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Slow growth of the overexploited milk shark *Rhizoprionodon* acutus affects its sustainability in West Africa

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Age and growth of *Rhizoprionodon acutus* were estimated from vertebrae age bands. From December 2009 to November 2010, 423 *R. acutus* between 37 and 112 cm total length $(L_{\rm T})$ were sampled along the Senegalese coast. Marginal increment ratio was used to check annual band deposition. Three growth models were adjusted to the length at age and compared using Akaike's information criterion. The Gompertz growth model with estimated size at birth appeared to be the best and resulted in growth parameters of $L_{\infty} = 139.55$ ($L_{\rm T}$) and K = 0.17 year⁻¹ for females and $L_{\infty} = 126.52$ ($L_{\rm T}$) and K = 0.18 year⁻¹ for males. The largest female and male examined were 8 and 9 years old, but the majority was between 1 and 3 years old. Ages at maturity estimated were 5.8 and 4.8 years for females and males, respectively. These results suggest that *R. acutus* is a slow-growing species, which render the species particularly vulnerable to heavy fishery exploitation. The growth parameters estimated in this study are crucial for stock assessments and for demographic analyses to evaluate the sustainability of commercial harvests.

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Key words: age; chondrichthyan; life-history traits; Senegal; vertebrae.

INTRODUCTION

Chondrichthyans play important functional roles in the top-down control of coastal and oceanic ecosystem structure and function (Ferretti *et al.*, 2010; Heithaus *et al.*, 2012). This group is unintentionally caught as by-catch in many fisheries and those sharks could reach 50% of all shark landings (Stevens *et al.*, 2000; Dulvy *et al.*, 2014). More than 700 000 t of cartilaginous fishes are harvested annually worldwide (Frisk *et al.*, 2001). The responses of sharks to an increased harvest pressure differs between species but there is a considerable concern about shark fisheries around the world as their declining catches are mostly attributed to overfishing (Baum *et al.*, 2003). Shark management and conservation are hindered by a lack of knowledge at the population level (Baum *et al.*, 2003), as well as by the lack of basic biological information (Frisk *et al.*, 2001). One of the most captured genera, *Rhizoprionodon*, is represented worldwide by seven species of small coastal carcharhinids (Compagno,

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1984), although only few studies have given consideration to their importance to fisheries (Castillo-Geniz *et al.*, 1998).

The milk shark *Rhizoprionodon acutus* (Rüppell 1837) is the most widely distributed species of this genus, extending all along the West African coast, in the Indo-Pacific, in the northern part of the Indian Ocean, from the Indonesian to the Philippine archipelagos and from Japan to Australia (Compagno, 1984; Capapé *et al.*, 2006; Harry *et al.*, 2010). The species is common on the West African continental shelf (Capapé *et al.*, 1994, 2006; Ba *et al.*, 2013*a*, *b*) in water depths up to 200 m (Compagno, 1984). It is the most landed shark species on the Senegalese coasts (Ba *et al.*, 2013*a*) and is also known in many countries, but there are very few available data on their populations status or fisheries (Simpfendorfer, 2003).

The life-history traits of this species vary significantly with its geographic location according to the literature: total length $(L_{\rm T})$ at first sexual maturity occurs between 65 and 95 cm, and at birth between 30 and 50 cm (Henderson et~al., 2006; Valadou et~al., 2006; Ba et~al., 2013a). The reproductive cycle ranges from seasonal (Bass et~al., 1975; Capapé et~al., 2006; Ba et~al., 2013a), partially seasonal (Henderson et~al., 2006), to fully aseasonal within northern Australian waters (Stevens & Mcloughlin, 1991).

The age and growth estimations are fundamental components of fisheries management, and maximizing their accuracy is critical for stock assessment (Cailliet *et al.*, 2006). The age of chondrichthyan species is usually estimated when interpreting growth increments deposited in calcified structures such as vertebrae, dorsal spines or caudal thorns (Goldman *et al.*, 2012). The timing and periodicity of growth increment deposition must be validated but periodicity is typically annual for chondrichthyans (McAuley *et al.*, 2006) as the mean birth dates are well defined most of the time by the viviparous type of reproduction (Cailliet & Goldman, 2004).

