Variation in cheliped form in two species of squat lobsters (Decapoda: Anomura) from Chile

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ABSTRACT

The current study describes the variation in cheliped form of two species of squat lobsters that inhabit the continental margin off Chile: Cervimunida johni and Pleuroncodes monodon. We compared their cheliped form in the context of the reproductive strategy. The general tendency of form variation of both species is similar: chelipeds change, on average, from longer and narrower pollex with short manus to a relatively shorter and wider pollex with longer manus from small to large individuals, respectively. The degree of cheliped arching was greater in males than in females of similar carapace length, and only C. johni males showed fully arched morphology. The allometric trajectories (cheliped shape vs. carapace length) were largely aligned with vectors of mean shape difference in both species. Cheliped form variation of C. johni (extended mate-guarding) resembles the cheliped morphology reported for the related species Munida rugosa more than P. monodon (short mate-guarding). Our results are consistent with previous finding and suggest that the cheliped form variation (from straight and slender to a fully arched morphology) is, or has been, subject to sexual selection through male-male competition for mates.

Descriptors: Cervimunida johni, Pleuroncodes monodon, Geometric morphometric, Reproductive strategy.

RESUMO

O presente estudo contém uma descrição da variação da forma dos quelípodos de duas espécies de lagostas que habitam a margem continental do Chile: Cervimunida johni e Pleuroncodes monodon. Foi comparada a forma dos quelípodos das espécies em relação à estratégia reprodutiva das espécies. Das espécies estudadas, C. johni apresenta um quelípodo arqueado, uma relação mais estreita entre a forma do quelípodo e comprimento da carapaça e um maior dimorfismo sexual. Apesar de que a tendência geral da variação da forma dos quelípodos de ambas as espécies é quase semelhante, a variação morfológica dos quelípodos C. johni ("extended mate-guarding") assemelha-se à relatada para M. rugosa mais do que assemelha-se P. monodon ("short mate-guarding"). Nossos resultados são consistentes com os resultados anteriores e sugerem que a variação na forma do quelípodo (de uma forma reta e fina para um formato arqueado) é ou foi sujeita à seleção sexual através da competição entre machos.

Descritores: Cervimunida johni, Pleuroncodes monodon, Morfometria geométrica, Estratégia reprodutiva.
INTRODUCTION

The yellow squat lobster *Cervimunida johni* PORTER 1903 and the red squat lobster *Pleuroncodes monodon* H. MILNE EDWARDS 1837 inhabit the oxygen minimum zone on the continental margin off Chile. In spite of being target of a trawl fishery operating on the continental margin along the Chilean coast, the general biology of these two species is only poorly known (THIEL and LOVRICH, 2011). Few studies have focused on the reproductive biology of these species, especially in *P. monodon*: PALMA (1994) and RIVERA and SANTANDER (2005) have reported the presence of larval stages in the plankton, and PALMA and ARANA (1997) have studied the developmental stages of embryos in the egg mass. ESPINOZA-FUENZALIDA et al. (2012) reported that females of both species mate during the intermolt period and become receptive shortly after having released larvae from a previous brood, when they form pairs with males. The interbrood interval (from larval release until having a subsequent brood) differs between the two species, being considerable longer in females of *P. monodon* than in *C. johni*. In addition, both species also differ in terms of sexual size dimorphism, duration of copulatory mate-guarding and mate-finding ability (ESPINOZA-FUENZALIDA et al., 2012). Based on differences in the reproductive biology, the latter authors suggested that males of *C. johni* aggressively compete for access to females while males of *P. monodon* might use the “pure searching” strategy to find receptive females. Accordingly, it is expected that the reported differences in the reproductive strategy of *C. johni* and *P. monodon* should be reflected in their morphologies, especially in those structures directly or indirectly related to reproduction.

