

ARTICLE

Relative growth and reproductive parameters in a population of *Microphrys bicornutus* (Brachyura, Majoidea) from Bocas del Toro, Caribbean Sea, Panama

Crecimiento relativo y dinámica reproductiva de una población de *Microphrys bicornutus* (Brachyura, Majoidea) de Bocas del Toro, Mar Caribe, Panamá

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Resumen.- Se estudió la estructura de la población, el crecimiento relativo de los caracteres sexuales secundarios de ambos sexos y la fecundidad de las hembras de *Microphrys bicornutus* de Bocas del Toro, Panamá. Se recolectó manualmente y con red un total de 135 individuos. Las hembras ovígeras midieron 6,5-14,4 mm de ancho de caparazón (AC), las no ovígeras 4,2-16 mm, y los machos 5,9-17,4 mm. El crecimiento relativo del largo del caparazón *versus* el ancho de caparazón fue isométrico en los machos y alométrico negativo en las hembras. El crecimiento relativo del largo de la quela y el ancho del abdomen respecto del ancho de caparazón mostró la presencia de 2 fases, madura e inmadura, para machos y hembras, respectivamente. La talla en la cual el 50% de los machos y las hembras alcanzaron la madurez morfométrica fue 13,6 y 8,5 mm, respectivamente. El valor medio del output reproductivo fue de 3,6, la fecundidad relativa de 2041,91, y la talla relativa de madurez sexual fue de 0,4. Como la mayoría de los cangrejos Brachyura, *M. bicornutus* mostró una fuerte correlación positiva entre la talla de la hembra y la fecundidad. Una comparación entre los parámetros reproductivos descritos en el presente trabajo y los de las poblaciones de *M. bicornutus* en Venezuela, Jamaica y Brasil mostró algunas diferencias, por lo tanto, podría sugerirse que las mismas se deberían a efectos latitudinales o ambientales.

Palabras clave: Mar Caribe, fecundidad, crecimiento relativo, madurez sexual, cangrejo araña

Abstract.- Size structure within a population, relative growth of secondary sexual characters in both sexes, and fecundity of females were assessed from field samples of *Microphrys bicornutus* in the vicinity of Bocas del Toro, Caribbean Sea, Panama. The study was based upon 135 individuals captured by hand and a kick net. Carapace width ranged from 6.5 to 14.4 mm for ovigerous females, 4.2 to 16 mm for non-ovigerous females, and 5.9 to 17.4 mm for males. The relative growth of carapace length *versus* carapace width was isometric in males and negatively allometric in females. Relative growth of the chela length and abdomen width, compared to the carapace width, exhibited 2 phases, immature and mature, for males and females, respectively. The carapace widths at which 50% of males and females reached morphometric maturity were 13.6 and 8.5 mm, respectively. The mean value of the reproductive output was 3.6, the relative fecundity was 2041.91 eggs, and the relative size at the onset of sexual maturity was 0.4. As in most brachyuran species, the studied population of *M. bicornutus* showed a strong positive correlation between fecundity and body size. Differences in those parameters for previously studied populations from Venezuela, Jamaica and Brazil may be due to latitudinal or regional environmental effects.

Key words: Caribbean Sea, fecundity, relative growth, sexual maturity, spider crab

INTRODUCTION

Comparative studies of population size structure and morphometrics in decapods have become common in recent years, especially due to interest in their variability and its possible relationship to environmental adaptation (Mantelatto *et al.* 2010). Characteristics such as the size at morphological maturity and reproductive potential are considered powerful tools to access population differences.