Very few data on the life history of *R. acutus* are available despite the fact that this species is heavily exploited (Simpfendorfer, 2003). Given the importance of life-history information in the development of age structured population models (Goldman *et al.*, 2012), the first objective of this study was to gather information about the age and growth of the *R. acutus* landing along the West African coast using interpretation of growth marks on vertebrae. These data will be useful for designing an effective management strategy for fisheries.

MATERIALS AND METHODS

SPECIMEN COLLECTION

From December 2009 to November 2010, R. acutus were sampled monthly from the artisanal fishery catches along the Senegalese coast (Fig. 1). Individuals were captured by using drift nets, surface longlines, sole gillnets and purse seines. Once fishermen landed the R. acutus, L_T (cm, stretched total length, from the tip of the snout to the extremity of the caudal fin, along the main axis of the body), total mass (W, W, W), sex and maturity information were recorded. Maturity of males and females were determined macroscopically (Ba et al., 2013a). Afterwards, the five largest vertebrae were removed in each individual from the region just below the first dorsal fin, stored on ice and then frozen before treatment at the laboratory.

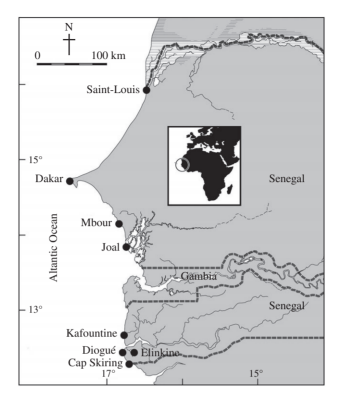


Fig. 1. Sampling locations of *Rhizoprionodon acutus* in Senegal.

PREPARATION AND INTERPRETATION OF VERTEBRAE GROWTH MARKS

In the laboratory, vertebrae were cleaned by immersion in a solution of 5% sodium hypochlorite and by removing adhering tissues under a stereoscope (Huveneers *et al.*, 2013). The minimum immersion time was 30 min in duration and varied depending on the vertebra size, and each one was then washed with water for 30-60 min and air-dried for 24-48 h also depending on its size. The preparation method of vertebrae was specifically developed for this study after different tests. Vertebrae were embedded in polyester resin before being sectioned along the sagittal axis with a low speed diamond saw (Isomet, Buehler; www.buehler.com). Vertebrae sections were then etched with 5% acetic acid for 2 min and then stained with toluidine blue for 5 min before being gently rinsed with water The stained vertebra sections were observed with a stereomicroscope under transmitted light coupled with a 3CCD video camera (Sony DXP; www.sony.com) and a calibrated image of each section was recorded. Vertebral radius (V_R in mm) was measured from the focus along the vertebral body to the edge using the TNPC software (http://www.tnpc.fr/en/visilog.html).

AGE ESTIMATION AND VALIDATION

To avoid bias interpretation, growth increments (thin stainable bands) were interpreted and counted without prior knowledge of the $L_{\rm T}$ or sex of the specimens. The same reader interpreted all vertebrae sections twice. The average percentage of error (E) between readings

was calculated using the Beamish–Fournier (1981) formula:
$$E = 100R^{-1}\sum_{i=1}^{R} \left| X_{ij} - \overline{X}_{j} \right| \overline{X}_{j}^{-1}$$
,

where X_{ij} is the *i*th age estimation of the *j*th fish, \overline{X}_j is the mean age of the *j*th fish and *R* is the number of times each fish is aged. Chang (1982) suggested incorporating the s.d. in the previous equation rather than the absolute deviation from the mean age. The resulting equation produces an estimate of the coefficient of variation (c.v., *X*) of consecutive readings:

$$X = 100 \left[\left(\sqrt{\sum_{i=1}^{R} \left(\left(X_{ij} - \overline{X}_{j} \right) (R-1)^{-1} \right)^{2}} \right) \overline{X}_{j}^{-1} \right].$$