Morphological variations related to different reproductive strategies have been extensively reported in decapods, particularly in cheliped morphology (see MARIAPPAN et al., 2000). In decapod crustaceans, the first pereopod generally bears a chela (claw) formed by a modification of the dactylus and propodus and is commonly known as cheliped. This appendage is typically used in feeding, defense against predators and agonistic behavior, and has therefore been considered a multifunctional organ (BROWN et al., 1979; LEE, 1995). Due to their simple basic structure, their various functions and their well-known mechanics, chelipeds are good models to study the evolution of morphology (CLAVERIE and SMITH, 2007). In squat lobsters, the patterns of cheliped form (shape and size) variation have been related to sexual dimorphism and mating interactions. Specifically, CLAVERIE and SMITH, (2010), reported an unusual cheliped size and shape dimorphism, in addition to a considerable variation in cheliped form, from straight and slender to an arched morphology in *Munida rugosa* (Fabricius), a common galatheid from the northeast Atlantic. These morphological differences, which are to some extent similar to differences between the well-known cutter and crusher cheliped shape in other decapods (YAMADA and BOULDING, 1998), has been related to sexual selection through male-male competition for mates. The arched morphology, which is stronger and is adapted to inflict puncture wounds during intense agonistic interactions, appeared only in larger individuals (usually above the minimal size of morphological maturity) and compared with equivalent-sized males, the arched morphology does not develop fully in females (see also CLAVERIE and SMITH, 2007).

Considering the reported difference in the reproductive strategies of *C. johni* and *P. monodon*, we hypothesized that cheliped form variation differs between the two species. Specifically we expected that cheliped form variation in the mate-guarding *C. johni* resembles that reported for *M. rugosa*, in which males also had been suggested to compete aggressively for reproductive females (CLAVERIE and SMITH, 2007, 2010). The main objective of this study was to evaluate the cheliped size and shape in order to: (1) determine whether the patterns of cheliped form described for *M. rugosa* were present in *C. johni* and in *P. monodon*, to (2) evaluate the relation between chelipeds shape and carapace length (allometry), and to (3) estimate the level of sexual dimorphism and whether the differences between sexes were caused by differences in cheliped size. In this purpose, we followed the general approach used by CLAVERIE and SMITH (2007, 2010).

MATERIAL AND METHODS

**Sample collection**

During May 2010, one hundred individuals each of *C. johni* (PORTER) and *P. monodon* (H. MILNE EDWARDS) were obtained from the commercial trawl fishery operating on the continental margin off the coast of Coquimbo, Chile (29°59’ S; 71°23’ W). We paid special attention that the selected individuals had the carapace and both chelipeds intact, in addition to having no injuries or marks of any previous molting problem. Specimens were sorted by sex based on the shape of the pleopods (BABA et al., 2011) and body size was indexed by means of the
carapace length (CL, linear measurement from the base of the rostral spine to the posterior margin of the carapace at the midline), measured with a digital calliper to 0.01 mm accuracy (Table 1).

Table 1. Information of sample composition. n: sample size; CL: carapace length range (in mm) and sex-specific percentage of correct assignation after leave-one-out cross-validation. The percentage of size-corrected variables is given in parentheses (further explanations in the results section).

<table>
<thead>
<tr>
<th>Species</th>
<th>Sex</th>
<th>n</th>
<th>CL (mm)</th>
<th>% of correct assignation</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. johni</td>
<td>female</td>
<td>43</td>
<td>25.3 - 38.3</td>
<td>97.6 (51.1)</td>
</tr>
<tr>
<td></td>
<td>male</td>
<td>57</td>
<td>29.8 - 49.2</td>
<td>94.7 (49.1)</td>
</tr>
<tr>
<td>P. monodon</td>
<td>female</td>
<td>47</td>
<td>29.4 - 36.9</td>
<td>87.2</td>
</tr>
<tr>
<td></td>
<td>male</td>
<td>53</td>
<td>32.2 - 39.9</td>
<td>86.7</td>
</tr>
</tbody>
</table>

Data acquisition and image analysis

The lateral surface of each cheliped (right and left) was photographed with a digital camera (NIKON Coolpix L20, 5 megapixels) in macro mode, setting the picture plane parallel to the cheliped lateral surface (Fig. 1). The photographic procedure (e.g. avoiding distortions and considering scaling procedure and cheliped orientation) was replicated from CLAVERIE and SMITH (2010). Briefly, each cheliped was held extended and horizontal for the photograph using a custom made jig, and a small translucent elastic band was used to hold the cheliped closed. Since the propodus and dactylus are only slightly curved laterally, analysis of the cheliped form in two dimensions was considered adequate and chelipeds were analyzed using the 2D landmark-based geometrics method (ROHLF and MARCUS, 1993). Images were compiled, scaled and digitized in the TpsUtil and TpsDig programs (ROHLF, 2010a, b). In order to achieve a good representation of the cheliped form, 6 landmarks (BOOKSTEIN, 1997) were digitized (Fig. 1). Landmark configuration was homologous to those used previously by ROSENBERG (2002) for fiddler crabs (Uca spp.) and by CLAVERIE and SMITH (2007, 2010) for Munida rugosa, and was chosen considering their homology and their ability to capture the general shape of chelipeds (Fig. 1). Landmark configurations were optimally aligned using a Generalized Procrustes Analysis (GPA; ROHLF and SLICE, 1990) to remove the effects of translation, rotation and scale. After GPA, shape variables (Procrustes coordinates) and centroid size (CS) were used in the morphometric and statistical analyses. CS was computed as the square root of the sum of squares of the distances from all landmarks to their centroid and was used as a proxy of cheliped size. In order to control and exclude any specimen strongly deviating from the average, both the total sample and within each sex-specific group were inspected for outliers according to the standard procedure reported in MorphoJ program (KLINGENBERG, 2011). Due to the fact that the preliminary screening for outliers (results not shown) showed a good correspondence between the observed data and shape distances expected under a multivariate normal distribution model, we did not exclude any specimens from the analysis.