Morphometric data such as those based upon relative size of the male chela and female abdomen have been used widely in studies of decapod crustacean relative growth to detect allometric changes reflecting sexual maturity (Hartnoll 1974). Fecundity of females, defined as the number of eggs per batch (clutch) produced by a female, is recognized as a strong indicator of reproductive potential. Both morphological maturity and fecundity of brachyurans from many different habitats and climatic regions have been reported previously (*e.g.*, Hartnoll 1985, Hines 1988, Mantelatto & Fransozo 1997, Moura & Coelho 2001, Litulo 2004), and many differences have been reported among conspecific populations. Such differences are usually related to latitudinal temperature gradients (Clarke *et al.* 1991, Lardies & Castilla 2001, Wehrmann & López 2003, Mantelatto *et al.* 2010), inter-annual temperature variation (Simons & Jones 1981, Ouellet & Plante 2004, Brillon *et al.* 2005), and regional variation in salinity, food resources, productivity (Bas *et al.* 2007) or other environmental factors (Mashiko 1990, Hancock *et al.* 1998, Carmona-Suárez 1992, 2003, 2007).

Microphrys bicornutus (Latreille, 1825) is a spider crab (Majoidea *sensu* Ng *et al.* 2008) living on coralline or other shallow subtidal rubble, grass flats, and red mangrove roots (Lemaitre 1981, Keith 1985). The carapace often carries a dense decoration of sponges, anemones, hydroids and algae for concealment (Ghetty & Hazlett 1978, Williams 1984). It is distributed from North Carolina and Bermuda to Florida, the Gulf of Mexico, Central America, the Antilles, Colombia, Venezuela, and Brazil (Melo 1996, Felder *et al.* 2009). Population characteristics and ecological features of *M. bicornutus* were studied previously in Venezuela (Buchuaco and Isla Margarita; Carmona-Suárez 1992, 2007, 2013 and López-Greco *et al.* 2000, respectively), Jamaica (Hartnoll 1965), Grand Cayman Island (Gore *et al.* 1882), and Brazil (Batista *et al.* 2009). In the course of these projects investigating inter-population size variations, relative growth, larval histories, and reproductive attributes such as fecundity, differences were reported among populations.

Here we report on a population from Bocas del Toro, on the Caribbean coast of Panama. Despite much of this region holding the status of a preserved area, anthropogenic activities (urbanization and tourism-related) in the archipelago are increasingly creating an urgent need for information on aquatic biodiversity (Torati *et al.* 2011). In particular, baseline data such as species lists and insights on reproductive biology are needed for future management and preservation of the region's marine biodiversity.

Given that *Microphrys bicornutus* has a wide distributional range, it represents an excellent candidate for comparative studies of regional adaptations. We hypothesize that there are differences in the morphometric and reproductive parameters among populations of the same species as regional adaptations. To facilitate latitudinal comparisons, we herewith investigate: [1] population structure, [2] relative growth of secondary sexual characters of both sexes, and [3] fecundity of females for a population from Bocas del Toro, Panama. Our findings are intended to provide a baseline for studies of inter-population variability in a species with a wide geographic range.

MATERIALS AND METHODS

Crabs were collected by hand from an area encompassing shoreline roots of mangroves (*Rizophora*) and *Thalassia* seagrass meadows near the Smithsonian Tropical Research Institute (STRI) marine station (09°20'N, 82°14'W) at Bocas del Toro on the Caribbean coast of Panama in August 2011. This region is located along the northern-most Atlantic side of Panama and comprises large and small islands that provide a number of micro-habitats among mangrove roots, corals, sponges, rubble, and other substrates.

Animals clinging to shoreline mangrove roots were collected by hand while lifting and carefully inspecting roots and associated epifauna, and those from the adjacent grass bed were taken by hand-pushing of a pole-mounted 22.5 x 45 cm fine meshed (1 mm) Wildco® kick net through the grass. This concentrated effort of sampling conducted over a full week (day and night) allowed the capture of diverse sizes of small crabs and full representation of the local population. Due to the limited number of individuals, samples were pooled and no differences between separately sampled biotopes were considered for the data analysis. Individuals were sacrificed by placement into a -15°C freezer for

