Meat yield of Bolinus brandaris (Gastropoda: **Muricidae):** Comparative assessment of the influence of sex, size and reproductive status

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Summary: The present study assessed the influence of sex, size and reproductive status on the meat yield (soft tissues proportion) of the purple dye murex (*Bolinus brandaris*) from the Ria Formosa lagoon (southern Portugal). During one year of monthly sampling (October 2008 - September 2009), average meat yield of *B. brandaris* was 40.5±6.1% (range: 25.8-56.1%) wet weight), with no significant differences between sexes. Relationships established between specimen size and soft parts weight indicated that both shell length and total weight are excellent indicators of meat yield. Significant differences in meat yield between size classes further reinforced the trend of increasing meat yield during ontogeny. Meat yield exhibited significant monthly variation and a similar temporal trend in both sexes, which were directly related to the reproductive status. Meat yield of B. brandaris was compared with that of other muricid species and the marked influence of the reproductive status on meat yield prompted a comparative assessment of the spawning season and peak of three sympatric muricids (*B. brandaris*, *Hexaplex trunculus* and *Stramonita haemastoma*). Overall, these findings have implications at diverse levels, including the management, regulation and inspection of this fishing/harvesting activity and the commercialization and consumption of this seafood product.

Keywords: purple dye murex; Bolinus brandaris; meat yield; sex; size; reproductive status; spawning season and peak.

Proporción de tejidos blandos de Bolinus brandaris (Gastropoda: Muricidae): Evaluación comparativa de la influencia del sexo, la talla y el estado reproductivo

Resumen: El presente estudio evaluó la influencia del sexo, la talla y el estado reproductivo en la proporción de tejidos blandos de la cañaílla (Bolinus brandaris) de la laguna Ria Formosa (sur de Portugal). Durante un año de muestreo mensual (octubre de 2008 - septiembre de 2009), la proporción global de B. brandaris fue de 40.5±6.1% (rango: 25.8-56.1% peso húmedo), sin diferencias significativas entre sexos. Las relaciones establecidas entre la talla de los ejemplares y el peso de las partes blandas indicaron que tanto la longitud de la concha como el peso total son excelentes indicadores de la proporción de tejidos blandos. Las diferencias significativas entre clases de talla reforzaron aún más la tendencia creciente durante la ontogenia. La proporción de tejidos blandos mostró una variación mensual significativa y una tendencia temporal similar en ambos sexos, que estaban directamente relacionadas con el estado reproductivo. La proporción de tejidos blandos de B. brandaris se comparó con otras especies de murícidos y la marcada influencia del estado reproductivo impulsó una evaluación comparativa de la época y pico de desove de tres murícidos simpátricos (B. brandaris, Hexaplex trunculus y Stramonita haemastoma). En general, estos hallazgos tienen implicaciones a diversos niveles, incluyendo la gestión, regulación e inspección de esta actividad de pesca/marisqueo y la comercialización y consumo de este marisco.

Palabras clave: cañaílla; Bolinus brandaris; proporción de tejidos blandos; sexo; talla; estado reproductivo; época y pico de desove.

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INTRODUCTION

The purple dye murex, Bolinus brandaris (Linnaeus, 1758), is a common muricid species that is extensively dispersed throughout the Mediterranean Sea, whereas its distributional range along the adjacent Atlantic Ocean is restricted mainly to the coasts of Portugal and Morocco (Poppe and Goto 1991, Macedo et al. 1999, Houart 2001). Probably as a consequence of accidental introduction of juveniles, B. brandaris has extended its distributional range northwards, and has already been recorded in Galicia (northwestern Spain) (Bañón et al. 2008). This species usually occurs in shallow waters of the sub-littoral zone (Dalla Via and Tappeiner 1981), but can also be found at 100 m (Muzavor and Morenito 1999), 150 m (Houart 2001) or even 200 m depth (Macedo et al. 1999, Malaquias 2007), inhabiting sandy, sandy-muddy and muddy bottoms (Macedo et al. 1999, Muzavor and Morenito 1999, Malaquias 2007).

The purple dye murex was harvested during the Roman Empire using baited wicker baskets (Spanier and Karmon 1987) in order to extract the purple dye that was greatly prized in the ancient Mediterranean (e.g. Reese 1980, 2010, Oliver 2015). Today, B. brandaris is fished for human consumption using diverse types of artisanal fishing gears (Poppe and Goto 1991, Houart 2001), mainly in Portugal (Vasconcelos et al. 2008a) and Spain (Martín et al. 1995, Tirado et al. 2002, Mallol et al. 2004), where this species is an important gastronomic resource and is highly valued in local seafood markets (Ramón et al. 2005, Vasconcelos et al. 2008a), but also in France (Bartolome 1985), Italy (Ramón and Amor 2001, Cecalupo et al. 2006) and Greece (Katsanevakis et al. 2011), and occasionally in Turkey (Ramón and Flos 2001) and Tunisia (Elhasni et al. 2013).

Knowledge on meat yield (weight and proportion of the raw edible content) of seafood products is important, both as general information for the consumers and as specific data for the seafood processing and marketing industry. However, despite its importance and the existence of specialized literature on seafood worldwide (e.g. FAO 1989), information on the meat yield of shellfish and specifically of gastropod species is still relatively scarce and restricted to a few commercial species (either fishery-exploited or aquacultureproduced species) (see compilation by Vasconcelos et al. 2009). In Portugal, the purple dye murex is fished along the Algarve coast (southern Portugal), mainly in the Ria Formosa lagoon. It is a greatly appreciated seafood with high demand and commercial value in local seafood markets, reaching prices of around 20-25 € kg⁻¹ for first sale (Vasconcelos et al. 2008a). In addition, the commercial value of *B. brandaris* has prompted a few studies to assess its potential as a candidate species for molluscan aquaculture in both Spain (Ramón and Flos 2001, Vela and Moreno 2004, Ramón et al. 2005) and Portugal (Vasconcelos et al. 2012a).

The present study reports the estimation of meat yield and analyses the influence of factors such as sex, size and reproductive status on the meat yield of *B*.

brandaris from the Ria Formosa lagoon (southern Portugal). Besides improving the overall knowledge on the species biology, this kind of information is important to increase the awareness on the need to sustainably harvest *B. brandaris*, as well as to support a responsible commercialization and consumption of this locally important shellfish resource.

MATERIALS AND METHODS

Study area and sampling

Commercial samples (\approx 50 individuals per month) of purple dye murex (*B. brandaris*) were purchased from a local shellfish supplier during a one-year study period (October 2008 - September 2009). Individuals were caught in the Ria Formosa lagoon (Algarve coast, southern Portugal) (Fig. 1), using an artisanal fishing gear baited with cockles (*Cerastoderma edule*), locally known as "wallet-line" (Vasconcelos et al. 2008a), which is a non-size-selective fishing gear and thus collects individuals with broad size and weight ranges.

Because fishing operations and handling procedures frequently damage *B. brandaris* shells, in order to avoid biased measurements and weight discrepancies, specimens with damaged shells (e.g. injured shell aperture and/or broken siphonal canal) were discarded for the purpose of the present analyses. In the laboratory, a few colonizing algae and/or encrusting epibionts were removed from the shells with a hard brush; then the shells were drained to remove excessive water retained inside the mantle cavity (sealed by the operculum) and blotted dry onto absorbent paper.

Estimation of raw meat yield

Live *B. brandaris* were measured for shell length (SL) using a digital calliper (precision of 0.01 mm) and weighed for total weight (TW) on a top-loading digital balance (precision of 0.01 g). Subsequently, individuals were de-shelled in a bench vice in order to remove the soft parts of the organism (edible content=foot and visceral mass) together with the attached operculum (Fig. 2), which were also drained and blotted dry onto absorbent paper to eliminate extra-visceral water.

Because *B. brandaris* lacks external sexual dimorphism, individuals were sexed after breaking the shells, removing the soft parts of the organism and exposing

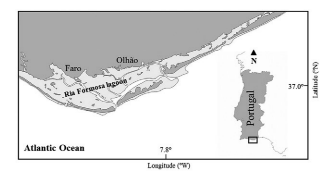


Fig. 1. – Map of the Ria Formosa lagoon (Algarve coast, southern Portugal).

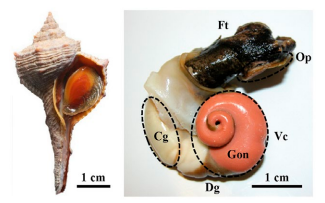


Fig. 2. – Purple dye murex (*Bolinus brandaris*): live specimen and soft parts of the organism (edible content) after shell breakage. Abbreviations: Cg, Capsule gland; Dg, Digestive gland; Ft, foot; Gon, gonad; Op, operculum; Vc, visceral coil (digestive gland and gonad complex).

the mantle cavity to observe the sexual organs in both sexes. Since *B. brandaris* is highly sensitive and severely affected by the imposex phenomenon in the Ria Formosa lagoon (Vasconcelos et al. 2010, 2011), which further complicates sexual identification, males were identified by the presence of penis and lack of capsule gland, while females were identified by the presence of vagina and capsule gland.

