INTRODUCTION

The amount of gene flow between populations is considered to be relatively proportional to the dispersal ability and geographic range of the organisms involved (Bohonak 1999, Lester & Ruttenberg 2005). For sessile marine invertebrates (fixed and those with a limited home range), the only effective means of dispersal is a planktonic larval stage. Thus, benthic species (those that do not undergo large migrations) are distributed in patches that can only be connected by planktonic larvae. Species that have long larval periods were thought to have extended geographic distributions and associated high levels of gene flow when compared to species

ABSTRACT: We studied the molecular population structure of the hermaphroditic caridean shrimp Lysmata wurdemanni (Gibbes, 1850), which is distributed along the east coast of the USA and in the Gulf of Mexico. Four populations within the reported range were sampled: North Carolina (NC), Florida east coast (FLE), Florida Keys (FLK), and Texas (TX). Of the 3 populations of L. wurdemanni tested for interpopulation breeding (FLE, FLK, TX), only the FLE × FLK cross was successful. Both Florida populations failed to successfully hybridize with the Texas population during the experimental period. Additionally, there appears to be strong directionality between Florida and Texas crosses: when TX females crossed with either male FLK or FLE shrimps, few eggs were present and no larval hatching occurred, whereas crosses between FLE or FLK females and TX males produced embryos, but no viable larvae were produced during the experimental period. Genealogical analyses of the mitochondrial genes cytochrome oxidase subunit I (COI) and the large subunit of ribosomal RNA (16S rRNA) genes yielded distinct and highly supported clades for each of the FLE, FLK, and TX populations. High pairwise $F_{ST}$ values between TX versus FLK, TX versus FLE, and FLK versus FLE suggested that the populations are genetically disconnected. The NC and FLE populations could not be distinguished phylogenetically. This is an example of a marine species for which the potential for larval dispersal is predominantly unrealized given the observed absence of gene flow and the strong reproductive isolation between populations on the southeast coast of the United States and on the Gulf coast.

KEY WORDS: Lysmata wurdemanni · Population connectivity · Reproductive compatibility · Interbreeding · Caridean shrimp

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with shorter larval durations (Shuto 1974, Jablonksi 1986). Those marine species with extended planktotrophic life cycles have been thought of as open populations, with regular or intermittent occurrences of gene flow from surrounding populations.

The amount of time larvae spend in the water column can vary dramatically within or across families and even within the same genus. This has provided contrasting examples of larval dispersion and biogeography within very closely related species (Kinlan & Gaines 2003, Bird et al. 2007, Kelly & Eernisse 2007). These studies have provided growing evidence that discounts the notion that long-lived planktonic larvae can be equated with broad dispersion and the absence of genetic structure across their geographic range. Populations that appear similar may in fact be distinct isolated populations of the same species or complexes of cryptic species (Knowlton 1993, Palumbi 1994). For example, Lee & Foighil (2004, 2005) examined the genetic structure of the scoured mussel Brachidontes exustus and established that ‘populations’ of this species were in fact a complex of species throughout its historical geographic range. Lee (2000) examined the global phylogeography of the copepod Eurytemora affinis and demonstrated with mitochondrial DNA and interpopulation mating experiments that the presumed uniformly distributed species was in fact several reproductively isolated sibling species. After detailed morphological examination, genetic lineages of the copepod Cletocamptus deitersi (see Rocha-Olivares et al. 2001) also proved to be different species (Gómez et al. 2004). This was illustrated in the case of the whelk Busycicon perversum as well, which is associated with high divergence between the Florida Atlantic and Gulf of Mexico populations. These populations are now considered sister taxa (Wise et al. 2004). Genetic divergence between marine populations may evolve as a result of intrinsic factors, extrinsic factors, or their interaction (Hilbish 1996). Divergence between populations may be maintained in the presence of selection. Sotka et al. (2004) suggested that, while populations of the barnacle Balanus glandula should be connected by dispersal alone (over a geological time scale), there is a strong likelihood that selection influences the patterns of genetic variation in populations and causes the sharp differences between northern and southern populations. Ocean currents and physical processes may indeed act as physical barriers promoting speciation. Selection for a specific genotype would further magnify this effect (Hilbish & Koehn 1985, Schizas et al. 2001). These genetic patterns have been mirrored in terrestrial environments, where physical barriers lead to localized extinction of haplotypes in widely distributed allopatric species, with differences only detectable at the genetic level (Vogler & DeSalle 1994, Barber 1999).