approximately 15 min. Sex and the presence of ovigerous females were recorded. Sexes were differentiated depending on the form of the abdomen following the criteria used by Hernández-Reyes *et al.* (2001) for *Mithrax forceps*. All measurements were made with digital calipers ± 0.01 mm. In both sexes, the carapace length (CL) and width (CW, used as the independent variable or reference measurement in all analyses) were measured. Maximum abdomen width was measured for females (AW, at a level of the fourth abdominal segment) and, chela length (ChL) was measured for males. Quantitative relationships between morphometric data were described by least square regressions (CL \times CW for both sexes, AW \times CW for females, and ChL \times CW for males). Allometric growth of all dimensions relative to CW was estimated using the allometric equation $Y = aX^b$ (Huxley 1950). These relationships were log-linearized (base 10): $\log_{10} \text{ChL}/\text{AW} = \log_{10} a + b \log_{10} \text{CW}$, where b is the regression slope value or allometric coefficient. Allometric coefficients between 0.9 and 1.1 were considered indicative of isometry, lower than 0.9 indicative of negative allometry, and above 1.1 indicative of positive allometry (= 'conservative definition' of Clayton 1990). Slope values were compared to the expected isometric values using Student's t -test, at a significance level of $\alpha = 0.05$. Slopes and intercepts of different regression lines were compared by ANCOVA covariance analysis (Zar 1999).

External morphological changes mark the end of all the processes that determine maturity in decapods, underpinning criteria commonly used to classify crabs as mature or immature (Somerton 1980, Conan & Comeau 1986). Estimates of the size at 50% maturity at the population level (CW_{50}) are most frequently used to define the reproductive state in decapods, this being based upon the classification of crabs in a sample into either mature or immature and subsequent estimation of the body size in which 50% of the specimens are mature. This approach assumes that there is a sigmoidal relation between size and the percentage of mature crabs (Roa *et al.* 1999). The CW_{50} may be interpreted as the size at which a randomly chosen specimen has a 50% chance of being mature (Somerton 1980). Thus, the relative size at which females and males reach morphometric maturity (CW at which 50% of females and males are morphometrically mature, CW_{50}) was calculated using the logistic equation. The logistic function used was $y = 1/(1 + e^{-r(\text{CW} - \text{CW}_{50})})$, where CW_{50} corresponds to the size at which 50% of the individuals are considered mature and r stands for the slope of the curve (*e.g.*, Corgos & Freire 2006, Sal-Moyano *et al.* 2011).

Ovigerous females were classified according to the stage of development of the eggs, following Mantelatto & Garcia (1999): Initial Stage - eyes of larvae not visible, eggs mostly filled with yolk and light-red colored; Intermediate Stage - eyes become visible, beginning pigmentation and segmentation of the larvae; Final Stage - zoea becomes visible, dark-brown eggs. Since all females captured had recently deposited eggs and thus belonged to initial stage, they were used for the fecundity analysis. The eggs were carefully removed from the pleopods and counted under a light stereomicroscope. The average volume (mm^3) of each egg was calculated according to Jones & Simons (1983) from the formula $1/6 \pi I^3$, where I is the mean of 2 egg diameters (including the chorionic membrane tightly adhering to the embryonic surface) for a subsample 15 eggs per female. The minimum and maximum diameters of each egg were measured under a light stereomicroscope. Wet weights of egg masses at all three stages were determined $\pm 1 \mu\text{g}$ using an analytical balance. Eggs from the female were washed in distilled water, after which water was removed by contact with filter paper. Egg masses were first dried until a constant dry weight was reached (50°C for 24 h) and then burned for 4 h at 500°C to obtain ash-free weights. Organic material was estimated by subtracting combusted material weight from dry weight. The female's body was treated similarly to their egg mass except that they were both dried and burned as above for 48 and 12 h, respectively (Lardies & Wehrtmann 1996). The following measurements were taken for each female: body wet weight (BWW), body dry weight (BDW), egg wet weight (EWW), egg dry weight (EDW) and number of eggs (NE). For each female, reproductive output (RO) was calculated as $\text{EDW}/\text{BDW} \times 100$ (Clarke *et al.* 1991), the relative fecundity (RF) was calculated as NE/BDW , and the relative size at the onset of sexual maturity (RSOM, Charnov 1990) was calculated as minimum ovigerous female size/maximum female size. A correlation between the number of eggs and the CW was performed. Voucher specimens were deposited in the Crustacean Collection of the Department of Biology (CCDB), Faculty of Philosophy, Sciences and Letters of Ribeirão Preto, University of São Paulo (FFCLRP/USP), Brazil and the Department of Biology (ULLZ), University of Louisiana at Lafayette, Lafayette, U.S.A., under catalog numbers CCDB 3557, 3564 and ULLZ 13376-13378, 13662.

