Abstract—Invasive red lionfish (Pterois volitans) have spread rapidly throughout the northern Gulf of Mexico (GOM) partly because of their high growth rate. Red lionfish were collected from the northern GOM across 3 ecological regions from 2012 through 2015. For male and female red lionfish, relationships between weight and total length (TL) were different by ecological region. Males achieved a greater mean weight adjusted for TL (333.6 g [standard error (SE) 3.6]) than females (195.1 g [SE 3.7]). A subsample of 1607 pairs of sagittal otoliths (from 744 males, 716 females, and 147 fish of unknown or undetermined sex) was used to assign ages. Ages ranged from 0.0 to 4.5 years (mean: 1.4 years), and these estimated ages and the dates of capture for specimens confirm the presence of red lionfish in the northern GOM in 2008, 2 years prior to the first detection of this species there. There were differences in age and growth between sexes within and among ecological regions, with males achieving higher growth rates and larger asymptotic lengths than females (all comparisons: \( P < 0.01 \)). These findings, coupled with other life history information, aid in discerning differences in distribution of red lionfish populations and are essential for creating management plans for mitigation of their effects on ecosystems.

Comparison of age and growth parameters of invasive red lionfish (Pterois volitans) across the northern Gulf of Mexico

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The invasive red lionfish (Pterois volitans) and devil firefish (P. miles) have been reported as established in U.S. waters of the western North Atlantic Ocean and in the Caribbean Sea, and the red lionfish has been found in the Gulf of Mexico (GOM) (Morris and Akins, 2009; Schofield, 2010; Fogg et al., 2017); however, the devil firefish has not yet been detected in the GOM (Johnson et al., 2016). It is difficult to distinguish between these 2 species except through genetic analysis, and data from previous studies are not necessarily separated by species. Therefore, throughout this paper, we use the term lionfish to refer to specimens that are not identified to species but are a potential combination of red lionfish, devil firefish, and hybrids between the 2 species.

Lionfish were first documented in the United States off Dania Beach, on the eastern coast of Florida, in 1985 (U.S. Geological Survey, Nonindigenous Aquatic Species Database, website, accessed December 2015) and later documented in the GOM in 2009 (Aguilar-Perera and Tuz-Sulub, 2010; Nonindigenous Aquatic Species Database, accessed December 2015). By 2012, lionfish were commonly captured in the northern GOM (Fogg et al., 2013), and histological evidence of spawning-capable lionfish in the northern GOM was first observed from fish captured in May 2012 (Brown-Peterson and Hendon, 2013). Efforts to research lionfish species have increased as their invasion has expanded, but most work has focused on the effects of lionfish...
species on native fish species and coral reef communities (Dahl and Patterson, 2014; Albins, 2015; Benkwitt, 2015; Ingeman and Webster, 2015; Rocha et al., 2015; Acero et al., 2019) and on aspects of landscape-level movements among reef systems and invasion control efforts (Frazer et al., 2012; Green et al., 2014; Tamburrolo and Côté, 2015). Results from a recent study indicate that red lionfish have unique morphological functional traits that reduce the theoretical ecological space of 5 coral reef mesopredators of the Caribbean Sea (Rojas-Vélez et al., 2019). However, data are limited for age and growth of lionfish species throughout the geographic range that they have invaded.

Knowledge of weight–length relationships, size and age structure, and growth patterns are important for the successful assessment and management of species. This information can also be used to assess the effects of invasive species on native species and the ecosystems they inhabit. For example, although data such as weight–length relationships are generally useful for quantifying changes in size or age structure that relate to potential overexploitation of a species (Berkeley et al., 2004; Dulvy et al., 2004), such data also could be used as indicators of success in management of invasive species (Pasko and Goldberg, 2014). Additionally, weight–length relationships could be used to examine changes in population structure during post-culling activities or following large-scale disease or environmental perturbation (e.g., an algal bloom). Although a number of studies have reported basic weight and length data for invasive lionfish (Barbour et al., 2011; Fogg et al., 2013; Dahl and Patterson, 2014; Edwards et al., 2014; Sabido-Itzá et al., 2016), few make comparisons between regions or sexes.

Estimations of age and growth relationships for species within invaded geographic ranges are important for describing spatially explicit variation in life history. Larger and older fish tend to affect ecosystems differently than smaller and younger individuals because diet and habitat use can change with age and size (Curtis et al., 2017; Mizrahi et al., 2017; García-Rivas et al., 2018). Age and growth patterns of lionfish vary geographically. Johnson and Swenarton (2016) verified their length-based model outputs with ages determined from a subsample of 100 sectioned otoliths from fish (age 0–3; maximum size of 342 mm in total length [TL]) captured offshore of Jacksonville, Florida. Additional information from other studies conducted outside of the GOM illustrates this variation. Lionfish captured in Onslow Bay, North Carolina, had a maximum age of 8 years, with more than 90% of these fish (<i>n</i>=814) <3 years old (Potts et al., 2010; Barbour et al., 2011). Two red lionfish collected off the coast of South Carolina, at sizes of 352 and 389 mm TL, were determined to be 5 and 6 years old, respectively (Meister et al., 2005).

