and Martínez Arbizu 2019; Vakati et al. 2019) where careful examination of material yields a higher level of standing diversity than previously thought.

#### CONCLUSION

Harpacticoid copepods play a key role in aquatic trophic dynamics, both as primary and secondary consumers, and as a significant food source for many larger invertebrates and vertebrates (Williamson and Reid 2009; Baguley et al. 2019). *Cletocamptus* copepods are highly abundant in the Cabo Rojo lagoons of Puerto Rico, reaching up to 1,300 copepods/cm<sup>2</sup>. Given the high abundances these copepods reach, they should be considered essential components of the trophic dynamics and carbon fluxes of these coastal lagoons, an important stopover for migratory shorebirds in the Caribbean (Miller and Lugo 2009). No significant correlation between their abundances and factors: lagoon, area, or sampling time was found, meaning that the variability of the abundances at each lagoon area was independent throughout sampling times. The physicochemical parameters considered in this study did not correlate with abundances either. As such, copepods were observed to be tolerant to wide fluctuations in salinity, i.e., 27

TABLE 4. Mean genetic distances between lineages of *Cletocamptus* found in Puerto Rico, including the Rocha-Olivares et al. (2001) sequences (*C. stimpsoni*, *C. fourchensis*, *C. deborahdexterae*, *C. sinaloensis*, and *C. helobius*). Numbers 1–5 denote groups yielded by ML phylogenetic analysis of COI gene. The lower matrix triangle shows the corrected distances, and the upper one shows the uncorrected *p*-distance.

	PR-1	PR-2	PR-3	PR-4	PR-5	<i>C. sinaloensis</i>	<i>C. deborah-dexterae</i>	<i>C. fourchensis</i>	<i>C. stimpsoni</i>	<i>C. helobius</i>	<i>Stenocaris</i> sp.
PR-1		0.26	0.26	0.25	0.26	0.27	0.26	0.27	0.27	0.23	0.27
PR-2	0.42		0.21	0.2	0.19	0.22	0.21	0.19	0.21	0.21	0.22
PR-3	0.43	0.3		0.22	0.2	0.22	0.22	0.22	0.24	0.25	0.25
PR-4	0.38	0.28	0.33		0.11	0.19	0.18	0.21	0.24	0.26	0.25
PR-5	0.42	0.27	0.29	0.13		0.21	0.19	0.23	0.24	0.27	0.25
<i>C. sinaloensis</i>	0.43	0.32	0.33	0.26	0.3		0.09	0.22	0.26	0.27	0.26
<i>C. deborahdexterae</i>	0.43	0.31	0.34	0.25	0.27	0.10		0.22	0.25	0.27	0.27
<i>C. fourchensis</i>	0.46	0.27	0.32	0.31	0.34	0.33	0.33		0.23	0.24	0.24
<i>C. stimpsoni</i>	0.44	0.3	0.38	0.38	0.39	0.43	0.39	0.35		0.25	0.26
<i>C. helobius</i>	0.35	0.3	0.39	0.42	0.42	0.43	0.44	0.36	0.38		0.23
<i>Stenocaris</i> sp.	0.44	0.32	0.4	0.39	0.38	0.42	0.46	0.37	0.43	0.34	