RESULTS

A total of 135 individuals were captured, including 54 (40%) non-ovigerous females, 18 (13.33%) ovigerous females, and 63 (46.66%) males. Non-ovigerous females ranged from 4.2 to 16 mm CW (9.08 ± 2.3 mm CW), ovigerous females ranged from 6.5 to 14.4 mm CW (9.93 ± 2.19 mm CW), and males ranged from 5.9 to 17.4 mm CW (10.8 ± 3.02 mm CW) (Fig. 1). Morphometrically, the relative growth of the CL *versus* the CW was isometric in males ($\log CL = 0.12 + 0.99 \log CW$, $R^2 = 0.99$, $b = 0.99$, $t = 153.7$, $P < 0.0001$) and negatively allometric in females ($\log CL = 0.42 + 0.71 \log CW$, $R^2 = 0.1$, $b = 0.71$, $t = 2.93$, $P < 0.01$).

In males, the relative growth of ChL with respect to CW differed markedly between immature and mature ones, the immature exhibiting a positively allometric phase ($\log ChL = 0.09 + 1.11 \log CW$, $R^2 = 0.88$, $b = 1.11$, $t = 18.3$, $P < 0.0001$), and the mature, showing a negatively allometric phase ($\log ChL = 0.48 + 0.68 \log CW$, $R^2 = 0.83$, $b = 0.68$, $t = 8.7$, $P < 0.0001$) (Fig. 2). Slopes of the regression lines over these two phases differed significantly (ANCOVA, $F = 9.3$, $P < 0.01$). The presence of an inflection point between these phases suggested a pubertal molt.

The analysis of the logistic equation showed that the size at which 50% of males reached morphometric maturity was $CW_{50} = 13.6$ mm. The smallest mature male measured 12.6 mm CW, and the largest immature male measured 13.8 mm CW.

In females, relative growth analysis showed that the growth of AW with respect to CW was negatively allometric in both immature ($\log AW = 0.58 + 0.62 \log CW$, $R^2 = 0.51$, $b = 0.62$, $t = 5.12$, $P < 0.001$) and mature females ($\log AW = 0.34 + 0.81 \log CW$, $R^2 = 0.77$, $b = 0.81$, $t = 11.6$, $P < 0.0001$) (Fig. 3). Although the slopes of regressions in this relationship did not differ between the female growth phases, differences in y-intercepts were significant (ANCOVA, $F = 323.7$, $P < 0.001$). The analysis of the logistic equation showed that the size at which 50% of females reached morphometric maturity was $CW_{50} = 8.5$ mm. The smallest mature female measured 6.5 mm CW, and the largest immature female 12.3 mm CW. In addition, the smallest size ovigerous female captured was 6.5 mm CW. A wide overlap between CW of large immature females and CW of those in the mature phase was observed; indicating females of the same carapace width can have undifferentiated or differentiated (mature) abdomens.