After removing the operculum, the whole meat (soft parts of the organism, i.e. foot and visceral mass) constitutes the edible portion of *B. brandaris* as it is consumed locally, so this assemblage of tissues was weighed for soft parts weight (SpW) on the top-loading digital balance (precision of 0.01 g). The raw meat yield (MY) was calculated as the proportion of the SpW relative to TW (MY=SpW/TW), with results expressed as mean±SD percentage of wet weight (i.e. g/100 g ww) (Vasconcelos et al. 2009).

Data treatment and statistical analysis

The relationships between specimen SL, TW and the respective SpW were assessed through regression analyses (least squares method), by fitting the power function to raw data (Y=aX^b) and assessing the degree of association between variables by the correlation coefficient (r). The relative growth between variables (isometry vs. allometry) was analysed through the allometry coefficient (regression slope - b) of the morphometric relationships. In relationships between the same type of variables (both ponderal, such as TW vs. SpW) isometry occurs for b=1, whereas in relationships between different types of variables (linear and ponderal, such as SL vs. TW and SL vs. SpW) isometry occurs for b=3, meaning that growth rates of both variables are identical throughout ontogeny (Huxley and Teissier 1936). Subsequently, a t-test (H_0 : b=1 or 3; H_A : b≠1 or 3) (Sokal and Rohlf 1987) was applied to confirm whether the slopes (b) of those morphometric relationships were isometric (b=1 or 3) or included in the allometric ranges (negative allometry: b<1 or 3; positive allometry: b>1 or 3) (Huxley and Teissier 1936). Finally, in order to assess possible differences in relative growth between sexes, the slopes (b) of the morphometric relationships of males and females were compared using a specific *t*-test for this purpose (H₀: $b_M=b_F$; H_A: $b_M\neq b_F$) (Zar 1996).

The sex ratio (M:F) of the overall samples was compared with parity (1:1) using the chi-square test $(\chi^2 \text{ test})$. In order to assess the influence of specimen size on the respective MY, individuals of both sexes were grouped into five size classes (<60, 60-70, 70-80, 80-90 and \geq 90 mm SL). In addition, individuals were further grouped into two broader size classes (<65 mm and ≥ 65 mm SL) defined according to the minimum landing size (MLS=65 mm SL) legally established for the catches of *B. brandaris* in the Ria Formosa lagoon (D.R. 1990, 2001). The influence of the reproductive status on MY was assessed using information previously obtained through histological techniques and the calculation of diverse reproductive condition indices of *B. brandaris* from the Ria Formosa lagoon during the same study period (October 2008 - September 2009). Based on the gonadal maturation stages detected in the histological sections of the gonads of B. brandaris (Vasconcelos et al. 2012b), the mean gonadal index $[GI=(\sum individuals each stage \times stage ranking)/total$ individuals each month] in both sexes was calculated (Seed 1976). For each stage of gonadal development, a numerical stage ranking was assigned as follows: I resting=0; II - pre-active=3; III - active=4; IV - ripe=5; V - partially spent=2; VI - spent=1. Accordingly, the GI ranged from 0 (all individuals in the monthly sample in the resting phase) to 5 (all individuals in the monthly sample in the ripe stage). In addition, based on data gathered on the visceral coil weight (VcW), female capsule gland weight (CgW) and male penis length (PL), the following bio-physiological indices of B. brandaris were calculated and expressed in percentage (weight or length): gonadosomatic index (GSI=VcW/ SpW) in both sexes (Vasconcelos et al. 2012b), the male penial index (PI=PL/SL) (Vasconcelos et al. 2011) and the female capsule gland index (CGI=CgW/ SpW) (Vasconcelos et al. 2012b).

Analyses of variance (ANOVA) were employed to compare specimen SL and TW between sexes and to assess the influence of sex, size class and reproductive status of the individuals on the respective MY. Whenever ANOVA assumptions (normality of data and homogeneity of variances) were not fulfilled, the non-parametric Kruskal-Wallis test (ANOVA on ranks) was performed. Each time significant differences among groups were detected by the ANOVA or Kruskal-Wallis test, pairwise multiple comparisons were made using Tukey or Dunn post-hoc tests, respectively. ANOVA's were performed following Zar (1996) and using the software package SigmaStat[©] (version 3.5). In all statistical analyses, significance level was considered for p<0.05.

RESULTS

A total of 563 individuals of *B. brandaris*, with broad ranges in both SL (48.6-107.7 mm) and TW (4.6-79.7 g), were subjected to analyses for estimating MY (Table 1). In the overall samples, females (n=308)

Table 1. – Descriptive statistics of the samples and meat yield of the purple dye murex (*Bolinus brandaris*) from the Ria Formosa lagoon. M, males; F, females; N, number of individuals; SL, shell length; TW, total weight; MY (%), meat yield. Data presented as mean±SD and range (minimum – maximum).

Sexes	N	SL (mm)	TW (g)	MY (%)
both	563	73.3±8.2 (48.6-107.7)	20.6±7.8 (4.6-79.7)	40.5±6.1 (25.8-56.1)
M	255	72.7±7.4 (51.6-96.7)	20.0±6.2 (5.0-46.8)	40.7±6.2 (25.8-55.3)
F	308	73.8±8.7 (48.6-107.7)	21.1±8.8 (4.6-79.7)	40.4±6.0 (27.2-56.1)

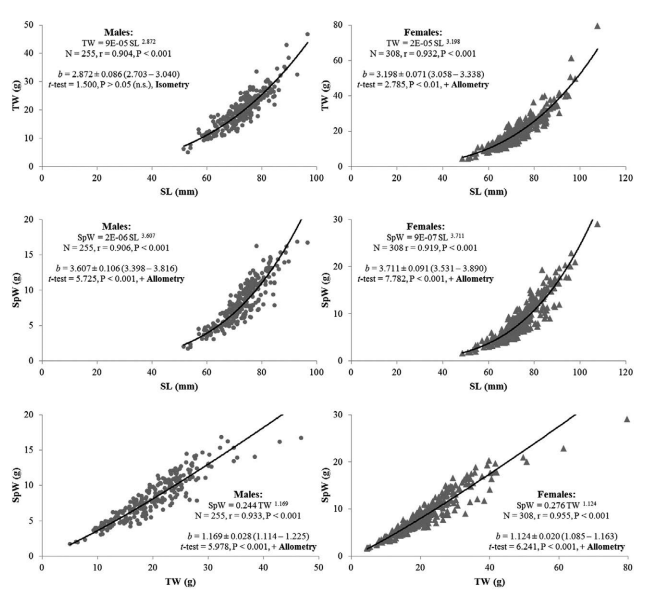


Fig. 3. - Relationships established between shell length (SL), total weight (TW) and soft parts weight (SpW) of Bolinus brandaris.

were significantly more abundant than males (n=255) (chi-square test: χ^2 =4.803, p<0.05), corresponding to a female-biased sex ratio of 1 M:1.2 F. Male and female *B. brandaris* showed similar mean shell length (males 72.7±7.4 mm SL; females 73.8±8.7 mm SL) and mean total weight (males 20.0±6.2 g TW; females 21.1±8.8 g TW), without statistically significant differences between sexes in either shell length (H=1.039, p>0.05) or total weight (H=0.326, p>0.05) (Table 1). Overall, average MY of *B. brandaris* was 40.5±6.1%, ranging from a minimum of 25.8% to a maximum of

56.1%. There were no statistically significant differences (H=0.667, p>0.05) in the average MY obtained from males ($40.7\pm6.2\%$; range=25.8-55.3%) and from females ($40.4\pm6.0\%$; range=27.2-56.1%) (Table 1).

The relationships established between specimen SL, TW and SpW in males and females of *B. brandaris* are presented in Figure 3. In both sexes, all relationships were highly significant (p<0.001) and invariably displayed high correlation coefficients (r=0.904 to 0.955), slightly higher in females (r=0.919 to 0.955) than in males (r=0.904 to 0.933) (Fig. 3). Overall, these

relationships indicated that specimen size (in both SL and TW) is an excellent indicator of SpW, and consequently of raw MY. Regarding the relative growth between variables (isometry vs. allometry), only the relationship SL vs. TW in males showed isometric growth (b=2.872±0.086; range=2.703-3.040), while all remaining relationships showed positive allometries (b=1.124 to 3.711) that revealed that throughout ontogeny the SpW increased at a proportionally faster rate than SL and TW (Fig. 3). Regarding the comparison of relative growth between males and females, the relationship TW vs. SpW displayed similar growth between sexes ($b_M=b_F$: t-test=1.514, p>0.05), whereas statistically significant differences between males and females were detected in the relationships SL vs. TW ($b_M < b_F$: t-test=8.704, p<0.001) and SL vs. SpW $(b_M < b_F: t-test=2.660, p<0.01)$, with females showing proportionally faster grow rates than males in both TW and SpW during ontogeny (Fig. 3).