Limited data from regions in the GOM and the Caribbean Sea indicate similar variation. Red lionfish collected from the Dry Tortugas in the Florida Keys had a maximum age of 7 years (Dubel1). Rodríguez-Cortés et al. (2015) provided the first growth and mortality estimates for the southern region of the GOM (Mexico), although the modeled lengths of red lionfish (<i>n</i>=776; range: 90–389 mm TL) from that study were not verified by using otoliths. Edwards et al. (2014), using otoliths from 110 male and 128 female lionfish captured off Little Cayman, found a maximum age of only 5 years and confirmed annual annuli formation for fish from that region of the Caribbean Sea. However, no ages determined from analysis of wild-caught lionfish are as old as the ages of lionfish held in captivity (30–33 years; Potts et al., 2010).

Lionfish species have invaded different regions of the GOM and the Caribbean Sea at different times (Schofield, 2010) and can be found in vastly different ecosystems (Barbour et al., 2010; Jud et al., 2011; Claydon et al., 2012; Ruttenberg et al., 2012) and different densities (Green and Côté, 2009; Darling et al., 2011; Dahl and Patterson, 2014). Therefore, we expected that age and growth parameters would vary by location, a notion that Villaseñor-Derbez and Fitzgerald (2019) have since verified. Additionally, rapid growth rates generally lead to successful invasion of an area by a species; therefore, it is important to estimate growth rates (Copp and Fox, 2007). Although age and growth data have been reported for portions of the invaded geographic range, growth of red lionfish could differ between the southern and northern regions of their non-native range (Barbour et al., 2011). Therefore, we hypothesized that the age structure of red lionfish is much younger in the northern GOM than in other invaded regions where they have been established for a longer period. The goal of this study was to determine if differences in weight and length and in age and growth relationships exist by sex or across the northern GOM.

Materials and methods

Initially, we did not identify our specimens to species, instead putting them in a complex that comprises both <i>Pterois volitans</i> and <i>P. miles</i> because it was unknown at the beginning of our study if both species occurred in the GOM (Hamner et al., 2007; Brown-Peterson and Hendon, 2013; Fogg et al., 2013) or if there were hybrids in the GOM. Results from subsequent work, including a study that used specimens we provided for genetic analysis, indicate that the red lionfish is the only species detected to date in the northern GOM (Johnson et al., 2016). Therefore, we identify all specimens that we collected in the northern GOM throughout our study as <i>P. volitans</i>. From 2012 through 2015, red lionfish were collected opportunistically every month (for details about specimen collection, see Fogg et al., 2017) across 3 ecological regions (or ecoregions) of the northern GOM: southeast (from the Florida Keys north to Anclote Keys, Florida), northeast (from the Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas; 1 fish was collected west of Galveston Bay) (Fig. 1). Clearly identified, ecologically relevant ecoregions in the GOM and the Caribbean Sea vary and are debated because authors use somewhat different environmental data and quantitative approaches to delineate ecoregions.

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Fogg et al.: Age and growth of invasive *Pterois volitans* in the northern Gulf of Mexico

**Figure 1**

Map of the locations (black dots) and ecological regions (ecoregions) where red lionfish (*Pterois volitans*) were sampled from 2012 through 2015 in the northern Gulf of Mexico. Three ecoregions were sampled in this study: southeast (Florida Keys north to Anclote Keys, Florida), northeast (Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas). The gray lines indicate depth contours from 10 to 200 m.

Most ecoregions are identified on the basis of known biogeographic criteria, such as thermal regimes, habitat, floral and faunal assemblage composition, sediment structure, currents and bathymetry, and coastal structural complexity. The northern tip of the Anclote Keys off the western coast of Florida, for example, has been identified as a point of a north–south ecoregion break (Beck and Odaya, 2001; Beck, 2003; Spalding et al., 2007) in the northern GOM. This boundary between the southeast and northeast ecoregions is used mostly because of the thermal differences between the 2 ecoregions; in the winter months, water temperatures are consistently lower in the northeast ecoregion than in the southeast ecoregion (temperature data were obtained from NOAA’s National Data Buoy Center, website). The Mississippi River often is used to divide the northern GOM into east and west ecoregions; however, we follow the ecoregion delineation based on Beck (2003) and Yáñez-Arancibia and Day (2004), dividing our study area into central and northeast ecoregions. Our northeast ecoregion, which is considered the east ecoregion by Beck and Odaya (2001), Beck (2003), and Yáñez-Arancibia and Day (2004), is characterized by less freshwater input, sandier sediments, and enhanced water clarity compared with the central ecoregion where increased freshwater input, lower salinities, and muddy sediment input dominate. The northeast ecoregion is dominated by seagrass meadows, but the central ecoregion is mainly salt marsh (Beck, 2003; Yanez-Aráncibia and Day, 2004).