Each growth cycle included a thin dark-stained band (translucent and more chromophilic band) and a large band (opaque and less chromophilic band) that were considered together as one annual cycle before the subsequent validation. The thin dark-stained band, ring, was counted for age estimation. The radius along the vertebral body was measured starting from the focus to the edge, the number of rings was counted along this axis and the distance between the vertebra focus and each ring was also measured. The 0+ year age class corresponding to the first distinct stained band defined as the birthmark was excluded from this analysis to ensure that growth from the birthmark did not affect the results. The distance from the last stained band to the vertebra edge was measured along the vertebral body in order to validate the timing of the band deposit. The mean marginal increment ratio $(I_{\rm M})$ for each month was plotted for females and males to determine if there was a yearly pattern in margin width (Natanson et al., 1995; Liu et al., 1999) using the following equation: $I_{\rm M} = (V_{\rm R} - R_n)(R_n - R_{n-1})^{-1}$, where $V_{\rm R}$ is the distance between the focus and the edge of the vertebra (in mm), R_n the distance between the focus and the last chromophilic band (in mm) and R_{n-1} is the distance from the centra to the penultimate band. Differences among monthly mean $I_{\rm M}$ were tested for heteroscedasticity by Bartlett's test and a non-parametric Kruskal-Wallis one-way ANOVA was used to test for differences in I_M by months (Simpfendorfer et al., 2000; Sulikowski et al., 2005). Monthly mean size was compared by one-way ANOVA. A Kolmogorov-Smirnov test was used to compare $L_{\rm T}$ frequency distributions between males and females. The relationship between L_T and V_R was calculated using linear regression analysis. An ANCOVA was used to compare the allometric relationships between sexes.

GROWTH MODELS

Since the parturition season of R. acutus may occur from May to June along the Senegalese coast (Capapé et al., 2006; Ba et al., 2013a), the mean birth date was set as 1 May in this study. Three growth models were adjusted to the length-at-age data for comparison and selection, and one to mass-at-age data. All models were fitted separately for males and females as sex differentiation is recognized for shark species. The first model was the von Bertalanffy growth function (VBGF) (von Bertalanffy, 1938): $L_t = L_{\infty} \left[1 - e^{-K(t-t_0)} \right]$, where L_t is the predicted length-at-age t, L_{∞} is the asymptotic length, K is the growth coefficient and t_0 is the x-intercept. Two Gompertz models (Mollet et al., 2002), one with fixed size at birth (GGF1) and the other with estimated size at birth (GGF2), were fitted to length-at-age data for comparison, and given by the following equation: $L_t = L_b e^{\left[\ln\left[L_{\infty}L_b^{-1}\right]\left[1-e^{(-Kt)}\right]\right]}$, where, in the first version of the Gompertz model (GGF1), L_b is the published size at birth 39 cm (L_T) (Capapé et al., 2006), and the other parameters are as described above. In the second version of the Gompertz model (GGF2), the estimated size at birth (L_b) replaced the published birth length (L_b), and was calculated from the published length at birth (L_b) using the estimated parameters (L_{∞} , K and t_0) of the VBGF, and given by the formula: $L_b = L_{\infty} \left[1 - e^{(K \times t_0)}\right]$, where K is the growth coefficient and t_0 is the x-intercept. Bias between published size at birth and estimated size at birth was defined as the ratio of L_b and L_0 (i.e. L_b L_0^{-1}) (Pardo et al., 2013).

x-intercept. Bias between published size at birth and estimated size at birth was defined as the ratio of $L_{\rm b}$ and L_0 (i.e. $L_{\rm b}$ L_0^{-1}) (Pardo et al., 2013). Mass-at-age data was modelled with a modified VBGF (Ricker, 1979): $W_t = W_{\infty} \left[1 - e^{-K(t-t_0)} \right]^3$, where W_t is the mass-at-age t, W_{∞} represents theoretical asymptotic mass and the other parameters are as previously defined.

Coefficient of determination values (r^2) were used in order to compare the fit of non-linear regression models described above with that of a linear intercept-only model, and as indicators

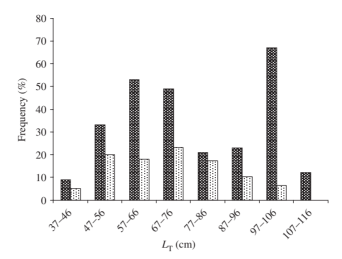


Fig. 2. Total length (L_T) frequency for *Rhizoprionodon acutus* sampled during this study [\mathbb{B} , females, n = 267; \mathbb{E} , males, n = 156].

of the proportion of the total variation in $L_{\rm T}$ that is explained (or accounted for) by regressions with age (Kvalseth, 1985): $r^2 = 1 - \left[\sum \left(y - \widehat{y}\right)^2\right] \left[\sum \left(y - \overline{y}\right)^2\right]^{-1}$.