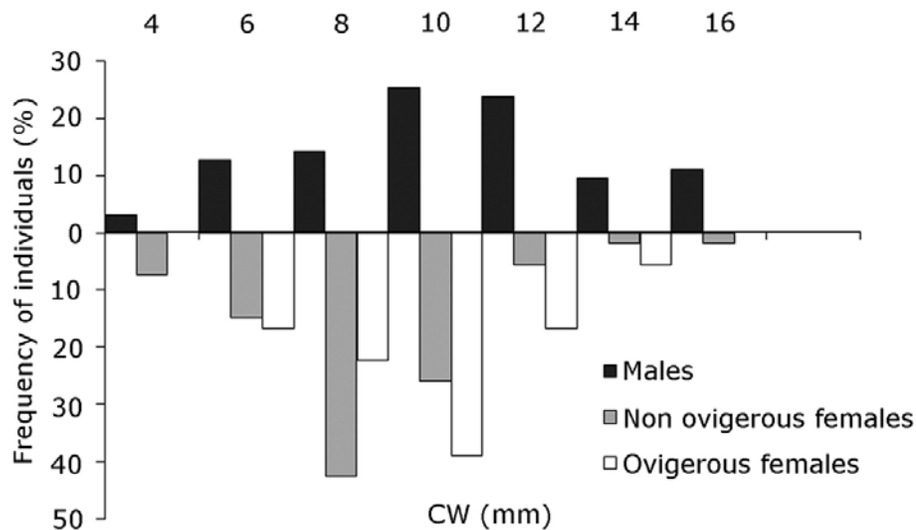


Figure 1. *Microphrys bicornutus*. Size frequency distributions of ovigerous females, non-ovigerous females and males. Numbers in the upper part of the graph represent the size ranges / *Microphrys bicornutus*. Distribución de frecuencia de talla de hembras ovígeras, no ovígeras y machos. Los números en la parte superior del gráfico representan los rangos de talla

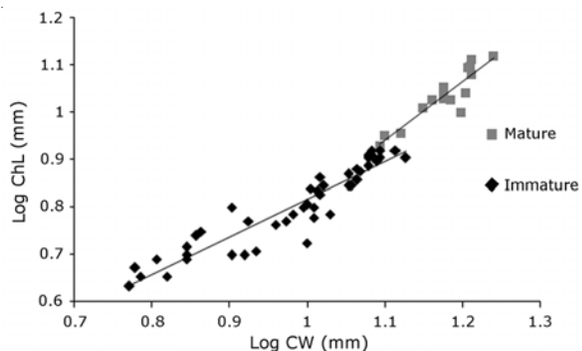


Figure 2. *Microphrys bicornutus*. Relationships between chela length (ChL) and carapace width (CW) of males; the regression lines correspond to immature and mature males / *Microphrys bicornutus*. Relaciones entre el largo de la quela (ChL) y el ancho de caparazón (CW) de los machos; las líneas de regresión corresponden a los machos inmaduros y maduros

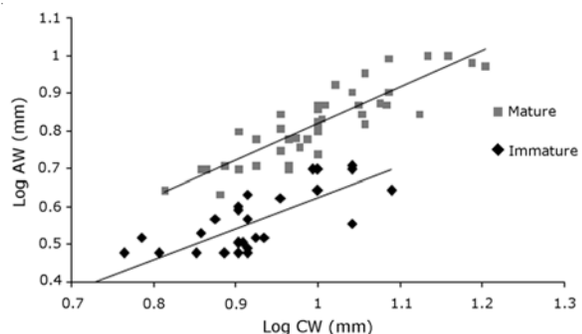


Figure 3. *Microphrys bicornutus*. Relationships between abdomen width (AW) and carapace width (CW) of females; the regression lines correspond to immature and mature females / *Microphrys bicornutus*. Relaciones entre el ancho de abdomen (AW) y el ancho de caparazón (CW) de las hembras, las líneas de regresión corresponden a las hembras inmaduras y maduras

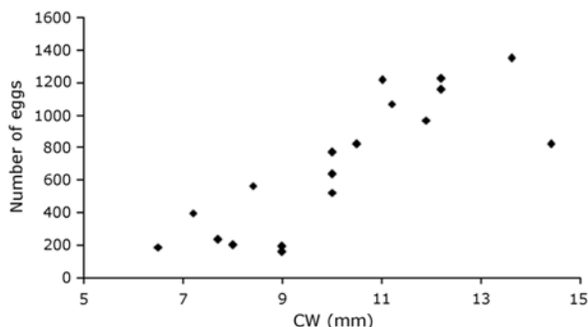


Figure 4. *Microphrys bicornutus*. Correlation between the number of eggs per batch and the carapace width (CW) in females / *Microphrys bicornutus*. Correlación entre el número de huevos por puesta y el ancho de caparazón (CW) de las hembras