The variation in MY as a function of the size of B. brandaris, with individuals of both sexes grouped into five SL size classes, is presented in Figure 4. Both males and females displayed highly significant differences in MY between size classes (males H=53.801, p<0.001; females H=56.595, p<0.001), clearly reflected in an evident trend of growing MY with increasing SL (Fig. 4). Similarly, the comparison of MY obtained from individuals below and above the MLS established for B. brandaris (MLS=65 mm SL) also showed highly significant differences in the MY obtained from individuals of both sexes below and above the MLS (males F=56.675, p<0.001; females H=27.004, p<0.001), reflecting much lower MYs in individuals below the MLS (males=34.4±4.7%; females=35.6±4.5%) than in individuals above the MLS (males=41.8±5.8%; females=41.0±5.9%) (Fig. 4). In both cases, i.e. with individuals grouped into five size SL classes and distributed below and above the MLS, there were no statistically significant differences (F or H, p>0.05) in MY between sexes within the same size class.

The monthly variation in the MY obtained from B. brandaris is presented in Figure 5. The MY of both males and females displayed highly significant monthly oscillation throughout the one-year study period (males F=37.583, p<0.001; females F=33.827, p<0.001), with both sexes reaching minimum yields in January (males 33.5±4.8%; females 34.2±3.7%) and maximum yields in June (males 49.4±2.6%; females $50.2\pm3.6\%$) (Fig. 5). Although the MY in both sexes exhibited a fairly similar and synchronous temporal trend, namely a marked decrease between December and January, followed by a clear increase from January to February, and another simultaneous sharp decrease between June and July, still some statistically significant differences in MY between sexes were detected in the monthly samples of October (females>males F=4.132, p<0.05), February (males>females F=6.860, p<0.05), March (females>males H=4.987, p<0.05) and September (males>females F=6.527, p<0.05) (Fig. 5).

The influence of diverse indicators of the reproductive status on the MY of *B. brandaris* is depicted in Figure 6. The monthly variation in MY in both sexes

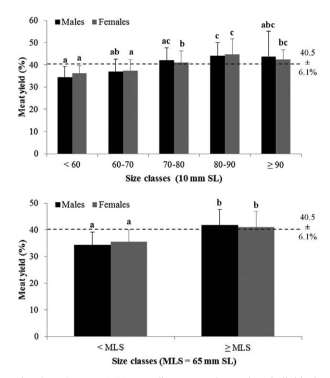


Fig. 4. – Average MY according to specimen size (individuals grouped into 10 mm SL classes) and as a function of the MLS=65 mm SL established for *Bolinus brandaris*. Interrupted lines denote average MY (mean±SD). In each sex, different superscript letters denote statistically significant differences (p<0.05) in MY between size classes.

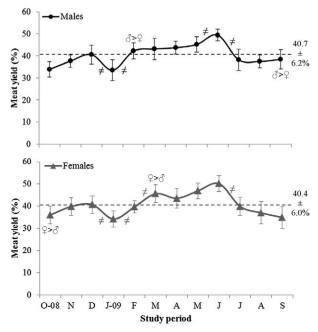


Fig. 5. – Monthly variation in the average MY of *Bolinus brandaris* males and females during the study period (October 2008 - September 2009). Interrupted lines denote average MY (mean±SD). In each sex, symbols denote statistically significant differences (p<0.05) in MY between consecutive months (\neq) or between sexes in each month (σ > \circ or \circ > σ).

slightly reflected the fluctuation in the mean GI during late spring - early summer. Indeed, sharp declines in male GI in May-August (from a maximum of 5.0 to a minimum of 1.1) and in female GI in June-August

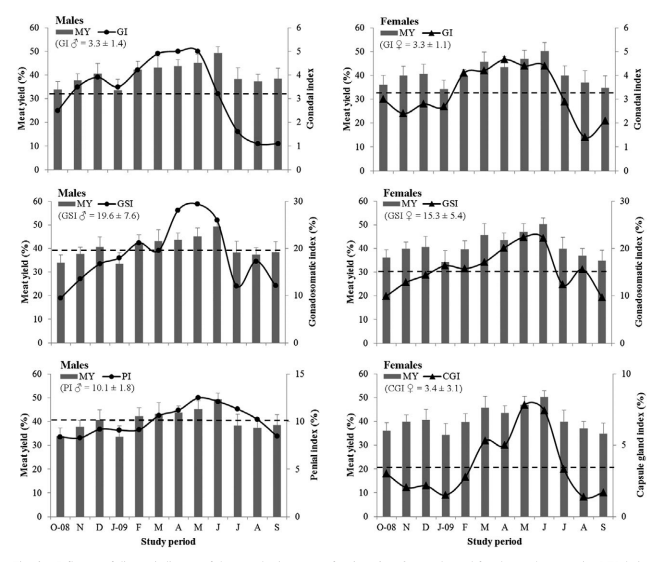


Fig. 6. – Influence of diverse indicators of the reproductive status of *Bolinus brandaris* males and females on the respective MY during the study period (October 2008 - September 2009). Interrupted lines denote average gonadal index, gonadosomatic index, penial index and capsule gland index (mean±SD).

(from 4.4 to a minimum of 1.4), reflecting the decrease in the proportion of ripe gonads and the subsequent increase in the proportion of partially spent and spent gonads, coincided with significant reductions in the MY of both males (from 45.1% in May to 37.4% in August) and females (from 50.2% in June to 37.0% in August) (Fig. 6). Similarly, the gonadosomatic index (GSI), indicative of gonadal maturation and inherent variation in gonad weight in both sexes, displayed abrupt decreases during the above period (males May-July; females June-July), reflecting gonad emptying during gamete release and female spawning. These decreases in both male GSI (from 29.4% in May to 12.2% in July) and female GSI (from 22.2% in June to 12.4% in July) were accompanied by marked reductions in MY in both males (from 45.1% in May to 38.2% in July) and females (from 50.2% in June to 39.9% in July) (Fig. 6). The sex-specific indices (male PI and female CGI) also exhibited the same general temporal trend of the previous bio-physiological indices. The PI, indicative of the copulative activity and consequent male gamete release, decreased continuously between the peak of copulation in May (12.5%) and September (8.5%), encompassing the period of marked reduction in male MY (from 49.4% in June to 37.4% in August). The CGI also displayed a sharp decrease between a maximum in May (7.8%) and a minimum in August (1.4%), indicating the deposition of egg capsules (oothecae) during the spawning season and consequent emptying of the capsule gland, which further contributed to the reduction in female MY recorded from June (maximum of 50.2%) to August (37.0%) (Fig. 6).

DISCUSSION

The average MY obtained from the purple dye murex (sexes combined) from the Ria Formosa lagoon was $40.5\pm6.1\%$, ranging between 25.8 and 56.1% during the study period. As stated above, information on the MY of gastropod species is still relatively scarce, and to the author's best knowledge in the available literature there is only one other study reporting the

Table 2. – Comparison of the meat yield of *Bolinus brandaris* from the Ria Formosa lagoon and other species within the family Muricidae, including some edible and/or commercially valuable muricids. Notes: Meat yield data presented as mean \pm SD and range (minimum – maximum); *, edible species; **, commercially valuable species; a, data converted from percentage shell weight to total weight; ^b, data expressed as percentage of soft-body total weight; ^c, variable depending on collecting site and date; N, number of individuals; SL, shell length range (mm); TW, total weight range (g); $\Diamond^+ \uparrow^{\bigcirc}$, sexes combined.

Species	Common nam	e Location	Meat yield (%)	Observations	Reference
Bolinus brandaris (Linnaeus, 1758)	purple dye murex**	Ria Formosa (Portugal), Atlantic Ocean	40.5±6.1 (25.8-56.1)	N=563; SL=48.6-107.7; TW=4.6-79.7; ♂+♀	present study
		Bay of Piran (Slovenia), Adriatic Sea	42.5ª	N=5	Dalla Via and Tappeiner (1981)
Chicoreus ramosus (Linnaeus, 1758)	branched murex**	Mandapan - Gulf of Mannar (India), Indian Ocean	≈50 ^b	_	Raghunathan et al. (1992)
Hexaplex trunculus (Linnaeus, 1758)	banded murex**	Bay of Piran (Slovenia), Adriatic Sea	33.6 ^a	N=20	Dalla Via and Tappeiner (1981)
		Kalithea Bay - Rhodes (Greece), Mediterranean Sea	34.2 (28.5-39.3) ^a	N=7; SL=39.0-8.0; TW=7.1-18.1	Alyakrinskaya (2005)
		Pathos Town (Cyprus), Mediterranean Sea	33.3 (31.6-37.6) ^a	N=5; SL=41.0-49.0; TW=8.3-14.3	Alyakrinskaya (2005)
		Ria Formosa (Portugal), Atlantic Ocean	37.9±4.6 (22.5-59.7)	N=1418; SL=40.2-82.8; TW=5.3-58.2; ♂+♀	Vasconcelos et al. (2009)
<i>Pinaxia coronata</i> A. Adams, 1853	rock shell**	Cross River Estuary (Nigeria), Atlantic Ocean	16.3 (11.9-19.5)	N≈500; SL=11.0-46.8	Udoh and Abiaobo (2014)
Rapana venosa (Valenciennes, 1846)	veined whelk	Novorossiysk Bay - Krasnodar (Russia), Black Sea	30.1 to 34.1 ^c (22.1-46.8) ^a	N=59; SL=43.0-80.0; TW=15.4-71.3	Alyakrinskaya (2005)
Stramonita haemas- toma (Linnaeus, 1767	red-mouthed)rock shell*	Magaluff Cape - Majorca (Spain), Mediterranean Sea	34.1 (22.1-53.3) ^a	N=3; SL=42.0-56.0; TW=11.3-27.7	Alyakrinskaya (2005)
		Las Galletas Bay - Tenerife (Spain), Atlantic Ocean	22.8 (16.4-26.9) ^a	N=6; SL=35.0-43.0; TW=5.0-10.1	Alyakrinskaya (2005)