Among decapod crustaceans, several studies have focused on the genealogy of populations and the phylogeny of closely related taxa. The best known are those dealing with transisthmian pairs of snapping shrimp (Alpheus). Knowlton et al. (1993) reported divergence in the mtDNA cytochrome oxidase subunit I (COI) gene among 7 pairs of Alpheus species across the Isthmus of Panama to be between 6.6 and 19.2%, with reproductive isolation, but not behavioral isolation, occurring in all but 1 replicate of the transisthmian crosses. In contrast, crosses between pairs of Alpheus species from the same ocean produced fertile clutches 60% of the time. It was inferred that reproductive isolation requires >6.6% divergence in COI sequences, or 4.5 to 5 million years of isolation, if the revised rates for divergence (1.4% per 10^6 years) across the Isthmus of Panama are used (Knowlton & Weigt 1998). Mathews et al. (2002) found divergence of 2 to 5% between 2 closely related species of Alpheus shrimps. Later, Mathews (2006) examined 16S sequences of the Alpheus shrimp over a wide geographic area and established distinct isolated populations and/or cryptic species.

While the separation of 2 bodies of water by a land mass is an obvious reproductive barrier for marine species, oceanographic features may also pose severe restrictions on a species’ ability to disperse. In addition to physical dispersal barriers, a few studies have experimentally demonstrated an unexpectedly high amount of local recruitment among species with extended larval development. This has been reported in Alpheus (see Knowlton & Keller 1986) and in some reef fish (Jones et al. 1999, Taylor & Hellberg 2003). Both biotic and abiotic factors can influence localized recruitment (Almany et al. 2007), e.g. larval behavior (Burton & Feldman 1982), temperature gradients (Wares et al. 2001) and ocean currents (McConaughha 1992, Baums et al. 2006) have been implicated in the restriction of larval dispersal.

As our study species we chose Lysmata spp.; these are small caridean shrimps that occupy wide geographic ranges and diverse ecological habitats (Bauer 2004). They differ from nearly all other decapods by having an unusual reproductive mode, protandric simultaneous hermaphroditism. Lysmata shrimp first develop as males and then may change to become a simultaneous hermaphrodite (Bauer & Holt 1998, Fiedler 1998, Bauer 2000). Within the genus, species delineations can be made on the basis of adult morphology, larval developmental characteristics, and ecological associations. Larvae within the Lysmata genus have been studied relatively well in recent years (e.g. Calado et al. 2005), and development times have been reported to range from approximately 18 to >160 d in the laboratory. Like many other caridean shrimp,
Lysmata may undergo a developmental delay (mark-time molting) (Gore 1985, Anger 2001), if conditions are not appropriate for an extended period of time, development (pre-competency) and settlement (post-competency) can be delayed for well over 30 d (A. Rhyne pers. obs.).

Until recently, the western Atlantic peppermint shrimp Lysmata wurdemannii was ubiquitous throughout its known range (New Jersey, USA, to Brazil). The species was reported to occupy multiple habitat types from rocky shorelines, hard-bottom reefs, tube sponges, to shell bottoms within inlets. However, a recent taxonomic revision (Rhyne & Lin 2006) identified 4 new species of Lysmata and clarified the distributional ranges for 6 species of western Atlantic Lysmata. Of these, L. wurdemanni was restricted to the Gulf of Mexico and the east coast of United States, with ranges extending from South Padre Island, Texas, to Long Island Sound, New York (Rhyne & Lin 2006). The revision utilized live color patterns, morphology, and interbreeding experiments to differentiate between species. No interbreeding trials were conducted between populations of the 6 species of western Atlantic Lysmata.

The goal of the present study was to decipher the genetic population structure of Lysmata wurdemanni across its range to determine if the species is genetically homogeneous or consists of isolated, distinct populations. To achieve this goal we chose several populations of the 6 species of western Atlantic Lysmata.

The goal of the present study was to decipher the genetic population structure of Lysmata wurdemanni across its range to determine if the species is genetically homogeneous or consists of isolated, distinct populations. To achieve this goal we chose several populations of L. wurdemanni along its reported geographic range and compared both interbreeding results and the molecular population structure derived from the mitochondrial genes COI and 16S.