Total length, measured in millimeters, and total weight, measured in grams, were recorded, and weight–length relationships were calculated by sex and ecoregion. These data were used to estimate the weight–length power functions. The weight–length data were log transformed (base 10) prior to an analysis of covariance (ANCOVA), with TL as the covariate. An ANCOVA was first completed for each sex separately, comparing the weight–length relationships found across each of the 3 ecoregions. If no significant differences were found across ecoregions, data were pooled and a second ANCOVA was completed to compare weight–length relationships for each ecoregion by sex. If the weight–length relationships for any ANCOVA violated the homogeneity of slopes assumption (parallelism), separate models were used. The estimated marginal means from these analyses were used to make comparisons between sexes of weight adjusted for mean TL. All ANCOVA were completed by using SPSS Statistics², vers. 20 (IBM Corp., Armonk, NY).

The sagittal otoliths were removed from red lionfish, and the left otolith was embedded and sectioned to a 300-µm thickness following Secor et al. (1991). Prepared otoliths

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2 Mention of trade names or commercial companies is for identification purposes only and does not imply endorsement by the National Marine Fisheries Service, NOAA.
Comparisons of weight–length relationships of red lionfish (Pterois volitans) for each ecological region (ecoregion) sampled in the northern Gulf of Mexico from 2012 through 2015: southeast (Florida Keys north to Anclote Keys, Florida), northeast (Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas). (A) Regression equations for comparisons by sex within each ecoregion. (B) Pairwise comparisons by sex between ecoregions. Between-sex and between-ecoregion comparisons of weight (W) were evaluated by using analysis of covariance with total length (TL) as the covariate.

### A

<table>
<thead>
<tr>
<th>Ecoregion</th>
<th>n</th>
<th>Regression equation</th>
<th>n</th>
<th>Regression equation</th>
<th>Comparison</th>
</tr>
</thead>
<tbody>
<tr>
<td>Southeast</td>
<td>857</td>
<td>$W = 2.00 \times 10^{-6} TL^{3.34}$</td>
<td>671</td>
<td>$W = 1.00 \times 10^{-6} TL^{3.44}$</td>
<td>$F_{1,1526}=12.68, P&lt;0.001$</td>
</tr>
<tr>
<td>Northeast</td>
<td>1181</td>
<td>$W = 3.00 \times 10^{-6} TL^{3.26}$</td>
<td>1239</td>
<td>$W = 3.00 \times 10^{-6} TL^{3.30}$</td>
<td>$F_{1,1423}=2.32, P=0.128$</td>
</tr>
<tr>
<td>Central</td>
<td>368</td>
<td>$W = 3.00 \times 10^{-6} TL^{3.30}$</td>
<td>354</td>
<td>$W = 2.00 \times 10^{-6} TL^{3.41}$</td>
<td>$F_{1,122}=6.84, P=0.090$</td>
</tr>
<tr>
<td>Pooled</td>
<td>2406</td>
<td>$W = 3.00 \times 10^{-6} TL^{3.29}$</td>
<td>2264</td>
<td>$W = 2.00 \times 10^{-6} TL^{3.37}$</td>
<td>$F_{1,4670}=21.96, P&lt;0.001$</td>
</tr>
</tbody>
</table>

### B

<table>
<thead>
<tr>
<th>Ecoregion comparison</th>
<th>$F_{1,203}=8.16, P=0.004$</th>
<th>$F_{1,1910}=27.72, P&lt;0.001$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Southeast vs. northeast</td>
<td>$F_{1,154}=1.42, P=0.223$</td>
<td>$F_{1,1590}=10.19, P&lt;0.001$</td>
</tr>
<tr>
<td>Northeast vs. central</td>
<td>$F_{1,122}=0.86, P=0.354$</td>
<td>$F_{1,1090}=0.61, P=0.436$</td>
</tr>
<tr>
<td>Southeast vs. central</td>
<td>$F_{2,2406}=4.17, P=0.015$</td>
<td>$F_{2,2284}=15.88, P&lt;0.001$</td>
</tr>
<tr>
<td>Pooled ecoregions</td>
<td>$F_{1,1910}=27.72, P&lt;0.001$</td>
<td></td>
</tr>
</tbody>
</table>

Results

The slopes of the weight–length relationships for male and female red lionfish were significantly different within the southeast ecoregion and did not differ in the 2 other ecoregions (ANCOVA: $F_{1,1526}=12.68, P<0.001$) (Table 1A, Fig. 2). The weight–length relationships from data pooled age data, and separate model parameters were determined for each sex and ecoregion for comparison (Diaz et al.).