MY of B. brandaris (42.5%) from the Adriatic Sea (Dalla Via and Tappeiner 1981). The comparison of the MY obtained from B. brandaris with that of other species of the family Muricidae, including some edible and/or commercially valuable muricids, is compiled in Table 2. It is worth stating that such comparisons in some muricids (e.g. B. brandaris, H. trunculus and S. haemastoma) should be cautiously interpreted because their edible content varies depending on the country where these species are used for human consumption. For instance, whereas in the northeastern Atlantic and northern Mediterranean the whole meat (foot and visceral mass) is ingested, in the southern Mediterranean only the foot is consumed. In general, the average MY of the purple dye murex from the Ria Formosa lagoon (40.5%) is higher than that obtained from other sympatric muricid species, namely H. trunculus, with values of 33.3% in Cyprus (Alyakrinskaya 2005), 33.6% in Slovenia (Dalla Via and Tappeiner 1981), 34.2% in Greece (Alyakrinskaya 2005) and 37.9% in Portugal (Vasconcelos et al. 2009), and S. haemastoma with values ranging from 22.8% in Tenerife (Alyakrinskaya 2005) to 34.1% in Majorca (Alyakrinskaya 2005) (Table 2). This higher MY of B. brandaris compared with H. trunculus and S. haemastoma is due to marked differences in their shells in terms of thickness, strength and ornamentation, and consequently, in their shell relative weight (e.g. Dalla Via and Tappeiner 1981, Alyakrinskaya 2005, Vasconcelos et al. 2016). For instance, the contribution of shell weight to TW in H. trunculus is almost 10% higher than that in B. brandaris (Dalla Via and Tappeiner 1981), whereas in S. haemastoma the contribution is over 10% higher than that in H. trunculus (Alyakrinskaya 2005).

The relationships established between *B. brandaris* size and the respective SpW revealed that both SL and TW are excellent indicators of SpW, and therefore of the raw MY that can be obtained from the purple dye murex. This information is most useful both for the seafood processing industry and for consumers, because the relationships SL vs. SpW and TW vs. SpW allow us to predict the edible content of B. brandaris as a function of the individual size (either in SL or TW), and thus estimate the MY that can be obtained from a batch of purple dye murexes with a certain size or weight range. Concerning relative growth, both males and females displayed positive allometries in the relationships SL vs. SpW and TW vs. SpW, revealing that during growth the edible content increased at a proportionally faster rate than SL and TW (i.e. the largest and heaviest individuals provide proportionally higher MYs), which in practice means that B. brandaris allocates more energy resources to somatic growth than to shell secretion throughout ontogeny. Inter-specific comparisons of the type of growth (isometry vs. allometry) are rather complicated due to high variability in shell morphology and structure between species and their variation during growth. Still, in B. brandaris the allometry coefficients of the relationships SL vs. SpW (males, b=3.607; females, b=3.711) and TW vs. SpW (males, b=1.169; females, b=1.124) are higher than those reported for other muricid species, such as Chicoreus ramosus (SL vs. meat weight, b=2.805) (Thapnu and Tantichodok 1991), Pinaxia coronata (SL vs. fresh tissue weight, b=2.801) (Udoh and Abiaobo 2014) and H. trunculus (SL vs. meat weight, b=3.557; TW vs. meat weight, b=1.118) (Vasconcelos et al. 2009). Concerning the comparison of relative growth between sexes, in the relationships SL vs. TW (males, b=2.872;

Craviac	I contion Contion Contion Control Contion Control C	Study namod	Spawning period	Deviod	Spawnir	Spawning events	Obcertvatione	R afaranca
Dolinus hum danie (1 in		normal barron	TO T	T CUIV		f mmmm h	10000	
poutnus pranaaris (LIII- naeus, 1758)		n.a.	May-Jun				n.a.	Lo Bianco (1888)
	Dor, Haifa, Palmahim, Tel-Aviv and Tel-Barukh, Israel ^{Med}	Jul 1945 – Jun 1977			May-Jul		FO (spawns cast ashore + dredged snawns)	Barash and Zenziper (1980)
	Golfe du Lion, France Med	n.a.	Spring				n.a.	Bartolome (1985)
	Sant Carles de la Ràpita, Catalo- nia Snain Med	Nov 1992 – Oct 1993	Jul-Jul		Aug		FO (dredged spawns)	Martín et al. (1995)
	Andalucía, Spain Att. Med	n.a.	May-Jun				n.a.	Anon. (2001)
	Catalonia, Spain Med	May-Dec 2000			May-Jun		FO (collected spawns)	Ramón and Flos (2001)
	Sant Carles de la Ràpita, Catalo- nia. Spain ^{Med}	Feb 1999 – Mar 2000	Apr; Jun-Jul	Jun-Jul			GH + GM; N=240; SLwS≈26-52	Ramón and Amor (2002)
	Marbella, Andalucía, Spain ^{Med}	Jun 1999- Mav 2000	May-Jul				GH + CI; N=2364; SI $\approx 30-60$	Tirado et al. (2002)
	Gulf of Roses, Catalonia, Spain Med Mar-Apr 2001	Mar-Apr 2001	May-Jul				FO (dredged spawns)	Mallol et al. (2004)
	Algarve coast, Portugal ^{Atl}	n.a.	May-Jun				n.a.	Malaquias (2007)
	Ria Formosa lagoon, Dortugal Atl	Jul 2005-Jun 2006			Jun		FO (observed spawns)	Vasconcelos et al. (2008a)
	Ria Formosa lagoon. Portugal ^{Atl}	Oct 2008-	Mav-Inl	lul-nul			CI; N=573;	Vasconcelos et al (2011)
	Dizarta lacton and small Gulf of	Sep 2009 Mar 2007	Mar/Am Main	TD: 110.			SL=51.6-97.9	
	Dizerte lagoon and small Gulf of Tunis, Tunisia Med	Feb 2008	Mar/Apr-May; Jun-Aug				SLwS=17.0-50.0	Abidli et al. (2012)
	Ria Formosa lagoon, Portugal ^{Atl}	Oct 2008- Sep 2009	May-Jul	Jun-Jul			GH + CI; N=567; SL=48.6-107.7	Vasconcelos et al. (2012b)
	Gulf of Gabès, Tunisia Med	Jan-Dec 2007	Apr-Jul	May-Jun			GH + GM + CI; N=1929; SI =31.0-95.1	Elhasni et al. (2013)
Hexaplex trunculus	Gulf of Naples, Italy ^{Med}	n.a.	May-Jun				n.a.	Lo Bianco (1888)
(Linnaeus, 1/38)	Banyuls-sur-Mer and Endoume, France ^{Med}	Apr- Jul 1932				mid-Jun – late-Jun	LO (spawns in aquaria)	Fischer and Raffy (1933)
	Lago di Faro, Sicily, Italy ^{Med}	n.a.				May-Jun	LO (spawns in aquaria); SI =50.0-100.0	Dulzetto (1946)
	Messina, Sicily, Italy Med	Jul 1949				Jul	LO (spawns in aquaria)	Dulzetto (1950)
	Istria coast, Croatia Adr	Jan-Feb 1974				Jan-Feb	LO (spawns in aquaria)	Wondrak (1974)
	Banyuls-sur-Mer, France Med	n.a.				May-Jun	LO (spawns in aquaria)	Bandel (1975)
	Dor, Herzliyya, Nahal Poleg, Tel- Aviv, Tel-Barukh, Israel ^{Med} and Cyprus ^{Med}	Jun 1965- Oct 1977	year-round	May-Jul			FO (spawns cast ashore)	Barash and Zenziper (1980)
	Akko Bay, Israel ^{Med}	May 1979-Sep 1980			May		FO (diving surveys)	Spanier (1981)
	Lago di Faro, Sicily, Italy ^{Med}	Jan-Dec 1983	Jun-Jul				GH	Albanese et al. (1983)
	Israel Med	n.a.			mid-Mar- Jun	mid-Mar-Jun	FO + LO (spawns in aquaria)	Spanier and Karmon (1987)
	Baie de Calvi, Corsica, France Med	Oct 1982- Sep 1984	Jul				GH; N=109	Martoja and Bouquegneau (1988)
	Italy ^{Adr} + Ionian and Tyrrhenian coasts ^{Med}	n.a.				May-Jun	LO (spawns in aquaria)	Terlizzi et al. (1999)