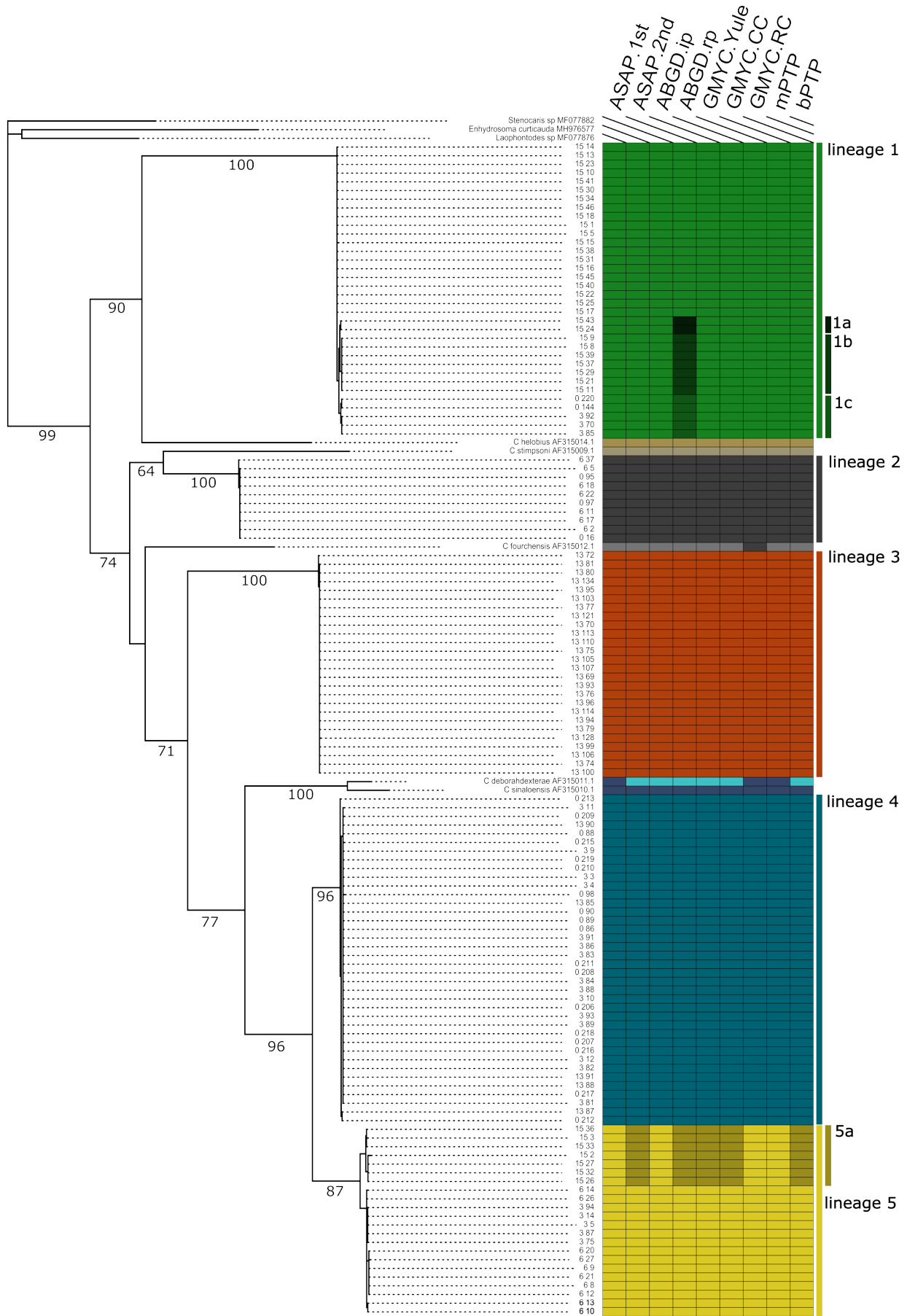
TABLE 5. Presence of each *Cletocamptus* lineage at each location in Puerto Rico as defined by the ML multi-gene phylogeny.

Lineage	Cabo Rojo	Providencia	Encarnación	Punta Petrona	Flamenco
1	✓	✓			✓
2	✓		✓		
3				✓	
4	✓	✓	✓	✓	✓
5		✓	✓		✓

to 135 psu, and can thrive in euhaline and hypersaline conditions.

Large genetic distances between the lineages, clear phylogenetic groupings, and consistent groupings yielded by species delimitation software indicate that coastal saline lagoons in Puerto Rico hold at least four genetic lineages/species of *Cletocamptus*. Some of these lineages exhibit patterns of genetic differentiation

attributed to spatial barriers, i.e., isolation of lagoons. Such diversity is an unexpected result, as we originally started this project to study *Cletocamptus dominicanus*, the only reported species at that time, with the initial hypothesis of no geographic differentiation due to the small sampling area and widespread distribution of these copepods. However, our finding of multiple putative species is congruent with recent studies of *Cleto-*



[Previous page] FIG. 9. Maximum likelihood phylogenetic tree constructed using COI sequences, annotated with species delimitation results. Colors represent putative lineages identified by the 1<sup>st</sup> and 2<sup>nd</sup> best ASAP results; initial and recursive partitions of ABGD; Yule, Constant Coalescent, and Relaxed Clock models for GMYC; mPTP; and bPTP. Numbers below branches indicate bootstrap values, while numbers at node tips correspond to individual copepod specimens. The tree was created with IQ-TREE (Minh et al. 2020), and annotations were made using the R v. 4.2.2 2 (RStudio Team 2020; R Core Team 2021) package of *Phytools* v. 1.5.3 (Revell 2012).