Because the von Bertalanffy growth curve is nonlinear, a sum of squares reduction test (Schabenberger and Pierce, 2002) was used with statistical software SAS (vers. 9.4; SAS Institute, Inc., Cary, NC), instead of a traditional analysis of variance to determine if there were differences in growth between ecoregions and sexes by comparing nonlinear trends between groups ($\alpha=0.05$). The sum of squares reduction test was conducted by fitting a full and reduced model to the data. The test statistic ($F$) was calculated following this equation: $F=[(SSRR-SSRF)/(DFRR-DFRF)]/MSRF$, where SSRR and SSRF are the residual sums of squares from the reduced and full model, respectively, $DFRR$ and $DFRF$ are the residual degrees of freedom for the reduced and full model, respectively, and $MSRF$ is the mean sum of squares from the full model. This test evaluates the SSRR when the SSRR is removed from the model.

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Figure 2

Weight–length relationships by ecological region (ecoregion) for (A) male and (B) female red lionfish (Pterois volitans) collected from 2012 through 2015 in the northern Gulf of Mexico. Circles indicate data for fish collected in the southeast ecoregion (Florida Keys north to Anclote Keys, Florida), squares indicate data for fish from the northeast ecoregion (Anclote Keys north to Mobile Bay, Alabama), and triangles indicate data for fish from the central ecoregion (west of Mobile Bay to Galveston Bay, Texas). $R^2$=coefficient of multiple determination.

across all 3 ecoregions for red lionfish had significantly different slopes by sex (ANCOVA: $F_{1,4670}=21.96$, $P<0.001$; Table 1A), indicating that females had a steeper weight–length relationship than males (Table 1A, Fig. 3) and, therefore, had a greater weight at a given TL. Pairwise comparisons of weight–length relationships by ecoregion (Table 1B) revealed a significant difference for male red lionfish only between the southeast and northeast ecoregions (ANCOVA: $F_{1,2038}=8.16$, $P=0.004$; Table 1B); whereas, significant differences were observed for female red lionfish in all comparisons between ecoregions except between the southeast and central ecoregions (ANCOVA: $F_{1,1025}=0.61$, $P=0.436$; Table 1B). However, in all ecoregions, males attained larger weights when adjusted for TL than females (Table 2). For a number of comparisons of ecoregions (Table 1B), weight–length relationships were significantly different, but the differences in weight were minimal on the basis of their estimated marginal means adjusted for TL (females: 18.21 g; males: 22.75 g; Table 2), and male red lionfish achieved a greater mean weight (333.62 g [standard error (SE) 3.58]) compared to females (195.13 [SE 3.69]) (Table 2).

From the 4250 pairs of otoliths that were extracted, a subsample of 1607 pairs of otoliths (744 males, 716 females, and 147 fish of unknown or undetermined sex) were randomly selected and processed from red lionfish ranging in size from 81 to 434 mm TL. Age agreement was reached for 1412 pairs of otoliths (87.9%) from that subsample. The APE for analysis of otoliths from the southeast ecoregion was 7.7, the APE for the northeast ecoregion was 14.2, and the APE for the central ecoregion was 11.7. With analysis results from all 3 ecoregions pooled, the APE between readers was 12.1. Annual increment formation was confirmed by using marginal increment analysis, with marginal increment widths most complete in May and gradually decreasing until a minimum index of completion
Figure 3
Weight–length relationships for male (squares) and female (triangles) red lionfish (*Pterois volitans*) collected from 2012 through 2015 in the northern Gulf of Mexico. Data are pooled for all sampled ecological regions. $R^2$=coefficient of multiple determination.

Table 2
Estimated marginal mean weight, in grams and adjusted for total length in millimeters, for invasive red lionfish (*Pterois volitans*) collected from 2012 through 2015 in 3 ecological regions (ecoregions) in the northern Gulf of Mexico: southeast (Florida Keys north to Anclote Keys, Florida), northeast (Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas). Estimates are given for females, males, and sexes pooled with standard errors of the mean (SEs) in parentheses.

<table>
<thead>
<tr>
<th>Sex</th>
<th>Ecoregion</th>
<th>n</th>
<th>Estimated weight (SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
<td>Southeast</td>
<td>668</td>
<td>187.98 (1.69)</td>
</tr>
<tr>
<td>Female</td>
<td>Northeast</td>
<td>1227</td>
<td>193.88 (1.22)</td>
</tr>
<tr>
<td>Female</td>
<td>Central</td>
<td>348</td>
<td>206.19 (2.35)</td>
</tr>
<tr>
<td>Female</td>
<td>Pooled</td>
<td>2243</td>
<td>195.13 (3.69)</td>
</tr>
<tr>
<td>Male</td>
<td>Southeast</td>
<td>844</td>
<td>325.20 (2.36)</td>
</tr>
<tr>
<td>Male</td>
<td>Northeast</td>
<td>1101</td>
<td>334.50 (2.01)</td>
</tr>
<tr>
<td>Male</td>
<td>Central</td>
<td>346</td>
<td>347.95 (3.61)</td>
</tr>
<tr>
<td>Male</td>
<td>Pooled</td>
<td>2291</td>
<td>333.62 (3.58)</td>
</tr>
<tr>
<td>Pooled</td>
<td>Southeast</td>
<td>1512</td>
<td>258.75 (4.79)</td>
</tr>
<tr>
<td>Pooled</td>
<td>Northeast</td>
<td>2328</td>
<td>253.52 (3.86)</td>
</tr>
<tr>
<td>Pooled</td>
<td>Central</td>
<td>694</td>
<td>294.96 (7.07)</td>
</tr>
</tbody>
</table>