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Anon. (2001) Tirado et al. (2002) Lahbib et al. (2004) Vasconcelos et al. (2004) Vasconcelos et al. (2004) Abidli et al. (2006) Lahbib et al. (2006) D Trigui El-Menif et al. (2006)	Malaquias (2007) Vasconcelos et al. (2008b) Vasconcelos et al. (2008c) Abidli et al. (2009) Lahbib et al. (2009) Lahbib et al. (2009) Elhasni et al. (2010) Gharsallah et al. (2010)	Lahbib et al. (2010) Lahbib et al. (2011a) Lahbib et al. (2011a) Güler and Lök (2014) Barash and Zenziper (1980) Lahbib et al. (2011b) El Ayari et al. (2015)
n.a. Anon. (2001) $GH + CI; N=2222;$ Tirado et al. (2002) $SL=37-90 \text{ mm}$ $CI; N=2880$ $CI; N=2880$ Lahbib et al. (2004) $CI; N=2880$ Lahbib et al. (2004) $D(spawns in aquaria);$ Vasconcelos et al. (2004) $PO(spawns in aquaria);$ Vasconcelos et al. (2004) $PO(spawns in aquaria);$ Vasconcelos et al. (2004) $DO(spawns in aquaria);$ Vasconcelos et al. (2006) $LO(spawns in aquaria);$ $Lahbib et al. (2006)$ $LO(spawns in aquaria);$ $Lahbib et al. (2006)$ $LO(spawns in aquaria);$ $Lahbib et al. (2006)$ $LO(spawns in aquaria); N=100$ - Trigui El-Menif et al. (2006) $LO(spawns in aquaria); N=100$ - Trigui El-Menif et al. (2006) $ISO(SSL=40-60)$ Trigui El-Menif et al. (2006)	n.a. Malaquias (2007) n.a. Malaquias (2007) GH; N=1183; Vasconcelos et al. (200) SL=40.2-82.8 Vasconcelos et al. (200) SL=40.2-79.8 Vasconcelos et al. (200) SL=40.2-79.8 Vasconcelos et al. (200) CI; N=903; Vasconcelos et al. (200) LO (spawns in aquaria); Abidli et al. (2009) N=600; SL=40-70 Lahbib et al. (2009) CI; N=720 Lahbib et al. (2009) CI; N=720 Lahbib et al. (2009) CI; N=720 Lahbib et al. (2009) Offshug survasin aquaria); Lahbib et al. (2009) GM + CI; N=1870; Elhasni et al. (2010) SL=20.8.80.0 GM + CI; N=1478; GM + CI; N=1478; Gharsallah et al. (2010) SL ≥ 50 Gharsallah et al. (2010) SL ≥ 50 Gharsallah et al. (2010)	LO (spawns in aquaria); N=100; SL=35.2-57.4 GM + CI; N=343; SL=40-60 FO (observed spawns) LO (spawns in aquaria); N=39; SL=57.0-67.3 FO (collected spawns) LO (spawns in aquaria); N=90; SL=40-80 FO (observed spawns) GM+CI; N=1035; SL=20.0-80.0
late-Apr- early-Jun Mar Apr-Jul Feb-Jul Mar-Jul	early-Mar mid- to late-Apr	late-Apr – mid-May May late-May – mid-Jun
Feb-Jun		late-Feb- Mar Jun-Aug late-Apr – early-Aug
Apr-Jul	May-Jun May-Jun Mar-Apr Apr-May	
May-Jun Apr-Jul Jan-May; Aug-Sep Feb-May; Jul-Oct	May-Jun Mar-Jun Apr-Jun Jan-Feb Apr-May Mar-May Mar-May	Feb-Mar Jun-Jul; Sep-Oct
n.a. Jun 1999- May 2000 Jul 2002- Jul 2003 Jul 2003 Apr- Oct 2004 n.a. n.a. n.a. Jan-Jul 2003 Jan-Jul 2003	200 Ja M H M	-Apr 2006 Jan 2005- Jun 2007 Jun 2007- May 2008 Jun 2007- May 2008 n.a. Aug 1957- Jun 1971 Mar- Aug 2007 Jun 2009- May 2010
Andalucía, Spain ^{Atl.Med} Marbella, Spain ^{Med} Bizerte lagoon, Tunisia ^{Med} Bizerte channel, Tunisia ^{Med} Ria Formosa lagoon, Portugal ^{Ad} Ria Formosa lagoon, Portugal ^{Ad} Bizerte lagoon, Tunisia ^{Med} Bizerte lagoon, Tunisia ^{Med} Bizerte channel, Tunisia ^{Med}	Algarve coast, Portugal ^{All} Ria Formosa lagoon, Portugal ^{All} Ria Formosa lagoon, Portugal ^{All} North lake of Tunis, Tunisia ^{Med} Menzel Jemil, Bizerte lagoon, Tunisia ^{Med} Bizerte channel, Bizerte lagoon, Tunisia ^{Med} Menzel Jemil and Bizerte channel, Bizerte lagoon, Tunisia ^{Med} Gulf of Gabès, Tunisia ^{Med} Bizerte lagoon, Tunisia ^{Med}	Menzel Jemil, Bizerte lagoon, Tunisia ^{Med} Boughrara lagoon, Tunisia ^{Med} Boughrara lagoon, Tunisia ^{Med} Urla, Izmir, Turkey ^{Aeg} <i>Stramonia haemastoma</i> Tel-Aviv, Israel ^{Med} (Linnaeus, 1767) Zarzouna, Bizerte lagoon, Tunisia ^{Med} Zarzouna, Bizerte lagoon, Tunisia ^{Med} Artificial channel, Bizerte lagoon, Tunisia ^{Med}

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females, b=3.198) and SL vs. SpW (males, b=3.607; females, b=3.711), females exhibited significantly higher allometry coefficients than males, indicating faster grow rates in both TW and SpW throughout ontogeny, which are probably due to particular features of the female reproductive system, namely a female-specific accessory sexual organ, the capsule gland that reaches a considerable size and weight during the spawning season (please see further details below about the influence of the reproductive status on the MY of *B. brandaris*).

The trend of increasing MY during the ontogeny of *B. brandaris* was further highlighted by grouping the individuals of both sexes into SL size classes. Indeed, highly significant differences in MY between size classes in both males and females constitute additional evidence of growing MY with increasing SL, i.e. smaller individuals yield significantly less edible content than larger individuals. Accordingly, individuals below the MLS legally stipulated for B. brandaris (<65 mm SL) provided significantly lower MYs (males=34.4%; females=35.6%) than individuals that achieved the minimum landing size ($\geq 65 \text{ mm SL}$) (males=41.8%; females=41.0%), further reinforcing the importance of compliance with the species MLS by fishermen/harvesters, seafood traders and shellfish consumers. In fact, since commercially undersized specimens yield significantly lower edible content than individuals above the MLS, this should also help discourage their commercialization and consumption, in order to avoid unconsciously paying for shells at the price of meat, considering the high commercial value of the purple dye murex (which fetches prices around 20-25 € kg⁻¹ for first sale) (Vasconcelos et al. 2008a). Moreover, the existence of commercially undersized B. brandaris available in local shellfish suppliers and seafood markets justifies the need to improve regulation measures and reinforce inspection activities focusing on this artisanal fishery.

The MY of B. brandaris exhibited significant monthly variations and an analogous and synchronous temporal trend in both males and females during the study period, which were undoubtedly related to their reproductive status, as evidenced by the monthly oscillation in diverse bio-physiological indices calculated for both sexes. In fact, the most noteworthy monthly variation in the MY of both males and females was a significant decrease recorded from June to July, which was accompanied by a simultaneous reduction in most indices of reproductive condition, namely the mean GI and the GSI in both sexes, the male PI, and the female CGI. Chronologically, such sharp decreases in these indices correspond to the following sequential phases in the reproductive cycle of *B. brandaris*: peak of male copulative activity in May (PI); maturation peak in May (males) and June (females) followed by sharp decreases in both GI and GSI, which denote decreasing proportions of ripe gonads and consequent replacement by increasing proportions of partially spent and spent gonads (GI) as a consequence of gamete release (GSI) in both sexes; and female spawning peak between June and July (CGI). Overall, the sharp decreases in these

bio-physiological indices reflect significant reductions in the size and weight of male and female gonads and in the female capsule gland, which are inevitably translated immediately into a substantial decrease in SpW and consequently a significant decline in the raw MY of the purple dye murex between June and July. Indeed, during the period of reproductive maturation, both male and female B. brandaris develop large gonads and females fill the capsule gland responsible for the formation of the egg capsules (oothecae) that encapsulate the developing eggs and embryos, an accessory sexual organ that also reaches considerable size during the spawning season (e.g. Amor et al. 2007, Vasconcelos et al. 2012b, Elhasni et al. 2013), and whose deflation and emptying after gamete release and spawning is noticeably reproduced as an abrupt decrease in the raw MY of the purple dye murex.