MATERIALS AND METHODS

Sample collection and preparation. Samples from 4 localities (Fig. 1) were collected with hand-nets along shallow-water rocky jetties during a nocturnal low tide at Sebastian Inlet (FLE) and Port Aransas Pass, Port Aransas, Texas (TX), with SCUBA in shallow (2 to 3 m) waters at Banks Channel, Wilmington, North Carolina (NC), and in rubble or with stone crab traps at 10 to 15 m depth, 35 to 50 km NNW of Marathon, Florida in Florida Bay (FLK). Hereafter, the terms samples and populations will be used interchangeably. The sampling method was determined by habitat availability and previous knowledge of resident populations. With the exception of the NC sample (2 preserved specimens), all other samples were transported to the Vero Beach Marine Laboratory alive and housed in 80 l aquaria for observation. Following 4 d of observation, the shrimp were given catalog numbers and moved into experimental tanks (20 l) for breeding experiments. Shrimps were fed 2 to 4 times daily an excess of a mixed diet of frozen mysid shrimp and adult brine shrimp. Entire shrimp or tissue clippings of pleopods were preserved in 95% ethanol 2 d prior to extraction of DNA, and fixed specimens were transported to the Isla Magueyes Laboratories (University of Puerto Rico, Mayagüez) for DNA processing.

Interbreeding trials. To determine reproductive compatibility, interbreeding experiments were conducted between all combinations of collected populations, except those from North Carolina (Table 1). For the remainder of the paper, reference to sex refers to the acting role of the shrimp during mating. Bauer & Holt (1998) have thoroughly reviewed the mating system of Lysmata wurdemanni. They elegantly demonstrated that individuals of L. wurdemanni are incapable of both self-fertilization and storing sperm. Females are available for mating immediately after molting and are not capable of making a mate choice because mating takes place before their soft body hardens (Bauer & Holt 1998). Males mate prior to molting; they react to the release of a sex pheromone and mate readily with freshly molted females. L. wurdemanni requires no acclimation time to potential mates, as they generally live in groups in the wild and are opportunistic in mating.
for 16S) annealing, finishing with 5 min incubation at 45°C (42°C for 16S) for 30 s, extension at 72°C for 1 min. We programmed an anchor-PCR procedure with 10 cycles of denaturation at 95°C for 15 s, annealing at 50°C for 30 s, and extension at 60°C for 45 s, repeated 60 times. Cycle sequence reactions were ethanol precipitated and loaded in either a ABI3130xl or an ABI 377 sequencing machine. DNA sequencing trace files were processed with the Phrap/Phred/Consed programs (Ewing & Green 1998, Ewing et al. 1998, Gordon 2004) for base calling, quality assessment, and, if necessary, manual editing. Edited DNA sequences were imported into MacClade (Maddison & Maddison 1992) for manual alignment and quality control. Aligned sequences were imported into DnaSP 4.10.9 (Rozas et al. 2003) for general statistical analysis, such as Tajima’s D (Tajima 1989) and Fu’s Fs (Fu 1997) to test for deviation from the neutral model of molecular evolution (Kimura 1968). For population structure analysis, $F_{ST}$ was calculated based on the average number of pairwise nucleotide differences within and between populations in Arlequin 3.11 (Excoffier et al. 2005). The appropriate model of nucleotide substitution for genealogical reconstruction using neighbor-joining, maximum likelihood and $F_{ST}$ calculations was estimated by a hierarchical likelihood ratio test (Huelsenbeck & Crandall 1997) in ModelTest 3.7 (Posada & Crandall 1998). If the DNA analysis program (i.e. Arlequin 3.11) did not provide the exact model of substitution, then, the more complex, most related model was used instead. The justification for this decision is based on the nested hierarchy of DNA substitution models, where simple models are nested within more complex models. Genealogies of the sequences were reconstructed using the neighbor-joining (NJ), maximum parsimony (MP), and maximum likelihood (ML) algorithms in PAUP* (Swofford 2002). Heuristic searches in MP reconstruction were performed using 10 random addition replicates with nearest neighbor interchange (NNI) branch swapping. Heuristic searches in ML reconstruction were performed as follows: starting tree(s) obtained via stepwise addition, random addition of sequences, 10 replicates, and tree bisection-reconnection (TBR) as the branch-swapping algorithm. Robustness of clades was evaluated with 1000 (MP and NJ) and 100 bootstrap replications (ML) (Felsenstein 1985). *Low level of F1 survival obtained; see ‘Discussion’. Note: L. wurdemanni are euhermaphrodites; the terms male and female are used to indicate the direction of the crosses.