*camptus* and of Harpacticoida in general, where careful examination of material yields high levels of genetic divergence.

In summary, this study contributes to the ecology of the genus *Cletocamptus* in Puerto Rico by demonstrating that these remarkable harpacticoid copepods thrive in extreme saline environments. These environments could serve as natural laboratories for studying mechanisms of adaptation to extreme environments, particularly in terms of salinity and temperature. Additionally, this research contributes to the study of biodiversity and systematics within the genus *Cletocamptus* by developing genetic resources for species that may lack formal morphological descriptions. Given the importance, ephemeral nature, and vulnerability of these coastal lagoons, it is crucial to study and protect their biota and habitat.

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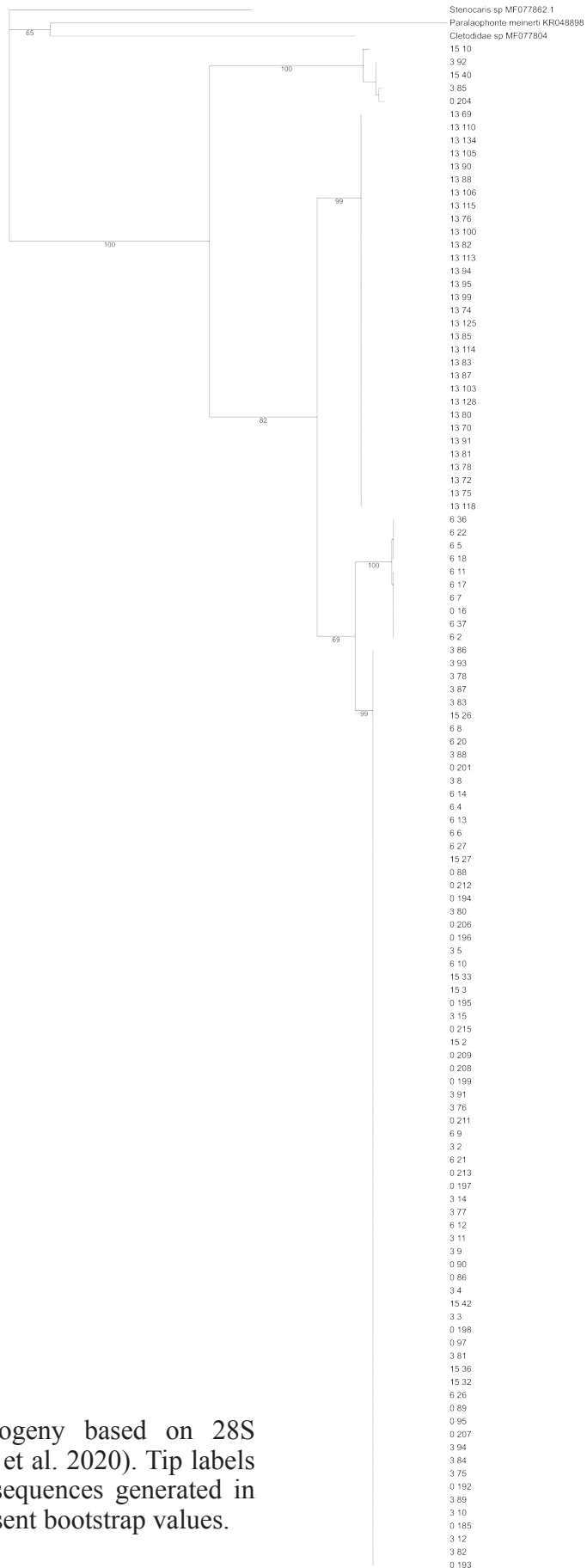


FIG. 10. Maximum likelihood phylogeny based on 28S sequences, created using IQ-TREE (Minh et al. 2020). Tip labels represent sequences from *Cletocamptus* sequences generated in this study, numbers next to branches represent bootstrap values.