was reached in October and November, indicating the beginning of annuli formation (Fig. 4). Ages of red lionfish ranged from 0.5 to 4.5 years (Fig. 5), with 93% of aged red lionfish <2 years old. The smallest and youngest specimen collected in this study was a 81-mm-TL, 0.5-year-old fish of unknown sex, and the oldest specimen was a 380-mm-TL, 4.5-year-old male. The largest fish collected was a 434-mm-TL male, but its estimated age was 3.5 years. The state record for the longest lionfish caught in the northern GOM (459 mm TL) was for a fish collected off Pensacola, Florida (record available from website).

There were significant differences in age and growth parameters by sex and by ecoregion (all comparisons: $P<0.001$; for comparisons of modeled estimates with the sum of squares reduction test, see Table 3; for observed values, see Figure 6). Female red lionfish from the southeast ecoregion had the highest growth rate ($K$) and asymptotic length ($L_\infty$) and achieved a greater length at age than...
females in the other 2 ecoregions (Table 3A, Fig. 6A). Similarly, male red lionfish from the southeast ecoregion also had the highest $K$ and $L_\infty$ values and achieved a greater length at age than males in the other 2 ecoregions (Table 3A, Fig. 6B). Data pooled by sex revealed a similar pattern, with the highest $K$ and $L_\infty$ values estimated for red lionfish from the southeast ecoregion and lowest values estimated for red lionfish from the central ecoregion (Table 3A, Fig. 6C). Separate comparisons were made by sex for each ecoregion, with male red lionfish achieving higher $K$ and $L_\infty$ values than females in all 3 ecoregions (all comparisons: $P<0.05$; Table 3B, Fig. 7, A–C). Model
Table 3

Estimates of von Bertalanffy growth curve parameters for invasive red lionfish (*Pterois volitans*) collected from 2012 through 2015 in 3 ecological regions (ecoregions) in the northern Gulf of Mexico: southeast (SE; Florida Keys north to Anclote Keys, Florida), northeast (NE; Anclote Keys north to Mobile Bay, Alabama), and central (C; west of Mobile Bay to Galveston Bay, Texas). The parameters are growth rate ($K$), asymptotic length ($L_\infty$), given in millimeters in total length, and theoretical age when the length is zero ($t_0$). (A) Comparisons among ecoregions by sex. (B) Comparisons between sexes by ecoregion. A sum of square reduction test was used to determine differences in growth between ecoregions and sex by comparing nonlinear trends between groups ($\alpha=0.05$).

A

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Pooled</th>
<th>Female</th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SE</td>
<td>NE</td>
<td>C</td>
</tr>
<tr>
<td>$K$</td>
<td>0.569</td>
<td>0.544</td>
<td>0.539</td>
</tr>
<tr>
<td>$L_\infty$</td>
<td>423.0</td>
<td>393.0</td>
<td>389.0</td>
</tr>
<tr>
<td>$t_0$</td>
<td>−0.155</td>
<td>−0.079</td>
<td>−0.341</td>
</tr>
<tr>
<td></td>
<td>$F_{12,1412}=27.143$, $P&lt;0.001$</td>
<td>$F_{12,626}=7.303$, $P&lt;0.001$</td>
<td>$F_{12,695}=12.606$, $P&lt;0.001$</td>
</tr>
</tbody>
</table>

B

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Southeast</th>
<th>Northeast</th>
<th>Central</th>
<th>Pooled</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male</td>
<td>Female</td>
<td>Male</td>
<td>Female</td>
</tr>
<tr>
<td>$K$</td>
<td>0.576</td>
<td>0.574</td>
<td>0.547</td>
<td>0.549</td>
</tr>
<tr>
<td>$L_\infty$</td>
<td>426.0</td>
<td>382.0</td>
<td>394.4</td>
<td>366.8</td>
</tr>
<tr>
<td>$t_0$</td>
<td>−0.170</td>
<td>−0.165</td>
<td>−0.086</td>
<td>−0.089</td>
</tr>
<tr>
<td></td>
<td>$F_{8,453}=2.412$, $P=0.008$</td>
<td>$F_{8,489}=2.012$, $P=0.030$</td>
<td>$F_{8,379}=2.362$, $P=0.010$</td>
<td>$F_{8,1321}=16.226$, $P&lt;0.001$</td>
</tr>
</tbody>
</table>

parameters determined from data pooled across all 3 ecoregions indicate that males also had higher $K$ and $L_\infty$ values than females (Table 3B, Fig. 7D).