Data analysis

For the purpose of this study, the geometric morphometric and the statistical analyses were performed separately for both species with the MorphoJ program v1.05a (KLINGENBERG, 2011). In order to quantify the relative magnitudes of size and shape variation at different levels (sexes, individuals and appendage sides), a preliminary Procrustes ANOVA (KLINGENBERG and MCINTYRE, 1998) was carried out. Briefly, differences in cheliped size and shape were tested by partitioning the total variance into its components, which were statistically compared in a hierarchical design (i.e. to assess if sex differences were larger than differences between individuals within each sex, and whether these were larger than differences between appendage sides of the same individual). Results revealed that appendage side (shape heterochely) only differ in C. johni (F = 14.68 and F = 11.23 for females and males, respectively; d.f = 8 for both). Nevertheless, the relative amount of shape variation (Mean Squares) due to appendage side was negligible in relation to the strong differences in cheliped shape between sexes and among individuals. Results for cheliped size largely mirrored those for shape, but did not differ among appendage side in any case. Based on these results, the
appendage sides were pooled within each individual and average cheliped data (average size and shape of the left and right cheliped) were used throughout all subsequent analyses.

**Form variation**

The patterns of form variation were assessed by performing a Principal Component Analysis (PCA) on Procrustes coordinates. PCA allows visualizing and exploring the trend in form variation by means of a graphical representation of the form across a morphospace of reduced dimensions defined by the scores of the first principal components (ZELDITCH et al., 2012).

**Cheliped allometry and sexual dimorphism**

The relationship between cheliped shape and carapace length was evaluated by means of a multivariate regression of shape variables on CL. The statistical significance was tested with permutation tests against the null hypothesis of independence (GOOD, 2000), and the strength of the association was assessed by means of the percentage of shape variance explained by CL (KLINGENBERG, 2011). The regression routine computes a regression score for each cheliped and the scatterplot of the regression scores against the CL provides a graphical means to examine the cheliped shape change as unit of carapace length change (DRAKE and KLINGENBERG, 2008).

The allometric trajectories were compared by means of a permutational multivariate analysis of covariance (PERMANCOV, ANDERSON et al., 2004) with CL as covariate. In order to exclude any bias from the analysis (for examples see ANDERSON, 2001), only specimens within the same carapace length range (from 29.7 to 38.3 mm for *C. johni* and from 31.5 to 36 mm for *P. monodon*) were included in PERMANCOVA. Given the presence of groups, prior to PERMANCOVA we assessed the differences in slopes of allometric trajectories. In both species the CL by sex interaction term was analyzed throughout the comparison of multivariate regression slopes between groups with TpsRegr (ROHLF, 2010c). Briefly, the test for slopes compares the amount of variance explained by two models: one is simultaneously fitting group-specific multivariate linear regressions with each population having its own slope, while the second one is also fitting group-specific lines but it does so by forcing them to be parallel (see VISCOSI and CARDINI, 2011 for details). As the differences among slopes were found to be not significant (*p* > 0.05), all subsequent analysis were performed by using the same allometric trajectory for groups. Sexual dimorphism in cheliped size was assessed by permutational tests, whereas form (shape and size) differences were tested by applying a Discriminant Analysis (DA). The reliability of the discrimination of shape variables was calculated by leave-one-out cross-validation (e.g. LACHENBRUCH, 1967). DA was repeated on non-allometric shape variables (size-corrected) to evaluate the sexual dimorphism without the allometric effect. Size-corrected variables were obtained as reported in LEZCANO et al. (2012).