In certain fishing resources, the optimum harvesting season should coincide with the period of highest MY (e.g. Udoh and Abiaobo 2014), which allows the weight of the catches/landings and the respective commercial value and resulting profits to be maximized. In the present case, besides the general concerns related to the sustainability of the fishing/ harvesting activity and the appropriate management of this fishing resource, data on the MY of the purple dye murex confirmed the importance and interest of accomplishing the MLS=65 mm SL legally stipulated for the catches of B. brandaris in the Ria Formosa lagoon (D.R. 1990, 2001), primarily by fishermen/ harvesters, but also by seafood traders and shellfish consumers. In addition, the present data also confirmed the influence of the reproductive status on the edible content of the purple dye murex, with the highest MYs being recorded in the period of reproductive maturation (i.e. prior to the gamete release and the spawning peak). In this particular, it should be taken into account that the fishing/harvesting activity targeting B. brandaris is subjected to a closed season (1 May - 30 June) (D.R. 2010), which was deliberately implemented during the spawning season in order to protect the female broodstock and collective spawns (Vasconcelos et al. 2012b). For this reason, although the highest MYs of B. brandaris coincide with the closed season of the fishing/harvesting activity, this temporary prohibition of the catches should be rigorously fulfilled by fishermen/harvesters in order to ensure a balanced and sustainable long-term exploitation of this locally important fishing resource of the Ria Formosa lagoon.

Finally, the marked influence of the reproductive status on the MY detected in the present study prompted a comparative assessment of the spawning season and peak, including spawning events observed in the field and in the laboratory, of three sympatric muricid species (*B. brandaris, H. trunculus* and *S. haemastoma*) throughout their geographic distributional range (Table 3). This multi-species comparative approach compiles most relevant baseline information consulted in virtually all literature available on this subject, providing supporting data from a long-time period (1888-2015) and thus allowing analysis of long-term trends. In ad-

dition, this dataset comprises information gathered at locations covering a wide geographical area (Atlantic Ocean, Mediterranean, Adriatic, Tyrrhenian, Ionian and Aegean Seas) and including ecologically diverse habitats (e.g. coastal areas vs. coastal lagoons). Overall, 55 data records on studies of the spawning season and peak of those three muricid species, obtained through biological sampling and analyses, field and laboratory observations, were collected from 41 bibliographic sources published by 27 different authors (Table 3).

According to the information available in the literature, these gastropod species display slight variation in the timing of reproduction throughout their distributional range, namely in the spawning season and peak. most probably associated with the colder seawater temperatures in the Atlantic Ocean compared with the Mediterranean Basin. As a consequence, Mediterranean populations of these species show a lengthy spawning season, generally ranging from spring to summer (sometimes even with an unusual longer duration), as well as more than one spawning period during the year (Table 3). Gonadal ripening and spawning are apparently triggered by the progressive rise in seawater temperatures from late spring to early summer. This reproductive dynamics induces a significant increase in the volume and weight of the gonads and accessory reproductive organs, subsequently reflected in the species MY, as is clearly demonstrated in the present study.

Indeed, gastropod reproductive dynamics, including the seasonality of gonadal ripening and spawning, are regulated by environmental cues that allow reproduction to occur at the most favourable time (Wayne 2001, Harding et al. 2008, Hotchkiss et al. 2008), with seawater temperature and food availability controlling the timing of spawning and being responsible for interannual variations in the reproductive cycle (Sternberg et al. 2010). Accordingly, as for the sympatric B. brandaris, H. trunculus and S. haemastoma, similar reproductive patterns (i.e. spawning season coinciding with rising seawater temperatures) have been reported for several other muricid species worldwide, such as Hexaplex nigritus (Cudney-Bueno et al. 2008), Rapana venosa (Saglam and Duzgunes 2007, Harding et al. 2008, Saglam et al. 2009), Reishia clavigera (Tong 1988, Lee 1999), Tenguella musiva (Tong 1988) and Trophon geversianus (Cumplido et al. 2010).

In conclusion, the present study showed that MY of the purple dye murex increases during ontogeny, as revealed by the relationships established between specimen size and SpW and by significant differences in MY between size classes. In addition, MY exhibited significant monthly variation directly related to reproductive status, as evidenced by the calculation of bio-physiological indices. Overall, these findings have implications at diverse levels, including the management, regulation and inspection of the artisanal fishing/harvesting activity and the commercialization and consumption of this valuable seafood product. In addition to *B. brandaris*, the same applies to other muricid species exploited for human consumption, including *H. trunculus* and *S. haemastoma*.

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REFERENCES

- Abidli S., Trigui El-Menif N., Boumaiza M. 2006. Variation de la concentration du TBT sur la fécondité et l'émission gamétique d'*Hexaplex trunculus*. In: Résumes des Communications du Troisième Congrès Franco-Tunisien de Zoologie, Tabarka, Tunisie, 3-7 November 2006, p. 73.
- Abidli S., Lahbib Y., Trigui El-Menif N. 2009. Effects of TBT on the imposex development, reproduction and mortality in *Hexaplex trunculus* (Gastropoda: Muricidae). J. Mar. Biol. Assoc. U.K. 89: 139-146. https://doi.org/10.1017/S0025315408002282
- Abidli S., Lahbib Y., Trigui El-Menif N. 2012. Relative growth and reproductive cycle in two populations of *Bolinus brandaris* (Gastropoda: Muricidae) from northern Tunisia (Bizerta Lagoon and small Gulf of Tunis). Biologia 67: 751-761. https://doi.org/10.2478/s11756-012-0060-7
- Albanese M.P., Calabrò C., Acosta Pomar M.L.C. 1983. Ciclo biológico di *Trunculariopsis trunculus* L. (Moll. Gast. Pros.) vivente nel Lago di Faro. Mem. Biol. Mar. Oceanogr. 13: 133-147.
- Alyakrinskaya I.O. 2005. Functional significance and weight properties of the shell in some mollusks. Biol. Bull. 32: 397-418. https://doi.org/10.1007/s10525-005-0118-y
- Amor M.J., Ramón M., Durfort M. 2007. Aspectos morfológicos y ultraestructurales de la glándula de la cápsula de *Bolinus brandaris* (Gastropoda, Prosobranchia). Boll. Malacol. 43: 78-86.
- Anon. 2001. Especies de Interés Pesquero en el Litoral de Andalucía. Vol. II - Invertebrados. Junta de Andalucía, Consejería de Agricultura y Pesca, Sevilla, 210 pp.
- Bandel K. 1975. Das embryonalgehäuse mariner prosobranchier der region von Banyuls-sur-Mer. Vie Milieu 25: 83-118.
- Bañón R., Rolán E., García-Tasende M. 2008. First record of the purple dye murex *Bolinus brandaris* (Gastropoda: Muricidae) and a revised list of non-native molluscs from Galician waters (Spain, NE Atlantic). Aquat. Invasions 3: 331-334. https://doi.org/10.3391/ai.2008.3.3.8
- Barash A., Zenziper Z. 1980. Egg masses of Mollusca from Mediterranean waters of Israel and notes on reproduction of the freshwater species *Theodoxus jordani* and *Melanoides tuberculata*. Veliger 22: 299-317.
- Bartolome C. 1985. Contribution à l'Étude du Gastéropode Murex brandaris (Linnaeus, 1758) dans le Golfe du Lion. Diplôme d'Etudes Approfondies de Sciences de l'Eau et Aménagement, Académie de Montpellier, Université des Sciences et Techniques du Languedoc, France, 92 pp.
 Cecalupo A., Vianello M., Perini L. 2006. Alcune forme aberranti
- Cecalupo A., Vianello M., Perini L. 2006. Alcune forme aberranti rinvenute nel Mare Adriatico di *Hexaplex trunculus* (Linnaeus, 1758) e *Bolinus brandaris* (Linnaeus, 1758). Not. S.I.M., Pubbl. Quadr. Soc. Malacol. 24: 13-15.
- Cudney-Bueno R., Prescott R., Hinojosa-Huerta O. 2008. The black murex snail, *Hexaplex nigritus* (Mollusca, Muricidae), in the Gulf of California, Mexico: I. Reproductive ecology and breeding aggregations. Bull. Mar. Sci. 83: 285-298.