The experimental layout for the interbreeding experiments consisted of 3 intra-population controls and 6 inter-population experimental crosses for each pair of populations (Table 1). Since Lysmata wurdemanni are euhermaphrodites, both male and female crosses were obtained from each pairing. Each shrimp was allowed to adjust to experimental conditions for at least a molt cycle, and the tanks were checked daily for the presence of molts. Because collection of eggs can be stressful on the shrimp, when egg masses were visually observed, a small sample was removed (approximately 50 eggs) to determine fertility. Two mortalities were obtained and those individuals were replaced with shrimp from the same population in similar breeding conditions. The replacement shrimp were maintained in identical conditions as those in the breeding experiments. When a shrimp molted, it was visually checked for the presence of eggs on the abdomen. The night of larval release, the shrimp was moved to a larval rearing system (Calado et al. 2003). The following data were collected using the general protocols by Fiedler (1998): (1) embryo development after 24 h, (2) retention of embryos beyond Day 5, (3) development of embryos beyond Day 10, and (4) successful larval hatching.

**Molecular analysis.** DNA was isolated with the PureGene DNA isolation kit (Gentra) for fixed tissue according the manufacturer’s guidelines. A portion of the mitochondrial COI and 16S genes were amplified with an Eppendorf MasterCycler. The COI region was amplified with the universal primers LCO1490 and HCO2198 (Folmer et al. 1994), and the 16S region was amplified with the 1471 to 1472 primers (Crandall & Fitzpatrick 1996). After the initial denaturation of 3 min at 95°C, we programmed an anchor-PCR procedure with 10 cycles of denaturation at 95°C for 15 s, annealing at 45°C (42°C for 16S) for 30 s, extension at 72°C for 60 s (45 s for 16S), and 25 more cycles at 50°C (45°C for 16S) annealing, finishing with 5 min incubation at 72°C. Successful PCR reactions were verified by running 5 µl of the amplicon on a 1% Tris-Borate-EDTA agarose gel. Amplified bands were visualized under UV light and captured digitally. PCR reactions were cleaned of excess dNTPs, primers, and other impurities by the enzymatic treatment EXOSAP. For every 15 µl of PCR product, 3 µl of the EXOSAP solution were added and incubated at 37°C for 15 min and at 80°C for 15 min. Sequencing reactions with each of the primers were prepared with the 3.1 BigDye Termination Kit under the following conditions: 94°C for 5 s, annealing at 50°C for 5 s, and extension at 60°C for 85 s, repeated 60 times. Cycle sequence reactions were ethanol precipitated and loaded in either a ABI3130xl or an ABI 377 sequencing machine. DNA sequencing trace files were processed with the Phrap/Phred/Consed programs (Ewing & Green 1998, Ewing et al. 1998, Gordon 2004) for base calling, quality assessment, and, if necessary, manual editing. Edited DNA sequences were imported into MacClade (Maddison & Maddison 1992) for manual alignment and quality control. Aligned sequences were imported into DnaSP 4.10.9 (Rozas et al. 2003) for general statistical analysis, such as Tajima’s D (Tajima 1989) and Fu’s Fs (Fu 1997) to test for deviation from the neutral model of molecular evolution (Kimura 1968). For population structure analysis, $F_{ST}$ was calculated based on the average number of pairwise nucleotide differences within and between populations in Arlequin 3.11 (Excoffier et al. 2005). The appropriate model of nucleotide substitution for genealogical reconstruction using neighbor-joining, maximum likelihood and $F_{ST}$ calculations was estimated by a hierarchical likelihood ratio test (Huelsenbeck & Crandall 1997) in ModelTest 3.7 (Posada & Crandall 1998). If the DNA analysis program (i.e. Arlequin 3.11) did not provide the exact model of substitution, then, the more complex, most related model was used instead. The justification for this decision is based on the nested hierarchy of DNA substitution models, where simple models are nested within more complex models. Genealogies of the sequences were reconstructed using the neighbor-joining (NJ), maximum parsimony (MP), and maximum likelihood (ML) algorithms in PAUP* (Swofford 2002). Heuristic searches in MP reconstruction were performed using 10 random addition replicates with nearest neighbor interchange (NNI) branch swapping. Heuristic searches in ML reconstruction were performed as follows: starting tree(s) obtained via stepwise addition, random addition of sequences, 10 replicates, and tree bisection-reconnection (TBR) as the branch-swapping algorithm. Robustness of clades was evaluated with 1000 (MP and NJ) and 100 bootstrap replications (ML) (Felsenstein 1985). Lysmata pederseni was used as the outgroup in both gene trees.