The three growth models fitted to length-at-age data were discriminated within a model selection framework using the small-sample corrected Akaike's information criterion (AIC_c) (Burnham & Anderson, 2002). AIC_c (X) is given by the following equation: $X = n \log(Z) n^{-1} + 2p[n(n-p-1)^{-1}]$, where Z is the residual sum of squares, p is the number of fitted parameters in the model (plus one for the estimated variance) and n is the sample size. The model with the lowest AIC_c is considered to fit the data best. All models were fitted separately for males and females as sex differentiation is recognized for shark species and all statistical analyses and model adjustments were preformed with the Statistica software (www.statsoft.com).

RESULTS

MORPHOLOGICAL MEASUREMENTS

A total of 267 females and 156 males of R. acutus were sampled. The $L_{\rm T}$ of females (63·1% of the total sample) was comprised between 38 and 112 cm and that of males (36·9% of the total sample) between 37 and 104 cm (Fig. 2). Mean \pm s.e. $L_{\rm T}$ for the whole sample was $78\cdot14\pm19\cdot86$ cm for females and $70\cdot26\pm14\cdot86$ cm for males. A significant difference was found in the monthly comparison of female sizes (ANOVA, $F_{1,11}=4\cdot76$, $P<0\cdot05$) and male sizes (ANOVA, $F_{1,11}=3\cdot11$, $P<0\cdot05$). The $L_{\rm T}$ frequency distributions were different between males and females, females being generally larger (Kolmogorov–Smirnov, $P<0\cdot05$). The relationship between $L_{\rm T}$ and $V_{\rm R}$ for males and females was significantly different (ANCOVA, $F_{2,425}=1\cdot03$, $P<0\cdot05$); therefore, all subsequent models were fitted separately for males and females. The linear regression was $L_{\rm T}=6\cdot843V_{\rm R}+28\cdot276$ ($r^2=0\cdot87$) for females and $L_{\rm T}=6\cdot364V_{\rm R}+29\cdot872$ ($r^2=0\cdot76$) for males.

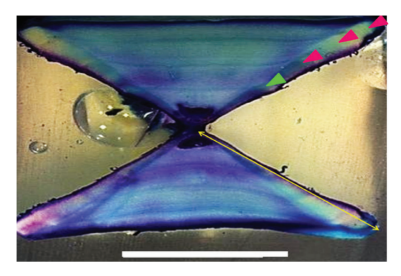


Fig. 3. Sagittal stained section of a vertebrae from a male *Rhizoprionodon acutus* (total length, *L*_T = 66 cm) with three interpreted growth bands. This individual was estimated to be 3 years old. →, birthmark; →, interpreted growth bands for age estimation; →, the vertebral body. White scale bar = 5 mm.

VERIFICATION OF GROWTH BAND PERIODICITY AND AGE ESTIMATION

Stained growth bands were visible on the vertebrae and were numbered along the vertebral body (Fig. 3). If bands were not clearly visible and interpretable, the corresponding vertebrae were excluded from the analysis. The average per cent error (APE) value ranged from 2.99% (for the 2+ year class, n=45) to 1.61% (for the 4+ year class, n=69), with a mean APE of 1.55% for the whole sample between the two readings. The c.v. value ranged from 4.22% (for the 2+ year class, n=45) to 2.28% (for the 4+ year class, n=69) with a mean c.v. of 2.20% for the whole sample between the two readings. These values gave enough precision of the results and then allowed age estimation using the chromophilic growth bands.

Significant differences were found in the marginal increment $I_{\rm M}$ among months for females (Kruskal–Wallis, $H=47\cdot98$, d.f. = 11, $P<0\cdot05$) and males (Kruskal–Wallis, $H=30\cdot50$, d.f. = 11, $P<0\cdot05$). The distance between the last stained band and the vertebra edge showed an annual increase until May for females and June for males followed by a decrease until August for females and September for males (Fig. 4). The lowest $I_{\rm M}$ was recorded in August and September for females and males, respectively, assuming that the stained band was formed annually in this period for both sexes.