Table 1. *Microphrys bicornutus*. Size (carapace width, CW) and weight of ovigerous females. n = number of females, sd = standard deviation / *Microphrys bicornutus*. Talla (ancho de caparazón, CW) y peso de las hembras ovígeras. n = número de hembras, sd = desviación estándar

n = 18	Mean value	sd	Minimum	Maximum
CW (mm)	9.930	2.190	6.50	14.400
Dry weight (g)	0.2118	0.1363	0.059	0.5495
Water content (g)	0.2754	0.1826	0.1067	0.7224
Organic matter (g)	0.094	0.053	0.031	0.205
Ash content (g)	0.117	0.085	0.028	0.345

Mean values of size and weight of ovigerous females were 9.93 mm CW and 0.2118 g dry weight, 0.2754 g water content, 0.094 g organic matter and 0.117 g ash content. The standard error, minimum and maximum values of those variables are shown in Table 1. The mean value/standard error for the egg volume of stage 1 was 0.0253/0.0113 mm³. The value of the RO was 3.6/2.02, while the RF was 2041.9/1291.7. The RSOM was 0.4 (6.5/16). A significant positive correlation was found between clutch size (number of eggs) in ovigerous females and their CW ($r = 0.68$, d.f. = 16, $P < 0.001$, $n = 18$, Fig. 4).

DISCUSSION

In Bocas del Toro, Panama, the majoid crab *Microphrys bicornutus* reproduces continuously throughout the year, as it does over a large part of its range (Gore *et al.* 1982). Thus, the present report must be regarded as a restricted initial evaluation of reproductive dynamics in this population. Although the number of ovigerous females captured in the present study is low ($n = 18$) we consider the results important since they constitute the first reproductive study for this common species in Panamanian waters.

The size frequency distribution reported here showed smaller ranges of female and male sizes than those reported for same species in Buchuaco, which presented ranging from 1.9 to 35.4 mm CL (Carmona-Suárez 2007), and in Isla Margarita, ranging from 4.1 to 33.2 mm CL (López-Greco *et al.* 2000). Hartnoll (1965) also reported that the population from Jamaica (17°43'N, 69°40'W) consisted of larger sizes, ranging from 11 to 33 mm of carapace length (extracted from Carmona-Suárez 1992). However, in Brazil, smaller size ranges compared to the ones recorded for Bocas del Toro were found for both sexes, reaching a maximum size of 11 mm (Batista *et al.* 2009). Size differences for populations of the same species have been most commonly attributed

to the variation in their latitudinal distribution (*e.g.*, Hines 1989, Conde & Diaz 1992). Moreover, size differences could also be related to environmental instability with periodic stress in shallow waters at the site [heavy rainfall, terrestrial runoff, and periodic 'natural eutrophication' as reported at Bocas del Toro (Carruthers *et al.* 2005, D'Croz *et al.* 2005, Guzman *et al.* 2005)]. These could favor extinctions of older, larger cohorts, or alter nutritional resources for growth and cause stunting. Specifically in the case of Brazil populations, the smaller size ranges corresponded to immature males and females, and since *M. bicornutus* lives on the phytal *Halimeda opuntia*, it was suggested that the complex microhabitat provided by the algae may favor the recruitment of immature crabs (Batista *et al.* 2009).