<table>
<thead>
<tr>
<th>Population</th>
<th>FLE (male)</th>
<th>FLK (male)</th>
<th>TX (male)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLE (female)</td>
<td>1–4</td>
<td>1–4</td>
<td>1–3*</td>
</tr>
<tr>
<td>FLK (female)</td>
<td>1–4</td>
<td>1–4</td>
<td>1–3*</td>
</tr>
<tr>
<td>TX (female)</td>
<td>1, 2</td>
<td>1, 2</td>
<td>1–4</td>
</tr>
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</table>
RESULTS

Interbreeding trials

Of the 3 populations tested during interpopulation breeding trials, 2 crosses failed to produce embryos that survived beyond Day 5. These 2 failed crosses occurred between the TX female shrimps and FLK or FLE male shrimps (Table 1). In these crosses, egg number was visibly reduced from normal fertilization (only a few hundred eggs as opposed to normal mating yields in which there is a full abdomen of eggs, often bulging out beyond the pleopods). Additionally, crosses between FLE or FLK females and TX males produced embryos that survived beyond Day 10; however, they were not subsequently viable as no larvae were produced. These embryos were characterized by having a higher than normal rate of mortality. While the embryos were not removed and counted, a noticeable number of embryos in these crosses appeared to be bacterially infected (indicated by a dark brown color). All other crosses produced viable larvae (Table 1).

Molecular analysis

A 540 bp portion of COI and a 497 bp portion of 16S were used for analysis after editing and trimming the DNA sequences. The resulting 16S segment of Lysmata pederseni was 4 bases longer than the 16S segment of L. wurdemanni. No insertions or deletions of amino acids in COI were found within the 4 populations of L. wurdemanni. There were no stop codons present in the amino acid translation, and no sequences contained double bands in both the forward and reverse reactions. Additionally, BLAST searches indicated that the closest hits were previously published sequences from related species of shrimps. Altogether, these results provide a high level of confidence that the sequences are mtDNA and not pseudogenes. The molecular data will be presented gene-by-gene, as a few shrimps did not yield a DNA sequence because of PCR and sequencing failures.

Sequence data from COI and 16S regions were obtained for shrimps from 4 localities/populations resulting in 17 (COI; Table 2) and 8 distinct haplotypes (16S; Table 3), respectively. Shrimps from the 4 sampling locations shared no haplotypes (except FLE and NC in 16S; Tables 2 & 3), but the nucleotide diversity within locations was low. Nucleotide diversity levels of COI/16S were estimated for NC (π = 0/0), FLE (π = 0.00446/0.00073), FLK (π = 0.00086/0.00106), and TX (π = 0.00365/0).

Tajima’s D and Fu’s Fs statistics did not reveal genetic patterns inconsistent with DNA neutrality for COI or 16S, indicating that the chosen genes are evolving neutrally and are therefore appropriate for population level inferences for Lysmata wurdemanni (data not shown). Nucleotide substitutions in COI within and among populations were all synonymous, occurring in the third and first codon positions. These substitutions consisted of 62 transitions (5 singletons) and 13 transversions. Three non-synonymous substitutions were recorded in L. pederseni, resulting in 3 amino acid differences compared to L. wurdemanni. There were 29 transitions and 16 transversions in the 16S sequences of L. wurdemanni. Genealogical trees were constructed in PAUP using HKY (Hasegawa-Kishino-Yano model of substitution) as the most appropriate model of substitution as the most appropriate model of substitution for both COI [Base = (0.2605, 0.2198, 0.1845), Nst = 2, TRatio = 5.6887, Γ = 0.1911], and 16S [Base = (0.3125,
Phylogenetic analysis of the COI and 16S data, in combination or separately, support a strong divergence between all populations, except NC and FLE, in which NC haplotypes were nested inside the FLE clade (Figs. 2 & 3). The extremely high \( F_{ST} \) statistics (Table 4), genealogical analysis (Figs. 2 & 3), and the COI parsimony network (Fig. 4) underline the distinct nature of the sampled populations of *L. wurdemanni*. These analyses demonstrated a lack of gene flow and strong isolation between all populations, except those on the east coast of the United States. There was a surprisingly large divergence (3.23% for COI and 2.55% for 16S, on average, 17.3 and 11.9 substitutions, respectively; Table 4) between the 2 populations in Florida, even though the locations are separated by only ~450 km and strong north-bound currents (Florida Current and Gulf Stream) exist within 50 km of the FLK and FLE populations. The highest divergence was observed in COI between the TX and the FLE and NC populations (>10%, 54 to 56 substitutions), followed by the...