Discussion

The results of this study reveal significant ecoregion- and sex-specific patterns in age, growth, and weight–length relationships of red lionfish. The $K$ values of red lionfish collected from the southeast ecoregion were higher than those of red lionfish from the northeast and central ecoregions. Although it was expected that $K$ values would be greater for the red lionfish collected in the southeast ecoregion, the $L_\infty$ values of many fish species are usually lower in the more southern regions of their geographic range than in the more northern regions (Boehlert and Kappenman, 1980) because fish species from northern latitudes typically achieve larger sizes than fish species from southern latitudes (Lindsey, 1966). The observed anomaly in $L_\infty$ values may also be a result of density-dependent growth, as has been documented in invasive red lionfish on small artificial reefs in the Bahamas (Benkwitt, 2013). Red lionfish collected from the northeast ecoregion in particular came from much smaller and isolated artificial and natural reefs and were found in much higher densities compared with densities observed on the reefs in the southeast ecoregion (senior author, unpubl. data).

The results of our marginal increment analysis indicate that annuli on otoliths from red lionfish were most complete in the spring (March–May) and least complete in the fall (September–October), a finding similar to that for another scorpaenid, the native blackbelly rosefish (*Helicolenus dactylopterus*), off the coasts of North and South Carolina (White et al., 1998). In contrast, black scorpionfish (*Scorpaena porcus*) from the Adriatic Sea had their most complete annuli in the late summer (July–September; La Mesa et al., 2010). Comparable thermal regimes between the northern GOM and North and South Carolina likely are the reason for the similar trends in annuli formation between invasive red lionfish and native blackbelly rosefish; similar trends have been documented in red snapper (*Lutjanus campechanus*) from similar thermal regimes (Nelson and Manooch, 1982). Black scorpionfish from the Adriatic Sea likely have delayed annuli completion compared with the annuli formation of the red lionfish in our study because of the relatively cooler sea-surface temperatures (~11°C) that they experience in the spring. Sea-surface temperatures in the Adriatic Sea warm to above 20°C in July (La Mesa et al., 2010), similar to sea-surface temperatures observed in March in the northern GOM. Therefore, water temperature may drive annuli completion in these species, as has been reported for other species and regions (Pearson, 1996).

Age estimation was challenging for some red lionfish in our study, with an APE of 12.1 between readers for
The von Bertalanffy growth curves and associated equations for (A) female and (B) male red lionfish (*Pterois volitans*) as well as for (C) sexes pooled. Also plotted are observed lengths at age, by the 3 ecological regions (ecoregions) where red lionfish were collected from 2012 through 2015 in the northern Gulf of Mexico: southeast (Florida Keys north to Anclote Keys, Florida), northeast (Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas).

Otoliths from fish from all ecoregions pooled. This value is similar to the APE of 7 reported for analysis of otoliths from fish collected off northeast Florida (Johnson and Swenarton et al., 2016). However, all of our APE values are low compared with those of Edwards et al. (2014), who reported an APE of 58 between readers for lionfish collected in the Caribbean Sea. Differences in regional age determination may be expected because annuli in lionfish collected in tropical waters are likely to be more difficult to distinguish than annuli in lionfish living in more temperate regions. Lionfish in tropical waters likely have relatively consistent growth because of minimal variation in water temperatures, and fish in temperate waters experience periods of slow and fast growth (Pitcher and Hart, 1982).

The age of red lionfish from the northern GOM ranged from 0 to 4.5 years ($\bar{x}$=1.35), ages that are markedly lower than the maximum reported age of 30–33 years for a specimen held in an aquarium (Potts et al., 2010). Interestingly, the oldest age estimated in this study was 4.5 years...
The von Bertalanffy growth curves and associated equations for red lionfish (*Pterois volitans*). Also plotted are observed lengths at age for female (triangles) and male (squares) red lionfish collected from 2012 through 2015 in the northern Gulf of Mexico, by ecological region (ecoregion), (A) southeast (Florida Keys north to Anclote Keys, Florida), (B) northeast (Anclote Keys north to Mobile Bay, Alabama), and (C) central (west of Mobile Bay to Galveston Bay, Texas), as well as for (D) all ecoregions pooled.

**Figure 7**
for a fish collected in November 2012, and the data from our study confirm the presence of red lionfish in the northern GOM as early as 2008, 2 years prior to the first detection in the region in 2010. Edwards et al. (2014) also found a single lionfish (Pterois sp.) with an estimated age that indicates it was present in the Caribbean Sea before the first detection off Little Cayman in 2010. Such delays in documented detections are expected because invasive species often are not detected immediately after introduction because of lag times associated with species expansion (Crooks and Soulé, 1999).