**RESULTS**

For *C. johni*, the first 2 principal components (PCs) of the PCA accounted for 75.4% of the total cheliped form variation (61% for PC1 and 14.4% for PC2; Fig. 2A). For *P. monodon*, PC1 accounted for 50.5% and PC2 for 20.6% (together they accounted for 71.1% of the total form variation; Fig. 2B). In both species, the examination of the wireframe representations of extreme forms on each axis indicates that PC1 mainly represents an inverse relationship between pollex (distal extension of the propodus) length and both manus (proximal part of the propodus) length and propodus height. Data points for females were associated with narrower cheliped, and males were related with high propodus, long manus and short pollex. Males of *C. johni* showed the widest variation in cheliped form and the fully arched morphology (i.e. with a relatively shorter pollex and a longer manus; Fig. 2A, B). The second axis represents the degree of bending of the cheliped in the vertical plane, with an inflection point at the base of the pollex.

The multivariate regression of cheliped shape on CL was significant (*p* < 0.05; permutation test with 10,000 random permutations) in both species (Fig. 2C, D), but CL accounted for a slightly larger amount of the total shape variation in *C. johni* than in *P. monodon* (13.4% and 9.2%, respectively). The cheliped shape change relative to CL was similar in both species: chelipeds change, on average from a long and narrow pollex with short manus to a relatively shorter and wider pollex with longer manus during growth (i.e. from small to large CL). The degree of cheliped arching was greater in males than in similar-sized females, and only *C. johni* males showed the fully arched
Figure 2. Plot of the principal components (PCs; upper) and of multivariate regression of cheliped shape onto carapace length (lower) of *Cervimunida johni* (A-C) and *Pleuroncodes monodon* (B-D). A-B: figures represent the displacement vectors (wireframe) from the overall mean shape (grey) to the positive and negative extreme shape (black) for PC1. Shape changes have been exaggerated (scale factor 0.1) for better visualization and percentages of explained variance for each axis are in parentheses. C-D: shape (Regression Score 1) as a function of carapace length, illustrating cheliped allometric growth of sexes. Wireframes represents the opposite extremes of allometric variation: the average cheliped shape and the change from the grey circles to the black dots indicates the predicted landmark shift corresponding to an increase and decrease of centroid size by 15 scale factor. Symbols: triangle for females; square for males.

Differences between sexes (intercept) were significant in both species (Fig. 2C, D) and the allometric trajectories were largely aligned with vectors of mean shape differences, with males of both species extending the cheliped trajectories to larger body sizes (Fig 2C, D).

Permutational tests confirmed that cheliped size (CS) differed significantly between sexes (*p* < 0.001; 10,000 random permutations) in both species. The results of the Discriminant Analysis showed that mean cheliped form differed between sexes (Fig. 3) and the accuracy of group discrimination was high in all sex-specific groups (Table 1). Because CS was only significantly related to cheliped shape in *C. johni*, size-corrected variables were calculated for sexes only in this species. The difference between size-corrected variables was not significant (*p* > 0.05) and the accuracy in the discrimination of groups strongly decreased (Table 1).

**DISCUSSION**

Here we present the first study comparing cheliped form variation of two species of squat lobsters of commercial importance which inhabit the continental margin of Chile: *Cervimunida johni* and *Pleuroncodes monodon*. Using the general approach of CLAVERIE and SMITH (2010), we confirmed that cheliped size and
shape of these species is related to known aspects of their reproductive biology, particularly the mating behavior.

In addition to duration of mate-guarding and resource finding and monopolization, *C. johni* and *P. monodon* also differ in terms of their mobility and intraspecific aggression (ESPINOZA-FUENZALIDA et al., 2012). *P. monodon* is much more active and aggressive than *C. johni*. Even more, pelagic phases have been reported from the genus *Pleuroncodes*, underlining their high mobility (BOYD, 1967; ROBINSON and GOMEZ-AGUIRRE 2004). Differences in aggregation patterns were also reported (AHUMADA et al., 2013) and the estimated average density of *P. monodon* was usually larger than that of *C. johni*. As the objective of this study was to examine the relationship between mate-guarding and cheliped form variation, the influence of all aforementioned factors (mobility, aggregation patterns, etc.) on cheliped form have not been taken into account, which should be done in future studies.