### Table 4. *Lysmata wurdemanni*. Average number of nucleotide differences between populations, divergence, and pairwise \( F_{ST} \) statistics for the COI/16S genes sequenced from 4 different populations; Wilmington, North Carolina (NC); Sebastian Inlet, Florida (FLE); Florida Bay, Marathon, Florida (FLK); and Port Aransas, Texas (TX). Average numbers of nucleotide differences and percentages of divergence were calculated in DNAsp. \( F_{ST} \) calculations were carried out with Arlequin, using Tamura-Nei distance (Tamura & Nei 1993; TrN). The specific settings were TrN + \( \Gamma \) (0.2) + TRatio = 5 for COI and TrN + TRatio = 2 for 16S. na: no tests were performed since only 1 NC sequence was available.

<table>
<thead>
<tr>
<th>Population Pairs</th>
<th>Average Divergence (%)</th>
<th>( F_{ST} )</th>
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<tbody>
<tr>
<td>FLE × NC</td>
<td>2.83 (0.2)</td>
<td>0.336 (na)</td>
</tr>
<tr>
<td>FLE × FLK</td>
<td>17.3 (11.9)</td>
<td>0.936 (0.967)</td>
</tr>
<tr>
<td>FLE × TX</td>
<td>54.8 (33.1)</td>
<td>0.973 (0.994)</td>
</tr>
<tr>
<td>FLK × NC</td>
<td>16.9 (11.7)</td>
<td>0.976 (na)</td>
</tr>
<tr>
<td>FLK × TX</td>
<td>51.0 (38.0)</td>
<td>0.986 (0.995)</td>
</tr>
<tr>
<td>TX × NC</td>
<td>56.0 (33.0)</td>
<td>0.980 (na)</td>
</tr>
</tbody>
</table>

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**Fig. 2. *Lysmata wurdemanni.* The 50% majority rule genealogical tree constructed with maximum-likelihood based on cytochrome oxidase subunit I (COI) sequences. Bootstrap values are presented for maximum-likelihood/maximum-parsimony/neighbor-joining analyses. Bootstrap values within the major clades have been deleted for clarity.**

**Fig. 3. *Lysmata wurdemanni.* The 50% majority rule genealogical tree constructed with maximum likelihood based on 16S sequences. Bootstrap values are presented for ML/MP/NJ analyses. * Bootstrap values <50%**
Texas–Florida Keys divergence (9.5%, 51 substitutions). FLE populations are closely related to NC populations (about 0.5%, 3 substitutions); however, the populations are indistinguishable with the 16S data. This is not surprising since 16S is regarded as a more conservative gene than COI and is routinely used for higher level phylogenetic studies (e.g. Braga et al. 1999, Goetze 2003, Porter et al. 2005). DNA sequences have been deposited in GenBank under Accession Numbers EU135796 to EU135869.

**DISCUSSION**

*Lysmata wurdemanni* has an extended planktonic larval stage, with ≥21 d in the water column and up to 90 d under less than ideal conditions in the laboratory (A. L. Rhyne & D. Zhang unpubl. data). The mitochondrial sequences delineate a striking separation between populations at multiple points along the species’ geographic range. This strongly supports an unrealized larval dispersal potential in the species. Populations that are a few hundred kilometers apart were significantly differentiated. Gene flow was absent across the geographic range of the species, with the possible exception of populations from the east coast of the United States. Because the sample size was small from the North Carolina population, inferences on the connectivity between the Florida east coast and those to the north, presumably connected by the Gulf Stream, are preliminary at the present time. Given the limited number of available specimens, there were no shared haplotypes between the North Carolina and Florida populations for COI (Fig. 4). An extended survey of populations along the east coast, coupled with a larger number of samples, will most likely result in shared haplotypes between these populations. Since COI is usually more variable than 16S, it is not surprising that the single 16S haplotype from the NC population was identical to one of the haplotypes sequenced from FLE (Table 3). The divergence measured between the 2 populations on the east coast of the United States was extremely low (0.5% in COI, absent in 16S), when compared to that between other populations.