Age distribution of marine fish species is an important factor for assessing the health of a population (Berkeley et al., 2004). Typically, an established, healthy population will exhibit a “well-balanced” age structure (Brunel and Piet, 2013) with numerous larger, older individuals. In the northern GOM, 93% of red lionfish in our study were ≤2 years old in all 3 ecoregions. Similar results have been reported off Little Cayman (Edwards et al., 2014) and in the Atlantic Ocean off northeast Florida (Johnson and Swenarton, 2016) and North Carolina (Barbour et al., 2011), where the majority (>90%) of lionfish were ≤3 years old. The higher proportion of fish between the ages of 2 and 3 years found in Little Cayman and off northeast Florida and North Carolina is likely the result of lionfish having invaded those locations earlier than the northern GOM (Schofield, 2010).

Although red lionfish can live much longer than what has been described in this study and in studies in other invaded geographic areas, the truncated age-class distribution observed in this study is a further indication that the population of red lionfish in the northern GOM may still be stabilizing in the region because older individuals are not present or not captured. Red lionfish from the southern GOM that were aged in 2012 appeared to be much younger, with age and growth parameters that were much lower than those reported for this study and previous studies (Rodriguez-Cortés et al., 2015). This difference in age structure is likely a result of red lionfish being collected within the first 2 years of their invasion in the southern GOM. The difference also could be due to aging techniques: we determined age on the basis of otolith analysis, but Rodriguez-Cortés et al. (2015) estimated ages on the basis of size-frequency analysis that was not verified by using otoliths. It is important to now establish up-to-date region- and sex-specific age and growth parameters so that, in the future when the invasion of red lionfish theoretically has stabilized across the region (Benkwitt et al., 2017; Côté and Smith, 2018), the same parameters can be estimated to evaluate the success of the invasion.

There was an inconsistent pattern of differences in weight–length relationships among ecoregions with only minor differences in weight among ecoregions: 18.21 g for females and 22.75 g for males.

Differences in weight–length relationships of lionfish have been reported throughout the range of the invasion among regions that are larger than the ecoregions used in our study (Suppl. Table). Some published comparisons of weight–length relationships have been based on data pooled across a range of locations within the sampled region (Barbour et al., 2011; Edwards et al., 2014; Sabido-Itzá et al., 2016), and the relationships derived from those pooled data differ from those presented here for the northern GOM. Another study compared weight–length data pooled by year, reporting a significant difference between the first and last year (2011–2013) (Dahl and Patterson, 2014). In contrast, Benkwitt et al. (2017) documented no changes in size structure in lionfish from the Bahamas over a 10-year period. Recently, Pusack et al. (2016) reported that red lionfish in their native range grow at a slower rate and achieve smaller maximum sizes than those in the range of their invasion. Finally, lionfish in the range they have invaded may be less susceptible to predation because of their larger size and would also be able to consume larger prey items (see review in Côté and Smith, 2018).

In this study, male red lionfish found in the northern GOM achieved greater weight and length than females. In contrast, results from early work in the northern GOM (Fogg et al., 2013) indicate no significant differences in weight–length relationships by sex for red lionfish. One explanation for these differences between studies may be that the data from Fogg et al. (2013) came from early in the invasion and, therefore, likely had not yet reached an asymptotic value. Although von Bertalanffy growth parameters were not reported by sex in other age and growth studies on invasive lionfish species (e.g., Potts et al. 2010; Barbour et al. 2011; Rodríguez-Cortés et al. 2015; Johnson and Swenarton 2016), age and growth and weight–length data quantified in our study confirm that sexual dimorphism exists with males growing larger and faster than females. Sexual dimorphic growth was documented also for invasive lionfish in Little Cayman by using otolith analysis, but weight–length relationships were not evaluated (Edwards et al., 2014). Males in our study achieved a greater length-at-age than females and, similar to males examined by Edwards et al. (2014), had significantly larger K and L∞ values than females. Males of the related scorpeanid species, blackbelly rosefish, also appear to grow faster and slightly larger than females (White et al., 1998). In contrast, female black scorpionfish, native to the eastern Atlantic Ocean, Mediterranean Sea, and Black Sea, attain larger sizes than males (Bilgin and Çelik, 2009).

The differences observed between male and female red lionfish in age and growth and in weight–length relationships are likely a result of the greater resources that females allocate to reproductive output (Gadgil and Bossert, 1970). Female red lionfish mature in their first year of life and are capable of reproducing every few days during 11 months of the year (Fogg et al., 2017). These elevated and constant reproductive rates likely result in more...
energy shunted to reproduction than to growth; therefore, growth in female red lionfish is reduced upon maturation. This trade-off of reduced growth in females as the energy is redirected to reproduction has been described in numerous fish species (Reznick, 1983; Roff, 1983; Parker, 1992).

Another potential reason that males grow larger than females is that males need to increase their chances of success in male rivalries and physical combat and, therefore, their opportunities for mating (Shine, 1989). Evidence of combat between large red lionfish (>350 mm TL, presumably males) has been observed in the northern GOM in the form of abrasions across the body. Recently, agonistic behavior between 2 large lionfish (Pterois spp.) has been observed and documented in Honduras (Fogg and Faletti, 2018).