Cumplido M., Averbuj A., Bigatti G. 2010. Reproductive seasonality and oviposition induction in Trophon geversianus (Gastropoda: Muricidae) from Golfo Nuevo, Argentina. J. Shellfish Res. 29: 423-428.

https://doi.org/10.2983/035.029.0219

Dalla Via G.-J., Tappeiner U. 1981. Morphological and functional correlates with distribution of Murex trunculus L. and Murex brandaris L. (Mollusca, Gastropoda) in the northern Adriatic. Boll. Zool. 48: 191-195.

oi.org/10.1080/11250008109439333

- D.R. 1990. Regulamento da pesca na Ria Formosa. Portaria N° 560/1990 de 19 de Julho. Diário da República, I Série, N° 165,
- pp. 3010-3012.
 D.R. 2001. Tamanhos mínimos de captura de espécies de peixes, crustáceos e moluscos. Portaria Nº 27/2001 de 15 de Janeiro. Diário da República, Iª Série-B, Nº 12, pp. 229-230.
- D.R. 2010. Regulamento da apanha de animais marinhos em águas oceânicas, águas interiores marítimas e não marítimas. Portaria Nº 1228/2010 de 6 de Dezembro. Diário da República, 1.ª Série, N° 235, pp. 5471-5477. Dulzetto F. 1946. Osservazioni sulla deposizione di *Murex truncu*-
- lus L. Atti Accad. Naz. Lincei 8: 1356-1361.
- Dulzetto F. 1950. Sull'origine e la struttura delle capsule ovifere di *Murex trunculus*. Arch. Zool. Ital. 35: 83-99. El Ayari T., Lahbib Y., Trigui El-Menif N. 2015. Associated fauna
- and effects of epibiotic barnacles on the relative growth and reproductive indices of *Stramonita haemastoma* (Gastropoda: Muricidae). Sci. Mar. 79: 223-232. 10.3989/scimar.04151.18A /doi.org/
- Elhasni K., Ghorbel M., Vasconcelos P., et al. 2010. Reproductive cycle and size at first sexual maturity of Hexaplex trunculus (Gastropoda: Muricidae) in the Gulf of Gabès (southern Tunisia). Invertebr. Reprod. Dev. 54: 213-225. ://doi.org/10.1080/0792 2010.96
- Elhasni K., Vasconcelos P., Ghorbel M., et al. 2013. Reproductive cycle of Bolinus brandaris (Gastropoda: Muricidae) in the Gulf of Gabès (southern Tunisia). Mediterr. Mar. Sci. 14: 24-35 loi.org/10.12681/mms
- FAO. 1989. Yield and nutritional value of the commercially more important fish species. FAO Fish. Tech. Pap. 309: 1-187
- Fischer P.H., Raffy A. 1933. La ponte du *Murex trunculus* L. Bull. Inst. Oceanogr. Monaco 618: 1-4.
 Gharsallah I.H., Vasconcelos P., Zamouri-Langar N., et al. 2010.
- Reproductive cycle and biochemical composition of Hexaplex trunculus (Gastropoda: Muricidae) from Bizerte lagoon, northern Tunisia. Aquat. Biol. 10: 155-166. doi.org/10.3
- Güler M., Lök A. 2014. Embryonic development and intracapsular feeding in *Hexaplex trunculus* (Gastropoda: Muricidae). Mar. Ecol. 35: 193-203. /doi.org/10.1111/maec.12066
- Harding J.M., Mann R., Kilduff C.W. 2008. Influence of environmental factors and female size on reproductive output in an invasive temperate marine gastropod Rapana venosa (Muricidae). Mar. Biol. 155: 571-581.

s://doi.org/10.1007/s00227-008-1044-6

Hotchkiss A.K., Sternberg R.M., LeBlanc G.A. 2008. Environmental cues trigger seasonal regression of primary and accessory sex organs of the mud snail, Ilyanassa obsoleta. J. Molluscan Stud. 74: 301-303. https://doi.org/10.1093/mollus/eyn009 Houart R. 2001. A Review of the Recent Mediterranean and North-

- eastern Atlantic Species of Muricidae. Ed. Evolver, Rome, 227
- Huxley J.S., Teissier G. 1936. Terminology of relative growth. Nature 137: 780-781. https://doi.org/10.1038/137780b0
- Katsanevakis S., Poursanidis D., Issaris Y., et al. 2011. "Protected" marine shelled molluscs: thriving in Greek seafood restaurants. Mediterr. Mar. Sci. 12: 429-438.
- Lahbib Y., Trigui El-Menif N., Le Pennec M., et al. 2004. Données sur le cycle reproducteur du mollusque gastéropode Hexaplex trunculus (Linné, 1758) de la Lagune de Bizerte (Tunisie). Bull. Soc. Zool. Fr. 129: 407-418.
- Lahbib Y., Trigui El-Menif N., Le Pennec M., et al. 2006. La ponte et la croissance absolue d'Hexaplex trunculus réalisée au laboratoire. In: Résumés des Communications du Troisième Congrès Franco-Tunisien de Zoologie, Tabarka, Tunisie, 3-7 Novembre 2006, p. 64.

- Lahbib Y., Abidli S., Trigui El-Menif N. 2009. Relative growth and reproduction in Tunisian populations of Hexaplex trunculus with contrasting imposex levels. J. Shellfish Res. 28: 891-898.
- Lahbib Y. Abidli S., Trigui El-Menif N. 2010. Laboratory study of the intracapsular development and juvenile growth of the banded murex, Hexaplex trunculus. J. World Aquacult. Soc. 41: 18-34.

https://doi.org/10.1111/j.1749-7345.2009.00310.x

Lahbib Y., Abidli S., Trigui El-Menif N. 2011a. Reproductive activity in the commercially exploited Mediterranean muricid Hexaplex trunculus collected from Boughrara lagoon (southern Tunisia). Russ. J. Mar. Biol. 37: 501-508.).1134

- Lahbib Y, Abidli S., Trigui-El Menif N. 2011b. Spawning and intracapsular development of Stramonita haemastoma haemastoma (Gastropoda: Muricidae) collected in northern Tunisia. Mar. Biol. Res. 7: 719-726. /doi.org/10.1080/17451000.2011.558099
- Lee J.H. 1999. Gametogenesis and reproductive cycle of the rock shell, *Reishia (Thais) clavigera* (Neogastropoda: Muricidae), on the west coast of Korea. Korean. J. Biol. Sci. 3: 375-383. 10.1080/122 5071.1999.96
- Lo Bianco S. 1888. Notizie biologiche riguardanti specialmente il periodo di maturità sessuale degli animali del Golfo di Napoli. Mitt. Zool. Stat. Neapel 8: 401-405. Macedo M.C.C., Macedo M.I.C., Borges J.P. 1999. Conchas
- Marinhas de Portugal (Seashells of Portugal). Ed. Verbo, Lis-
- boa, 516 pp. Malaquias M.A. 2007. Gastropoda. In: Borges T.C. (ed.), Biodiversity in the Fisheries of Algarve (South Portugal). Universidade do Algarve, Faro, pp. 196-241.
- Mallol S., Muñoz M., Hernández M.R., et al. 2004. Evaluation of the purple dye murex Bolinus brandaris (Mollusca: Gastropoda) population as a new fishery resource in the Gulf of Roses (Catalan coast, NW Mediterranean). Rapp. Comm. Int. Explor. Sci. Mer Mediterr. 37: 394-395
- Martín P., Sánchez P., Ramón M. 1995. Population structure and exploitation of Bolinus brandaris (Mollusca: Gastropoda) off the Catalan coast (northwestern Mediterranean). Fish. Res. 23: 319-331.

- https://doi.org/10.1016/0165-7836(94)00346-X Martoja M., Bouquegneau J.-M. 1988. *Murex trunculus*: un nouveau cas de pseudo-hermaphrodisme chez un gastéropode prosobranche. Bull. Soc. R. Sci. Liège 57: 45-58. Muzavor S., Morenito P.M. 1999. Roteiro Ecológico da Ria Formo-
- sa. Vol. IV Moluscos Gastrópodos. Universidade do Algarve, Faro, 85 pp. Oliver A.V. 2015. An ancient fishery of banded dye-murex (*Hexa*-
- plex trunculus): zooarchaeological evidence from the Roman city of Pollentia (Mallorca, Western Mediterranean). J. Archaeol. Sci. 54: 1-7.

https://doi.org/10.1016/j.jas.2014.11.026 Poppe G.T., Goto Y. 1991. European Seashells. Vol. 1 (Polypla-

- cophora, Caudofoveata, Solenogastra, Gastropoda). Verlag Christa Hemmen, Wiesbaden, 352 pp.
- Raghunathan C., Patterson Edward J.K., Ayyakkannu K. 1992. Utilization of the non-edible meat of the gastropods *Chicoreus* ramosus and Fasciolaria trapezium as a supplementary diet for penaeid prawn Penaeus indicus. Phuket Mar. Biol. Cent. Spec. Publ. 11: 9-15.
- Ramón M., Amor M.J. 2001. Increasing imposex in populations of Bolinus brandaris (Gastropoda: Muricidae) in the north-western Mediterranean. Mar. Environ. Res. 52: 463-475.
- https://doi.org/10.1016/S0141-1136(01)00101-5 Ramón M., Amor M.J. 2002. Reproductive cycle of *Bolinus* brandaris and penis and genital duct size variations in a population affected by imposex. J. Mar. Biol. Assoc. U.K. 82: 435-442. https://doi.org/10.1017/S002531
- Ramón M., Flos R. 2001. First trials to cultivate the muricid gastropod Bolinus brandaris (Linnaeus). Eur. Aquacult. Soc. Spec. Publ. 29: 219-220.
- Ramón M., Cano J., Peña J.B., et al. 2005. Current status and perspectives of mollusc (bivalves and gastropods) culture in the Spanish Mediterranean. Bol. Inst. Esp. Oceanogr. 21: 361-373.
- Reese D.S. 1980. Industrial exploitation of murex shells: purple-dye and lime production at Sidi Khrebish, Benghazi (Berenice). Lybian Stud. 11: 79-93.

doi.org/10.1017/S026371890000858X

Reese D.S. 2010. Shells from Sarepta (Lebanon) and East Mediter-

ranean purple-dye production. Mediterr. Archaeol. Archaeometry 10: 113-141.