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**Fig. 4. Lysmata wurdemanni.** Parsimony haplotype network based on COI sequences. Each large circle represents a unique haplotype. The haplotype identification number and the number of specimens sharing each haplotype are indicated within each circle. The minimum number of steps is represented by the small empty circles between each haplotype. Closed loops represent homoplasy, as indicated by the statistical parsimony analysis.
The complete lack of gene flow between populations within Florida was surprising given the close proximity to strong east-then north-bound currents in the Florida Keys, and the long larval duration in the water column. One might have expected the populations to be connected in a stepping-stone fashion, with limited or no genetic connectivity between the northern and southern populations of Florida. However, the data strongly support a series of subdivided populations. Given the geographic distance between the populations, population differentiation was expected between Texas and all other populations surveyed, but not between the 2 Floridian locations. The high level of divergence (>10%), coupled with unsuccessful hybridization (Table 1) between Texas and Florida populations, verifies that comparison of COI genes provides an acceptable level of resolution to distinguish unique populations. At the same time populations with <3% divergence readily interbreed without any modality. Although no experimental data were available, it is very likely that shrimp from Florida could readily interbreed with shrimp from North Carolina.

The Texan population demonstrated a high level of mitochondrial DNA divergence relative to the other populations and low interbreeding ability with other populations. During interbreeding experiments, the Texas shrimps failed to outcross with other populations. The modality observed is interesting given that this species is a simultaneous hermaphrodite and shrimp were used in both male and female roles. Additional breeding experiments will have to be conducted using a higher replicate number to confirm that there is a total lack of interbreeding ability between Texas populations and those found in Florida. In the genus *Lysmata* it is common for bacterial infections to spread from dead embryos to compromise viable eggs; in addition the presence of eggs on a molt is an indication of high bacterial levels or stress on the female (A. L. Rhyne unpubl. obs.). These factors must be considered when determining if viable offspring can be produced. The results of the present study demonstrate both a difference in viability that is dependent on the modality of the cross and a disconnection between the interbreeding ability of populations and a high level of molecular divergence.

A high level of divergence is not unexpected when compared to other studies in the literature (for review see Avise 2004). A number of species have displayed a high degree of separation between the Gulf of Mexico and eastern Florida populations (Saunders et al. 1986, Reeb & Avise 1990, Vogler & DeSalle 1993, 1994, Schizas et al. 1999). These examples are not limited to species with low dispersal ability or sessile adult populations. The black sea bass *Centropristis striata* has been shown to exhibit a distinct mtDNA break between the Atlantic and Gulf populations (Bowen & Avise 1990). Some decapod species have established differences between the Gulf of Mexico and Atlantic Ocean populations. For example, the crab genera *Uca* and *Sesarma* both display ‘trans-Floridian’ species complexes (Felder & Staton 1994). These differences are most likely due to vicariant distribution, associated with the emergence of the Florida Peninsula during past glaciation events. Felder & Staton (1994) surmised that these ‘populations’ were in fact sister taxa, as reflected by the fact that species of both genera display remarkable differences in coloration pattern throughout their range. Other studies have reported less divergence between Gulf of Mexico and Florida populations. For example, the blue crab *Callinectes sapidus* displays a certain level of population structure over a wide geographic range, but lacks the degree of divergence observed in other species (McMillen-Jackson & Bert 2004). In *Lysmata wurdemanni*, every sampled population is distinct; the highest degrees of divergence were observed between the Texas population and those in Florida and North Carolina (~10% in COI). While this might suggest a cryptic species complex, interbreeding results demonstrate speciation in progress. Other crustaceans, such as the tidepool copepod *Tigriopus californicus*, exhibit even higher divergence (17 to 22% in COI) between central and southern Californian populations (Burton & Lee 1994, Burton 1996), but can still interbreed. There is still the possibility that hybridization between Florida Keys and Texas populations results in *F₂* breakdown, similar to hybrid breakdown observed after the first generation in the copepod *Eurytemora affinis* (Lee 2000). Additional sampling of *L. wurdemanni* may identify reproductively incompatible populations resembling the degree of genetic differentiation and reproductive incompatibility observed between some southern California and Mexican populations of *T. californicus* (Ganz & Burton 1995).

Sequence divergence at the mtDNA level has been used to infer reproductive isolation between closely related species. Knowlton et al. (1993) established that a divergence >6% in COI sequences suffices to produce reproductive isolation in the snapping shrimp *Alpheus*. The present study fails to find total reproductive isolation at ~10% divergence. However, comparisons of the amount of genetic divergence necessary for hybrid breakdown may not be appropriate between phylogenetically different taxa. Additionally, the stochasticity associated with coalescence times in mitochondrial genes is high (Hudson & Turelli 2003), and predictions or extrapolations based on a single gene may be highly inaccurate.