In males of several majoid species, relationships between the carapace width and the size of the chelae characterize immature and mature phases, separated by a pubertal molt (Watson 1970, Conan & Comeau 1986, Schejter & Spivak 2005, Barón *et al.* 2009, Sal Moyano *et al.* 2011). In other cases, they provide evidence of three phases, those being usually labeled as juveniles, adolescents and adults, separated by a pre-pubertal and a pubertal (or terminal) molt respectively (Sainte-Marie *et al.* 1995, Sampedro *et al.* 1999, Carmona-Suárez 2003). Here, we report 2 phases, showing the immature phase to end in a discontinuity marking a pubertal molt. These 2 phases were described for the same species from Venezuela, in Buchuaco (Carmona-Suárez 2007) and Isla Margarita (López-Greco *et al.* 2000), although in the former a small overlap between phases could be distinguished. The estimated size of 50% morphometric maturity calculated in the present report (13.6 mm CW or 17.4 mm CL) was 13.4% smaller (a 2.5 mm difference) than the 19.9 mm CL calculated by Carmona-Suarez (2007) for the Buchuaco population. However, this value was similar to the one (16.5 mm CL) reported by López-Greco *et al.* (2000) for the Isla Margarita population. Hence, it is suggested that males from Bocas del Toro and Isla Margarita attain morphometric maturity earlier than males from Buchuaco, showing that the environmental conditions may be acting in a different way in terms of supply for the growth and attainment of maturity in both populations. However, since males from Buchuaco apparently reached larger sizes than those from either Isla Margarita or Bocas del Toro, the acquisition of morphometric maturity at a larger size could also simply suggest that male sizes vary among populations.

In females, similar to other majoid species, the presence of 2 phases was recognized, with separation by a puberty molt (Sainte-Marie & Hazel 1992, Carmona-Suárez 2003, Schejter & Spivak 2005, Barón *et al.* 2009, Sal Moyano *et al.* 2011). However, in our study, *M. bicornutus* showed a wide size overlap between immature and mature female phases, similar to that described by Carmona-Suárez (2007) and López-Greco *et al.* (2000) for the same species from Venezuela. Although the number of individuals used in the present work to calculate morphometric maturity was less than in the aforementioned studies, the estimated size of 50% morphometric maturity calculated in the present study (8.5 mm CW or 11.7 mm CL) was similar to the one (12 and 13.6 mm CL) calculated by Carmona-Suárez (2007) and López-Greco *et al.* (2000). Hence, minimal size difference at morphometric maturity was found between the Bocas del Toro population and those from Buchuaco and Isla Margarita, which likely reflects similar environmental conditions for the growth of these decapods in the Caribbean region. It is noteworthy that males differ in size at morphometric maturity between the Buchuaco and the Bocas del Toro populations, while females do not. It may indicate that females are more conservative in terms of body size, while males are more plastic in responding to varied environmental conditions.

Studies reporting reproductive parameters such as fecundity and/or reproductive output in majoids are scarce considering diversity of this group (Table 2). The mean value of the RF recorded in the present study was similar to that reported for the same species in Venezuela by López-Greco *et al.* (2000) (2041.9 eggs in present study *versus* 2391.5), although the mean value of the RO in the present study was less than half the previous report (3.6 in present study *versus* 8.9). This difference may be caused because of the smaller mean value of egg diameters in stage 1 reported here as compared to that recorded in the aforementioned study. By contrast, in the Buchuaco population, a lower number of eggs was reported (mean = 1067.4), although egg size was larger. Thus, this population may produce less eggs but with higher quantities of yolk (thus, larger size), and thereby yield embryos and larvae optimized to local environmental conditions (Carmona-Suárez 2013). It is known that the RO can vary for a decapod species between Decapod species between different latitudes (Terossi *et al.* 2010a, b). Similarly, *M. bicornutus* may be responding by having smaller and more numerous eggs in the Bocas del Toro population due to the stress that characterizes this habitat where

Table 2. Studies conducted on American Majoidea reporting fecundity (mean or maximum and minimum number off eggs) and/or reproductive output (RO). CW = carapace width, CL= carapace length of ovigerous females / Estudios realizados en Majoidea de América reportando la fecundidad (media o máximo y mínimo del número de huevos) y/o el output reproductivo (RO). CW = ancho de caparazón, CL = largo de caparazón de hembras ovígeras