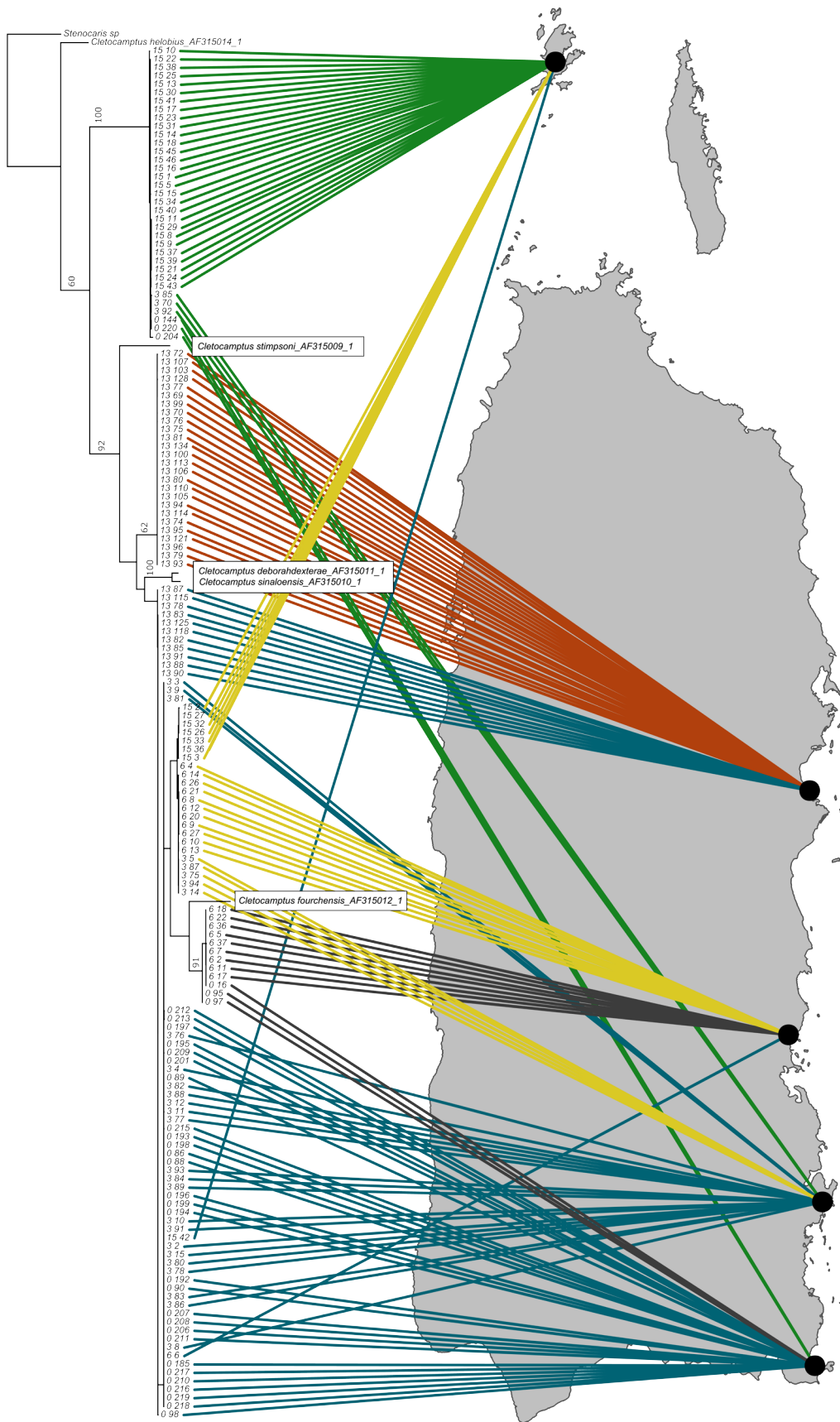


FIG. 11. Multigene ML phylogeny overlaid on a map of Puerto Rico. The phylogenetic tree was constructed using IQ-TREE (Minh et al. 2020) with concatenated COI and 28S sequences, including those from Rocha-Olivares et al. (2001). *Stenocaris* sp. was used as an outgroup. Each tip represents a single copepod specimen, with colored lines linking specimens to their sampling locations. Green, gray, red, blue, and yellow represent lineages 1 through 5, respectively; phylogenetic groups were rearranged for clarity on the map. Numbers next to branches represent bootstrap values. The figure was created using the R v. 4.2.2.2 (RStudio Team 2020; R Core Team 2021) package of Phytools v. 1.5.3 (Revell 2012).