Growth rates and other life history traits vary by region in other marine fish species (Choat and Axe, 1996; Ruttenberg et al., 2005). The K value for red lionfish in the northern GOM is greater than that reported for fish from Little Cayman (Edwards et al., 2014), North Carolina (Barbour et al., 2011), and northeast Florida (Johnson and Swenarton et al., 2016), although K values reported for fish from the Florida Keys (Swenarton et al., 2015) and Yucatan, Mexico (Rodriguez-Cortés et al., 2015), were much greater than those in all other studies (Table 4). The higher K value observed in the northern GOM could explain why densities of red lionfish are higher there than anywhere else in the geographic range that they have invaded (Dahl and Patterson, 2014). The pattern for age and growth of red lionfish in the northern GOM appears to be most similar to that of lionfish from the Atlantic Ocean off northeast Florida and North Carolina (Table 4). Similarities and differences in age and growth among studies could, in part, be driven by environmental thermal regimes (Lyons et al., 2017; Barker et al., 2018) in addition to other biological and ecological factors (South et al., 2017). However, age and growth will need to be reevaluated in the future because our study examined samples collected early in the invasion of the northern GOM region.

Age and growth data are important to assess potential changes to the population structure of red lionfish that result from implementation of management plans, disease events, and environmental incidents such as an algal bloom. For example, Chagaris et al. (2017) modeled how potential strategies for management of lionfish may affect several recreationally and commercially important native fish species on the West Florida Shelf. The age and growth relationships of lionfish used in their model were based on fish from outside of the region of the West Florida Shelf and, therefore, may not have correctly reflected population dynamics. Moreover, other management plans from agencies around the invaded region specifically mention the need for regional age and growth data (Morris5; ANSTF6, Johnston et al., 2015), and these plans noted the need for ongoing research. Accurate and region-specific weight–length, age, and growth data are vital for developing age-structured population models that can be used to evaluate potential effects of targeted removals on the red lionfish population (Barbour et al., 2011) and the potential population-level effects of large-scale disease for red lionfish and devil firefish (Harris et al.7). These data can also be applied to behavioral studies for which only size information is available (e.g., Garcia-Rivas et al., 2018).

Graham and Fanning (2017) reviewed management plans for invasive lionfish species in the Caribbean Sea, but research updates are needed so that future management plans for invasive lionfish prevention and management plan, 48 p. Prepared by the Invasive Lionfish Control Ad-hoc Committee of the ANSTF [Available from website.]


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### Table 4

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Southeast</th>
<th>Northeast</th>
<th>Central</th>
<th>Pooled</th>
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<tr>
<td>L&lt;sub&gt;∞&lt;/sub&gt;</td>
<td>423.0</td>
<td>393.0</td>
<td>389.0</td>
<td>400.2</td>
</tr>
<tr>
<td>K</td>
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<td>0.544</td>
<td>0.539</td>
<td>0.560</td>
</tr>
<tr>
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<td>−0.155</td>
<td>−0.079</td>
<td>−0.341</td>
<td>−0.210</td>
</tr>
</tbody>
</table>

Estimates of von Bertalanffy growth curve parameters, with sexes pooled, for invasive red lionfish (Pterois volitans) collected from 2012 through 2015 in 3 ecological regions (ecoregions) in the northern Gulf of Mexico: southeast (Florida Keys north to Anclote Keys, Florida), northeast (Anclote Keys north to Mobile Bay, Alabama), and central (west of Mobile Bay to Galveston Bay, Texas). The parameters are growth rate (K), asymptotic length (L<sub>∞</sub>, given in millimeters in total length [TL]), and theoretical age when the length is zero (t<sub>0</sub>). Because of truncated data sets (i.e., a lack of red lionfish <100 mm TL), a sum of squares reduction test was used to compare model parameters between all ecoregions and sexes. All comparisons were significantly different (P<0.01). For comparison, parameters pooled by sex are included for specimens from North Carolina (Barbour et al., 2011), Little Cayman (Edwards et al., 2014), Florida Keys (Swenarton et al., 2016), northeast (NE) Florida (Johnson and Swenarton et al., 2016), and Yucatan, Mexico (Rodriguez-Cortés et al., 2015).
activities, including local “culling” programs (Green et al., 2017; Mizrahi et al., 2017), lead to more accurate forecasting (Johnston et al., 2017) and effective regional management. Information from regional demographic data sets, coupled with identification of spatial and temporal patterns in the harvest vulnerability of older and larger lionfish, can be used to identify regions or seasons that need better protection or regulation (Zhou et al., 2010; Tobin et al., 2013). Data from this study can inform development of region-specific management strategies for controlling the invasion of lionfish in the GOM. This study provides the first regional estimates of life history parameters for red lionfish in the northern GOM, where populations of this invasive species appear to be not only established but also expanding their range (Dahl and Patterson, 2014; Dahl et al., 2016; Fogg et al., 2017). Life history information provided here can be incorporated into studies of competition between lionfish and reef fish species (see Rojas-Vélez et al., 2019) that will enhance management decision-making and stock assessments.

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