Species	Study site	Fecundity	RO	Size	References
<i>Collodes tenuirostris</i>	Mazatlán, México	1302	-	4.9-19.2 mm CW	García- Guerrero & Hendrickx (2004)
<i>Euprognatha bifida</i>	Mazatlán, México	83	-	3.4-8.8 mm CW	García- Guerrero & Hendrickx (2004)
<i>Leurocyclus tuberculosus</i>	Rio de Janeiro, Brazil	10327.3	-	15.3-55.4 mm CW	Stauffer <i>et al.</i> (2011)
<i>Libidoclaea granaria</i>	Atlantic Ocean, Argentina	1200-9300	-	16-48 mm CW	Schejter & Spivak (2005)
<i>Microphrys bicornutus</i>	Isla Margarita, Venezuela	2391.5	8.90	11.2-22.2 mm CL	López- Greco <i>et al.</i> (2000)
<i>Microphrys bicornutus</i>	Buchuaco, Venezuela	1067.4	-	7-23.6 mm CL	Carmona- Suárez (2013)
<i>Microphrys bicornutus</i>	Bocas del Toro, Panamá	2041.9	3.60	6.5-14.4 mm CW	Present study
<i>Mithraculus forceps</i>	São Paulo, Brazil	402.8	-	9.4-14.0 mm CW	Cobo & Okamori (2008)
<i>Mithrax forceps</i>	Isla de Margarita, Venezuela	34-4777	-	11.2-30.3 mm CL	Hernández- Reyes <i>et al.</i> (2001)
<i>Paradasygius depressus</i>	Mazatlán, México	1528	-	13.0-31.6 mm CW	García- Guerrero & Hendrickx (2004)
<i>Pyromaia tuberculata</i>	Mazatlán, México	910	-	3.8-19.0 mm CW	García- Guerrero & Hendrickx (2004)
<i>Stenorhynchus debilis</i>	Mazatlán, México	1470	-	4.8-13.1 mm CW	García- Guerrero & Hendrickx (2004)
<i>Stenorhynchus seticornis</i>	São Paulo, Brazil	621.1	-	6.0-14.0 mm CW	Okamori & Cobo (2003)
<i>Eurypodius latreillii</i>	Straits of Magellan, Chile	1627-9384	0.13	29-62.9 mm CL	Navarrete <i>et al.</i> (1999)
<i>Epialtus bituberculatus</i>	Grande and Domingas Dias, Brazil	149	-	4.8-7.9 mm CW	Cobo & Barros (2009)
<i>Chionoecetes opilio</i>	Bonne Bay, Canada	54000 (max)	-	37-90 mm CW	Comeau <i>et al.</i> (1999)

heavy terrestrial runoff and turbidity surges can be frequent. Moreover, it was proposed that for the majoid crab, *Mithraculus forceps* (A. Milne-Edwards, 1875), certain environmental conditions, such as the availability of food for adults, may have some effect on intra-specific differences related to the values of the RO of this crab (Cobo & Okamori 2008). In the case of the RSOM, a similar value was found to that reported by López-Greco *et al.* (2000), also coinciding with values earlier reported for several other brachyurans, particularly majoid crabs (Hines 1982). The positive correlation between the CW and the number of eggs reported here for *Microphrys bicornutus* is a trend commonly evident in most brachyurans (*e.g.*, Hines 1982, García-Guerrero & Hendrickx 2004).

The present study, in combination with comparative data available in the literature, shows a relatively variable pattern in fecundity among majoids. Reproductive adaptation may depend on divergent strategies in terms of reproductive performance and output, these perhaps being in turn related to variable patterns in population structures, sizes at maturation, seasonality, sex ratios and other factors bearing on reproduction. Thus, more thorough study of these topics for related species and regional populations of majoids is required, both for appreciation of phylogenetic diversity and understanding of environmental constraints that likely bear on majoid reproductive adaptation.

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