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APPENDIX 1a. COI GenBank accession numbers and associated collection data; all specimens are from Puerto Rico; JPD = J. P. Domenech, AG = A. González, LV = Liz Vega, NS = N. Schizas.

Voucher #	Coordinates	Sex	Collected by	Date	Locality	Accession	GI
0_144	17.956305, -67.211543	female	JPD, AG	08-DEC-2019	Cabo Rojo: Laguna Candelaria	OQ682378.1	2463908584
0_16	17.944309, -67.192303	female	JPD, AG	08-DEC-2019	Cabo Rojo: Laguna Sucia	OQ682339.1	2463908468
0_206	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682315.1	2463908405
0_207	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682316.1	2463908408
0_208	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682317.1	2463908410
0_209	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682336.1	2463908460
0_210	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682318.1	2463908413
0_211	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682319.1	2463908415
0_212	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682301.1	2463908370
0_213	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682303.1	2463908375
0_215	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682337.1	2463908463
0_216	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682320.1	2463908417
0_217	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682321.1	2463908420
0_218	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ682322.1	2463908422
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13_105	17.96006, -66.36575	female	JPD, LV	02-APR-2021	Santa Isabel: Laguna Punta Petrona	OQ682351.1	2463908508
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13_107	17.96006, -66.36575	female	JPD, LV	02-APR-2021	Santa Isabel: Laguna Punta Petrona	OQ682353.1	2463908513
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6_14	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682287.1	2463908334
6_17	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682342.1	2463908476
6_18	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682345.1	2463908483
6_2	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682340.1	2463908471
6_20	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682289.1	2463908339
6_21	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682290.1	2463908342
6_22	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682346.1	2463908494
6_26	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682281.1	2463908319
6_27	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682293.1	2463908350

6_37	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682348.1	2463908500
6_5	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682341.1	2463908473
6_8	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682291.1	2463908345
6_9	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ682292.1	2463908347

APPENDIX 1b. 28S GenBank accession numbers and associated collection data; all specimens are from Puerto Rico; JPD = J. P. Domenech, AG = A. González, LV = Liz Vega, NS = N. Schizas.

Voucher #	Coordinates	Sex	Collected by	Date	Locality	Accession	GI
0_16	17.944309, -67.192303	female	JPD, AG	08-DEC-2019	Cabo Rojo: Laguna Fraternidad	OQ819183.1	2486667153
0_185	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819193.1	2486667163
0_192	17.947031, -67.192559	male	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819194.1	2486667164
0_193	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819195.1	2486667165
0_194	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819196.1	2486667166
0_195	17.947031, -67.192559	male	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819197.1	2486667167
0_196	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819198.1	2486667168
0_197	17.947031, -67.192559	male	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819199.1	2486667169
0_198	17.947031, -67.192559	male	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819200.1	2486667170
0_199	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819201.1	2486667171
0_201	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819202.1	2486667172
0_204	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819295.1	2486667265
0_206	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819203.1	2486667173
0_207	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819204.1	2486667174
0_208	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819205.1	2486667175
0_209	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819206.1	2486667176
0_211	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819207.1	2486667177
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0_89	17.947031, -67.192559		JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819213.1	2486667183
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0_95	17.947031, -67.192559	female	JPD	14-SEP-2018	Cabo Rojo: Laguna Fraternidad	OQ819215.1	2486667185
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13_82	17.96006, -66.36575		JPD, LV	02-APR-2021	Santa Isabel: Laguna Punta Petrona	OQ819285.1	2486667255
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13_99	17.96006, -66.36575	female	JPD, LV	02-APR-2021	Santa Isabel: Laguna Punta Petrona	OQ819294.1	2486667264
15_10	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819299.1	2486667269
15_2	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819219.1	2486667189
15_26	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819217.1	2486667187
15_27	18.32665, -65.31479		JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819218.1	2486667188
15_3	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819223.1	2486667193
15_32	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819220.1	2486667190
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15_36	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819222.1	2486667192
15_40	18.32665, -65.31479	female	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819296.1	2486667266
15_42	18.32665, -65.31479	male	JPD, NS	17-AUG-2022	Culebra: Laguna Flamenco	OQ819224.1	2486667194
3_10	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819225.1	2486667195
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3_12	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819226.1	2486667196
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3_5	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819231.1	2486667201
3_75	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819232.1	2486667202
3_76	17.9417, -66.95926		JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819233.1	2486667203
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3_78	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819235.1	2486667205
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3_80	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819236.1	2486667206
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3_83	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819239.1	2486667209
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3_87	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819242.1	2486667212
3_88	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819243.1	2486667213
3_89	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819244.1	2486667214
3_9	17.9417, -66.95926	female	JPD, LV	02-APR-2021	Guánica: Laguna Providencia	OQ819249.1	2486667219
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6_10	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819250.1	2486667220
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6_12	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819251.1	2486667221
6_13	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819252.1	2486667222
6_14	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819253.1	2486667223
6_17	17.99063, -66.7182		JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819185.1	2486667155
6_18	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819189.1	2486667159
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6_37	17.99063, -66.7182		JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819187.1	2486667157
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6_5	17.99063, -66.7182	male	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819192.1	2486667162
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6_8	17.99063, -66.7182	female	JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819260.1	2486667230
6_9	17.99063, -66.7182		JPD, LV	02-APR-2021	Peñuelas: Laguna Encarnación	OQ819261.1	2486667231