GROWTH ESTIMATION

Age estimation showed that the young individuals dominated the whole sample (Fig. 5). The three growth models used in this study (VBGF, GGF1 and GGF2) fitted the data well for growth in length (0·65 < r^2 < 0·85; Table I). Based on the AIC_c values incorporating separate parameter estimates for each sex, the GGF2 outperformed the GGF1 and the VBGF (Table I) and fitted the best model length-at-age data. For both the sexes, the standard VBGF-estimated L_{∞} , 152 and 117 cm for females and

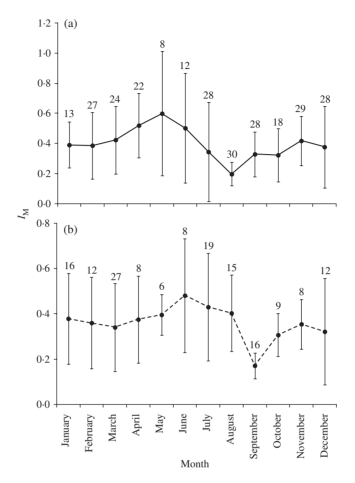


Fig. 4. Mean \pm s.E. monthly marginal increment ratio $(I_{\rm M})$ for (a) female (n=267) and (b) male (n=156) Rhizoprionodon acutus. Numbers indicate sample size,

males, was higher than the observed maximum, 112 and 104 cm for females and males [Table I and Fig. 6(a)]. The GGF models produced a larger asymptotic size for females than males [Table I and Fig. 6(b), (c)]. For females, both estimated and fixed size at birth in the GGF models, L_{∞} (139 and 125 cm), were significantly higher than the observed maximum length of 112 cm. For males, the L_{∞} estimated with GGF2 (126 cm) was significantly larger than the observed maximum length of 104 cm, while for the GGF1 (105·06 cm), it was almost similar to the observed maximum length. The maximum observed age was 9 years for females (106 cm $L_{\rm T}$) and 8 years for males (104 cm $L_{\rm T}$) (Fig. 6). Bias between published size at birth and estimated size at birth given by the ratio $L_{\rm b} L_{\rm 0}^{-1}$ were 0·86 and 0·82 for females and males.

For both forms of the GGF model, the growth coefficient in length at age was higher for males (0.29 and 0.18 for GGF1 and GGF2) than females (0.23 and 0.17 for GGF1 and GGF2; Table I). The growth model used in this study fitted the data well for growth in mass (0.68 < r^2 < 0.70; Table I). The mass asymptotic value of females was 9150 g, whereas that of males was 7755 g (Fig. 7). For both sexes, asymptotic mass estimated

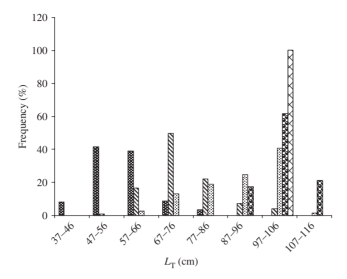


Fig. 5. Total length (L_T) -frequency of *Rhizoprionodon acutus* by age classes ($\bowtie 3, 1+2$ years; $\bowtie 3, 3+4$ years; $\bowtie 3, 5+6$ years; $\bowtie 3, 7+8$ years; $\bowtie 3, 9+$ years; n=423).

values were higher than the maximum observed masses (7000 g for females and 5300 g for males). The growth coefficients in mass for both males and females were almost similar (Table I). After the fifth year, mass gain of females was higher than males (Fig. 7). The evolution of the monthly mean mass showed the smaller increase during the months of June and July. This period of reduced overall growth corresponded with the deposition of a stainable band in the vertebrae in August and September.

DISCUSSION

RHIZOPRIONODON ACUTUS AGE ESTIMATION AND VERIFICATION

This study has provided estimates of age and growth for *R. acutus*, a coastal shark species, which is the most landed shark species along the Senegalese coast and easily accessible to small-scale fisheries (Capapé *et al.*, 2006; Ba *et al.*, 2013*a*). This is the first comprehensive study of age and growth for age estimation in this species in West African waters. It used stained growth bands on vertebrae for this purpose.