One of our a priori expectations was that the shrimps from the Florida Keys, which were collected in Florida
Bay, would be genetically similar to the other Floridian samples given that the Florida Current and Gulf Stream are close to the sampling areas. It now seems possible that *Lysmata wurdemanni* larvae in Florida Bay do not enter the Florida Current. Higher resolution sampling will be required to elucidate the dynamics of the population structure within the Florida Keys and to determine whether gene flow in this small geographic area is extensive or is limited, resulting in population subdivision along the eastern Florida coast. Perhaps there are several distinct populations within the Florida Keys and Florida Bay, where Florida Bay is disconnected from the Florida Keys and populations in Florida Bay are related to those in western Florida. Other marine taxa, such as the blue mussel *Mytilus edulis* have long-living larval stages, but the larval dispersal is limited to 10s of kilometers in southwest England (Gill & Hibish 2003). If the realized dispersal of *L. wurdemanni* is similar to that of *M. edulis*, then the significant population structure of *L. wurdemanni* could be explained and additional distinct populations should be expected. There are *L. wurdemanni* shrimp in the Florida Keys that exist in very close proximity to the Florida Current. Rhyne & Lin (2006) discuss the presence of several species of *Lysmata* occupying small rock ledges around the Key West Lakes, a small group of mangrove islands west of Key West, which is subdivided by the Northwest and Boca Grand Channels. This population could broadcast larvae into both the Gulf of Mexico and the Atlantic Ocean, and larvae would be expected to easily migrate out to the Florida Current. A population at this locality might hold the key to unlocking a possible genetic connectivity between populations of the Florida Bay and those along the eastern United States coast. Given that significant Gulf Stream transport is expected for populations along the eastern Florida coast and the southeastern United States, it is possible that populations along the east coast are the least divergent (0.56%).

There has been a great deal of attention paid to population connectivity in recent literature (Palumbi 2003, Cowen et al. 2006, Becker et al. 2007). This is primarily due to the importance of gene flow to the conservation and biodiversity of marine systems (Cowen et al. 2006). Studies have focused on the need to understand population structure before establishing marine reserves for the purpose of maintaining viable local populations. In the present study, *Lysmata wurdemanni*, a highly traded decapod in the marine aquarium industry, exhibits a significant genetic structure comprised of 3 distinct mitochondrial lineages at a large geographical scale. Restricted or closed populations are highly susceptible to local fishing pressures; thus, they must be managed differently from those with open populations. The population in Florida Bay is re-supplied by local recruitment and probably by other adjacent populations from the eastern Gulf of Mexico that were not sampled. Given that this population is under intense fishing pressure, with little to no fishery management, steps should be taken to control the exploitation of this species. Sound management plans for marine ornamental species should thus use information from genetic studies identifying species comprised of several restricted populations.

Beyond applications to fishery management, the present study provides an opportunity to use *Lysmata wurdemanni* as a model for species with long larval durations. Because there are several modes of larval development within the genus *Lysmata*, the population structure of additional species should also be examined. The closely related species *L. boggessi* Rhyne & Lin, 2006 and *L. pederseni* Rhyne & Lin, 2006 provide an interesting genetic comparison to *L. wurdemanni* because of their somewhat overlapping distribution (e.g. *L. boggessi*) or obligate symbiosis with tube sponges (e.g. *L. pederseni*).

In conclusion, *Lysmata wurdemanni* displays a marked population structure among the sampled locations, despite having an extended planktonic larval stage and a large reproductive capacity. The molecular data are consistent with the Gulf of Mexico/Florida genetic disconnection reported in several intertidal marine species (Roberts 1997). Our data support the distinctiveness of the Texas population as a subspecies or sibling species compared to those found in Florida or on the east coast of the United States. The role of the FLK populations remains unclear, and the inclusion of more populations, especially from the eastern Gulf of Mexico, is required in order to unravel the distribution of each genetically distinct population. It is likely that the populations along the northwestern Florida coast could be closely related to those in Texas through a series of stepping stones or that there are other intermediate populations between Florida and Texas maintained by highly localized recruitment. Both outcomes have a significant impact on the evolutionary genetics and the conservation of this species.

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