Cheliped form variation was similar in *P. monodon* and *C. johni*, but the degree of arching was greater in males, especially in those of *C. johni*. Similar wide variation in cheliped morphology (from straight to arched) was previously revealed in other squat lobsters, particularly in *Munida* spp. (CLAVERIE and SMITH 2007; 2010 and references therein). These authors suggested that this morphology was present in a common ancestor or the extant species had convergent morphological evolution due to similar selection pressures. The arched morphology is considered a sexually selected trait, adapted to inflict puncture wounds on opponents during agonistic interactions. Males with arched chelipeds are thought to be more successful in agonistic interactions and/or in obtaining and handling mates than those with straight chelipeds. Our results support the hypothesis reported by CLAVERIE and SMITH (2007), since males of *C. johni*, a species with extended mate-guarding, presents fully arched chelipeds that serve as a weapon during antagonistic encounters, while *P. monodon* with brief mate-guarding did not develop this morphology.

Asymmetry in chelipeds (cheliped laterality or handedness) of lobsters and other decapod crustaceans has been extensively reported (e.g., GOVIND, 1989; LEE, 1995; SCHENK and WAINWRIGHT, 2001; DENNENMOSER and CHRISTY, 2013). Asymmetry involves both, functional and structural differences among appendage sides and is mainly due to differences in feeding and locomotion, but also to sexual behavior (for examples see MARIAPPAN et al., 2000). In otherwise symmetric specimens, regeneration after autotomy could also cause asymmetry (CLAVERIE and SMITH, 2010). In *M. rugosa*, CLAVERIE and SMITH (2010) have found significant differences among cheliped sides and suggest that chelipeds play different roles in fighting and/or feeding. In our study, we found that while *P. monodon* showed neither cheliped shape nor size differences among
appendage sides, *C. johni* showed small (negligible compared with other levels of cheliped variation) but significant cheliped shape asymmetry. Until further studies in *C. johni* assess the strength of asymmetry, and whether it is the result of differences in function/structure or autotomy, we assume that chelipeds in both *C. johni* and *P. monodon* are symmetric among sides and play similar functions.

Even when the general patterns of within-sex cheliped shape variation was almost similar in both species (wide cheliped variation in males of *C. johni*; see above), sexual dimorphism was clearly different, particularly in the strength of differences and in the relation with both, carapace length and cheliped size. Sexual dimorphism in carapace length and in cheliped size was previously reported for both species (see CLAVERIE and SMITH, 2010 and references therein) and confirmed in our study, where males are larger than females and develop larger chelipeds. For both species, the proportion of specimens with arched chelipeds increased, on average, with increasing cheliped and body size. However, a considerable proportion of large males maintained straight chelipeds, especially in *C. johni*. Cheliped size was related to cheliped shape only in *C. johni*. Moreover, sexual dimorphism in cheliped shape is mainly due to differences in cheliped size; after correcting for size effects, no sexual dimorphism was found, similar as in some brachyuran crabs (LEE, 1995). The allometric patterns of cheliped shape variation (cheliped shape vs. carapace length) follow the same developmental pathways in both sexes, with males extending the cheliped trajectories to larger body sizes. Nonetheless, the allometric trajectories were largely aligned with vectors of mean shape differences, suggesting that chelipeds differ among sexes even in small specimens.

Sexual dimorphism in chela features (arched morphology) appears to be more pronounced in the species with long mate-guarding (*C. johni*) than in the species with short mate-guarding (*P. monodon*). These findings are in accordance with the conclusions by LEE (1995) who suggested that species with prolonged mate-guarding have stronger sexual dimorphism in chela size than species without or with short mate-guarding. This may result from strong sexual selection on chela size and shape in males that guard females for long time periods. During prolonged mate-guarding males may frequently have to fend off other males, generating advantages for males with large and powerful chelae. In the crab *Carcinus maenas*, where males guard premolt females for extensive time periods (BERRILL and ARSENAULT, 1982), males with larger chelae were significantly more successful in winning fights than males with small chelae (SNEDDON et al., 1997). In crayfish *Orconectes rusticus* males with larger claws were more successful in winning intrasexual encounters, and their matings with females lasted significantly longer than those of small claw males (SNEDDEN 1990). Thus, in species with prolonged mate-guarding sexual selection seems to drive intersexual differences in chela morphology.

In summary, the main results of our study is that cheliped form variation reflect the reported differences in both species: *C. johni* which is an extended mate-guarding species, presents a wide cheliped form variation (specifically in males), strong sexual dimorphism (highly related to cheliped size) and allometry with carapace length that match the morphology reported for *M. rugosa* more than *P. monodon*, a species with short mate-guarding.

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