The values of indices comparing consecutive interpretations of vertebrae growth bands (APE and c.v. in this study) indicated a very good precision for the age estimation of *R. acutus* with stained bands distinguishable on vertebrae. Cailliet & Goldman (2004) mentioned that there has been an increase in the use of both verification and validation methodologies in chondrichthyan age and growth estimation studies, such as the use of the marginal increment ratio. The hypothesis of annual growth increment deposition has been supported in this study by the marginal increment ratio on the vertebrae edge. This method demonstrated that a unique growth band, consisting of one thin stained band, was formed annually. The minimal width of the marginal

Table I. Growth model parameters [asymptotic length (L_{∞}) and mass (W_{∞}) , growth coefficient (K)]: von Bertalanffy growth function (VBGF) for total length $(L_{\rm T},{\rm cm})$ and mass (g), and Gompertz growth model (with fixed size at birth (GGF1) and estimated size at birth (GGF2) for $L_{\rm T}$, for male and female *Rhizoprionodon acutus*

Models	Sexes	Parameters	Estimatio	n S.E.	t-values	P	r^2	AIC_c
VBGF	Females	L_{∞}	152-21	15.89	9.58	<0.001		1098-90
		\widetilde{K}	0.11	0.03	4.42	< 0.001	0.85	
		t_0	-2.72	0.44	-6.24	< 0.001		
	Males	L_{∞}	116.83	13.29	8.79	< 0.001	0.76	627.11
		\widetilde{K}	0.17	0.05	3.27	< 0.001		
		t_0	-2.27	0.59	-3.82	< 0.001		
GGF1	Females	L_{∞}	125.98	4.05	31.03	< 0.001	0.85	480.87
		K	0.23	0.01	16.97	< 0.001		
	Males	L_{∞}	105.06	4.64	22.631	< 0.001	0.77	478.99
		K	0.29	0.02	11.00	< 0.001		
GGF2	Females	L_{∞}	139.55	6.9	20.05	< 0.001	0.85	469.49
		K	0.17	0.01	12.87	< 0.001		
	Males	L_{∞}	126.52	12.52	10.10	< 0.001	0.78	466.34
		K	0.18	0.02	6.63	< 0.001		
Mass VBGF	Females	W_{∞}	9148.59	1809.37	5.06	< 0.001	0.68	1739-36
		K	0.19	0.04	4.47	< 0.001		
		t_0	-0.51	0.55	-0.93	>0.05		
	Males		7756-27	2528-27	3.07	< 0.001	0.70	1621.19
		K	0.20	0.07	2.94	< 0.001		
		t_0	-0.57	0.73	-0.78	>0.05		

AIC, Akaike information criteria.

increment was recorded for *R. acutus* captured in August for females and in September for males. The edge analysis supported the hypothesis that bands were formed at the end of the rainy season (May to September). An annual periodicity for growth band formation is common in temperate and subtropical shark species (Branstetter, 1987*a*; Simpfendorfer *et al.*, 2002; Wintner *et al.*, 2002; Joung *et al.*, 2004; McAuley *et al.*, 2006). This was also verified for different species of the *Rhizoprionodon* genus: *Rhizoprionodon taylori* (Ogilby 1915) in Australia (Simpfendorfer, 1993), the Atlantic sharpnose shark *Rhizoprionodon terraenovae* (Richardson 1836) in the Gulf of Mexico (Branstetter, 1987*b*; Carlson & Baremore, 2003) and in the south-eastern U.S.A. (Loefer & Sedberry, 2003). This annual growth band deposition is often seen in other species having a seasonal reproductive cycle similar to *R. acutus* (Capapé *et al.*, 2006; Ba *et al.*, 2013*a*). Despite this result, further validation of the annual periodicity of the ring pattern observed in *R. acutus* could also be added through techniques such as chemical marking, bomb carbon dating or tag-recapture studies.

RHIZOPRIONODON ACUTUS GROWTH

Multiple growth functions are often calculated and compared in order to characterize adequately the growth of a given species (Cailliet *et al.*, 2006; Katsanevakis, 2006; Diouf *et al.*, 2009). The three growth models used in this study (VBGF, GGF1 and

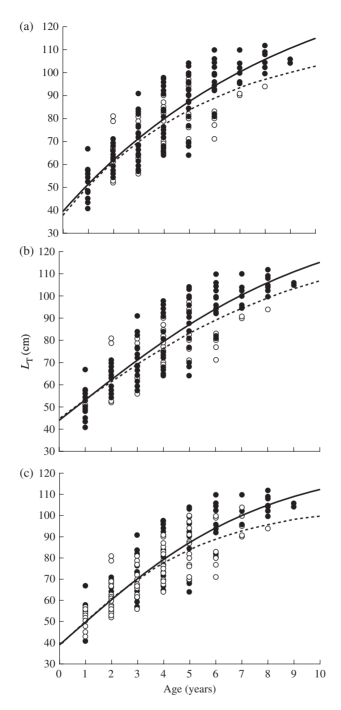


Fig. 6. Growth model curves (a) von Bertalanffy growth function (VBGF), (b) GGF1 and (c) GGF2 of females (\bullet —; n = 267) and males (O—:; n = 156 from observed total length (L_T)-at-age data for *Rhizoprion-odon acutus* in West African waters (see Table I).