Saglam H., Duzgunes E. 2007. Deposition of egg capsule and larval development of *Rapana venosa* (Gastropoda: Muricidae) from the south-eastern Black Sea. J. Mar. Biol. Assoc. U.K. 87: 953-957.

https://doi.org/10.1017/S0025315407056330

Saglam H., Duzgunes E., Ogut H. 2009. Reproductive ecology of the invasive whelk *Rapana venosa* Valenciennes, 1846, in the southeastern Black Sea (Gastropoda: Muricidae). ICES J. Mar. Sci. 66: 1865-1867.

https://doi.org/10.1093/icesjms/fsp184

- Seed R. 1976. Ecology. In: Bayne B.L. (ed), Marine Mussels: their Ecology and Physiology. Cambridge University Press, Cambridge, pp. 13-65.Sokal R.R., Rohlf F.J. 1987. Introduction to Biostatistics, 2nd edi-
- Sokal R.R., Rohlf F.J. 1987. Introduction to Biostatistics, 2nd edition. Freeman, New York, 363 pp.
- Spanier E. 1981. Behavioral ecology of the marine snail *Trunculariopsis (Murex) trunculus*. In: Shuval H. (ed.), Developments in Arid Zone Ecology and Environmental Quality. Balaban ISS, Philadelphia, pp. 65-70.
- Spanier E., Karmon N. 1987. Muricid snails and the ancient dye industries. In: Spanier E. (ed.), The Royal Purple and the Biblical Blue: Argaman and Tekhelet. Keter Publishing House Jerusalem Ltd., Jerusalem, pp. 179-192.
 Sternberg R.M., Gooding M.P., Hotchkiss A.K., et al. 2010. Envi-
- Sternberg R.M., Gooding M.P., Hotchkiss A.K., et al. 2010. Environmental-endocrine control of reproductive maturation in gastropods: implications for the mechanism of tributyltin-induced imposex in prosobranchs. Ecotoxicology 19: 4-23. https://doi.org/10.1007/s10646-009-0397-z
- Thapnu P., Tantichodok P. 1991. Basic data on *Chicoreus ramo-sus*: some length-weight relationships. Phuket Mar. Biol. Cent. Spec. Publ. 9: 94-96.
- Terlizzi A., Geraci S., Gibbs P.E. 1999. Tributyltin (TBT)-induced imposex in the neogastropod *Hexaplex trunculus* in Italian coastal waters: morphological aspects and ecological implications. Ital. J. Zool. 66: 141-146. https://doi.org/10.1080/11250009909356248
- Tirado C., Rodríguez de la Rua A., Bruzón M.A., et al. 2002. La Reproducción de Bivalvos y Gasterópodos de Interés Pesquero en Andalucía. Junta de Andalucía, Consejería de Agricultura y

Pesca, Huelva, 129 pp. Tong L.K.Y. 1988. The reproductive biology of *Thais clavigera* and

- Morula musiva (Gastropoda: Muricidae) in Hong Kong. Asian Mar. Biol. 5: 65-75.
- Trigui El-Menif N., Lahbib Y., Le Pennec M., et al. 2006. Intensity of the imposex phenomenon - impact on growth and fecundity in *Hexaplex trunculus* (Mollusca: Gastropoda) collected in Bizerta lagoon and channel (Tunisia). Cah. Biol. Mar. 47: 165-175.
- Udoh J.P., Abiaobo N.O. 2014. Condition index, meat yield and population structure of the marine gastropod, *Thais coronata*, off Cross River Estuary, Nigeria. Adv. Life Sci. Technol. 23: 24-32.
- Vasconcelos P., Gaspar M.B., Joaquim S., et al. 2004. Spawning of *Hexaplex (Trunculariopsis) trunculus* (Gastropoda: Muricidae) in the laboratory: description of spawning behaviour, egg masses, embryonic development, hatchling and juvenile growth

rates. Invertebr. Reprod. Dev. 46: 125-138. https://doi.org/10.1080/07924259.2004.9652616

- Vasconcelos P., Carvalho S., Castro M., et al. 2008a. The artisanal fishery for muricid gastropods (banded murex and purple dye murex) in the Ria Formosa lagoon (Algarve coast, southern Portugal). Sci. Mar. 72: 287-298.
- Vasconcelos P., Lopes B., Castro M., et al. 2008b. Gametogenic cycle of *Hexaplex (Trunculariopsis) trunculus* (Gastropoda: Muricidae) in the Ria Formosa Iagoon (Algarve coast, southern Portugal). J. Mar. Biol. Assoc. U.K. 82: 321-329. https://doi.org/10.1017/s0025315408000593
- Vasconcelos P., Lopes B., Castro M., et al. 2008c. Comparison of indices for the assessment of reproductive activity in *Hexaplex trunculus* (Gastropoda: Muricidae). Mar. Biol. Res. 4: 392-399. https://doi.org/10.1080/17451000802020436
- Vasconcelos P., Gaspar M.B., Castro M., et al. 2009. Influence of growth and reproductive cycle on the meat yield and proximate composition of *Hexaplex trunculus* (Gastropoda: Muricidae). J. Mar. Biol. Assoc. U.K. 89: 1223-1231. https://doi.org/10.1017/S00253154090003026
- https://doi.org/10.1017/S0025315409003026 Vasconcelos P., Gaspar M.B., Barroso C.M. 2010. Imposex in *Bolinus brandaris* from the Ria Formosa lagoon (southern Portugal): usefulness of "single-site baselines" for environmental monitoring. J. Environ. Monit. 12: 1823-1832. https://doi.org/10.1039/c0em00261e
- Vasconcelos P., Moura P., Barroso C.M., et al. 2011. Size matters: importance of penis length variation on reproduction studies and imposex monitoring in *Bolinus brandaris* (Gastropoda: Muricidae). Hydrobiologia 661: 363-375. https://doi.org/10.1007/s10750.010.0544-9
- https://doi.org/10.1007/s10750-010-0544-9 Vasconcelos P., Pereira A.M., Constantino R., et al. 2012a. Growth of the purple dye murex, *Bolinus brandaris* (Gastropoda: Muricidae), marked and released in a semi-intensive fish culture earthen pond. Sci. Mar. 76: 67-78. https://doi.org/10.3989/scimar.03313.21B
- Vasconcelos P., Moura P., Barroso C.M., et al. 2012b. Reproductive cycle of *Bolinus brandaris* (Gastropoda: Muricidae) in the Ria Formosa lagoon (southern Portugal). Aquat. Biol. 16: 69-83. https://doi.org/10.3354/ab00434
- Vasconcelos P., Barroso C.M., Gaspar M.B. 2016. Morphometric relationships and relative growth of *Hexaplex trunculus* and *Bolinus brandaris* (Gastropoda: Muricidae) from the Ria Formosa lagoon (southern Portugal). J. Mar. Biol. Assoc. U.K. 96: 1417-1425.
- https://doi.org/10.1017/S0025315415001472
 Vela J.M., Moreno Ó. 2004. Posibilidades del cultivo de gasterópodos murícidos en la región Suratlántica Ibérica. In: Seminário sobre Moluscicultura no Litoral Sul-Atlântico - Diversificação (DiverAqua). Tavira, 19-20 February 2004.
 Wayne N.L. 2001. Regulation of seasonal reproduction in mollusks.
- Wayne N.L. 2001. Regulation of seasonal reproduction in mollusks. J. Biol. Rhythms 16: 391-402.

https://doi.org/10.1177/074873001129002097

- Wondrak G. 1974. Laichverhalten und gelegebildung bei Murex trunculus L. (Gastropoda: Prosobranchia). Arch. Mollusk. 104: 115-119.
- Zar J.H. 1996. Biostatistical Analysis, 3rd Edition. Prentice-Hall International Inc., New Jersey, 662 pp.