APPENDIX 2. Average salinity (psu), pH, and temperature for each lagoon at each sampling time. The number in parentheses is the standard deviation.

Date	Lagoon	Salinity (psu)	pH	Temperature (° C)
2020-01-23	Candelaria	58 (12)	8.79 (0.22)	32 (0)
	Fraternidad	76 (3)	8.51 (0.05)	32 (1)
	Playa Sucia	60 (0)	8.96 (0.01)	33 (2)
2020-02-21	Candelaria	82 (43)	8.34 (0.12)	32 (3)
	Fraternidad	100 (11)	8.26 (0.02)	25 (2)
	Playa Sucia	79 (1)	8.51 (0.11)	27 (2)
2020-03-16	Candelaria	86 (6)	8.36 (0.11)	33 (1)
	Fraternidad	105 (13)	8.28 (0.07)	28 (5)
	Playa Sucia	87 (2)	8.51 (0.11)	33 (1)
2020-07-27	Candelaria	97 (19)	8.46 (0.18)	40 (2)
	Fraternidad	101 (55)	7.74 (0.38)	36 (3)

	Playa Sucia	90 (5)	8.29 (0.05)	38 (0)
2020-08-19	Candelaria	40 (8)	8.51 (0.15)	34 (2)
	Fraternidad	55 (11)	8.31 (0.12)	32 (1)
	Playa Sucia	55 (8)	8.09 (0.18)	31 (1)
2020-09-09	Candelaria	60 (17)	8.34 (0.15)	35 (2)
	Fraternidad	72 (17)	8.15 (0.15)	31 (1)
	Playa Sucia	58 (3)	8.26 (0.11)	31 (2)
2020-10-21	Candelaria	52 (16)	8.06 (0.04)	34 (3)
	Fraternidad	64 (2)	8.5 (0.08)	33 (3)
	Playa Sucia	59 (2)	8.51 (0.04)	37 (4)
2020-11-11	Candelaria	31 (4)	8.36 (0.17)	28 (1)
	Fraternidad	44 (6)	8.44 (0.02)	26 (1)
	Playa Sucia	41 (6)	8.58 (0.35)	26 (1)
2020-12-23	Candelaria	42 (7)	8.57 (0.63)	31 (3)
	Fraternidad	49 (12)	8.77 (0.16)	32 (2)
	Playa Sucia	46 (1)	8.54 (0.03)	33 (1)
2021-01-26	Candelaria	53 (18)	8.25 (0.31)	27 (3)
	Fraternidad	51 (14)	8.37 (0.29)	29 (1)
	Playa Sucia	53 (2)	8.28 (0.03)	31 (1)
2021-02-26	Candelaria	79 (41)	8.35 (0.23)	32 (3)
	Fraternidad	81 (38)	7.81 (0.53)	32 (3)
	Playa Sucia	81 (3)	8.33 (0.02)	32 (3)

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