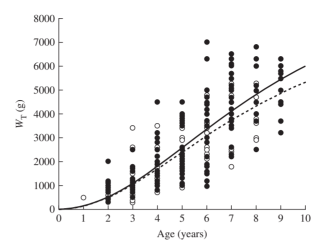


Fig. 7. von Bertalanffy growth modelling in mass of females (\bullet ——; n = 267) and males (\bigcirc ——; n = 156) Rhizoprionodon acutus in West African waters (see Table I).

GGF2) fitted the data well for growth in length and mass. The Gompertz model with an estimated size at birth (GGF2), however, was the best model with no uncertainty as indicated by AIC_c values (Table I). Even if the von Bertalanffy model is most often used to describe fish growth, an increasing number of elasmobranch studies indicated that the Gompertz model might better describe their growth (Mollet *et al.*, 2002; Wintner *et al.*, 2002; Cailliet *et al.*, 2006; Katsanevakis, 2006). The Gompertz model is then commonly used as growth function for Chondrichthyans (Mollet *et al.*, 2002; Braccini *et al.*, 2007) and is then particularly relevant for comparisons. Cailliet *et al.* (2006) recommended using the size at birth in the growth models instead of t_0 , because it can be biologically meaningful. The L_0 values calculated in this study for both sexes were biologically reasonable compared with the published data on size at birth for this species in this area (Capapé *et al.*, 2006; Ba *et al.*, 2013a). The ratio of L_b and estimated L_0 with values <1 for both sexes indicated no severely biased growth estimates as it was shown for other shark species (Pardo *et al.*, 2013).

Based on the growth rate values, *R. acutus* have a relatively slow growth rate along the Senegalese coast. Other studies of *Rhizoprionodon* spp. have estimated a wide range of *K* values, from 0·17 to 1·34 for males and 0·16 to 1·01 for females (Table II). Comparison of *R. acutus* growth rates from this study with others suggested that *R. acutus* is a comparatively slow or intermediate growing species. Growth rates also vary extensively within species depending on local water temperature and productivity (Barker *et al.*, 2005). The *K* values of the GGF2 estimated in this study were different from those estimated in Australia for *R. acutus* and from other species of the genus (Harry *et al.*, 2010). Geographic variation is a possible factor which could account for growth differences between areas as it was documented for other shark species, *Sphyrna tiburo* (L. 1758) (Lombardi-Carlson *et al.*, 2003) and *Carcharhinus limbatus* (Müller & Henle 1839) (Carlson & Sulikowski, 2006). Growth curves change in response to environmental conditions and fishing pressure. *Rhizoprionodon acutus* in West Africa with a slower growth rate, size and age of maturity higher in comparison with the population in Australia (Harry *et al.*, 2010) and all other species of the genus, is likely to become

TABLE II. Growth parameters of Rhizoprionodon species reported in the literature

Species	L_{∞} (cm)	K (year ⁻¹)	n	Authors	Region (or oceanic basin)
Rhizoprionodon	126.5 (M)/139.5 (F)	126-5 (M)/139-5 (F) 0.18 (M)/ 0.17 (F) 156 (M)/ 267 (F)	156 (M)/267 (F)	This study	Senegalese coast
acutus R. acutus	82·1 (M)/86·1 (F)	0.94 (M)/0.63 (F)	127 (M)/38 (F)	Harry <i>et al.</i> (2010)	North-eastern coast of
Rhizoprionodon	78.1	0.3	84	Lessa <i>et al.</i> (2009)	Australia Northern coast of Brazil
talanati Rhizoprionodon	112.9	0.17	134	Lessa <i>et al.</i> (2009)	
porosus Rhizoprionodon	92.5	0.45	215	Parsons (1985)	North central Gulf of
ierraenovae R. terraenovae	108	0.36	20	Branstetter (1987a)	Mexico Laboratory aquaria
R. terraenovae	94	0.85 (M)/0.63 (F)	304	Carlson & Baremore (2003)	Gulf of Mexico
R. terraenovae Rhizoprionodon	98·3 (M)/98·8 (F) 65·2 (M)/73·2 (F)	0.50 (M)/0.49 (F) 1.34 (M)/1.01 (F)	116 (M)/123 (F) 52 (M)/85 (F)	Loefer & Sedberry (2003) Simpfendorfer (1993)	South-eastern U.S.A. North Queensland,
taytori					Austrana

M, males; F, females.

heavily exploited and more sensitive to overexploitation. Carlson & Baremore (2003) have shown a decrease in the size at maturity and an opposite trend in the growth rate of R. terraenovae. In this study, both parameters have shown a downward trend for R. acutus. For Carlson & Baremore (2003), the increase in growth rate of R. terraenovae resulted from a decrease in the competition in the Gulf of Mexico with an increase of food availability; this does not appear to be the case in this study for R. acutus. Rhizoprionodon acutus, similar to the lemon shark Negaprion brevirostris (Poey 1868), might be an exception to slow-growing species, which are found in subtropical and tropical waters (Simpfendorfer et al., 2002). According to those authors, carcharhinid species with slower growth rates spend some or all of their time in temperate waters, whereas those with faster growth rates are more often found in tropical waters. According to Branstetter (1990), slow-growing and coastal species, such as R. acutus, protect their nurseries as it is the case for other sharks along the Senegalese coast (Capapé et al., 2006). Length-selective fishing mortality could explain the observed differences in growth curves (Walker et al., 1998). The decrease of growth rates could result in an artificial selection by the fishing gears (Moulton et al., 1992). Many data have highlighted and reinforced the understanding that species with slower growth rates and later ages of maturity, such as R. acutus, are more susceptible to possible extinction risks (Cortés, 2000; García et al., 2008). Consequently, management measures must be implemented for this species.

The older females observed in this study were 9 years ($L_{\rm T} = 106\,{\rm cm}$) and the older males were 8 years ($L_{\rm T} = 104\,{\rm cm}$). In Australia, maximum ages were younger for females (8·1 years) and for males (4·5 years) (Harry *et al.*, 2010), the individual sizes being smaller in this area. Asymptotic lengths predicted by the GGF2 were higher than the observed maximum lengths for both sexes; this pattern could result from the lack of large specimens in the landings. The 1–3 year-old fish, due to either sampling effects or fishing pressure, dominated the age structure (Fig. 5).

The data on growth in mass of R. acutus are the first available and will be very useful for management of this species. The results of this study suggest that R. acutus tended towards W_{∞} but grew very little in mass at older ages. According to this study, growth gain of females was greater than that of males, confirming the sexual dimorphism for this species (Capapé et al., 2006; Ba et al., 2013a). The variation of asymptotic mass between males and females might be related to the sex-specific preferences in the food-searching sites. As females have larger stomach, this may influence the mass of the animals (Ba et al., 2013a). This feature could also be explained by the need for females to reach a larger size to support embryos because of their viviparity (Walmsley-Hart et al., 1999; Ba et al., 2013a).

CONSEQUENCES FOR R. ACUTUS EXPLOITATION

The results of this study, and particularly the slow estimated growth rate, provide evidence of variable growth rate of this species over its distribution area (Walker et al., 1998). Stevens et al. (2000) reported a possible change in the growth of sharks and batoids resulting in selective fishing mortality. Length-selective fishing mortality with the effect of altering the length-frequency composition of sharks in particular age classes of the wild population could explain the differences in the growth rates of *R. acutus* between locations (Moulton et al., 1992). As observed by Walker et al. (1998), small sharks swim through gillnets but become progressively more vulnerable

to capture as they grow. The gillnets, the main fishing gear used along the Senegalese coast to capture *R. acutus* (Ba *et al.*, 2013*a*), have the effect of culling a higher proportion of slower-growing sharks among the older population (Stevens *et al.*, 2000). Density-dependent factors could cause a change in the growth of a population due to exploitation. This effect could explain the size and age structure of *R. acutus* in this study (Svedäng & Hornborg, 2014). Reduction of the growth rate could also result from differences in fishing intensity, with greater exploitation rates causing smaller sizes (at maturity and maximum) through density-dependent compensation or simply artificial selection for smaller sizes (Olsen *et al.*, 2004). Ba *et al.* (2013*a*) observed a reduction in the size range of *R. acutus* in the study area as it has been shown in other species of the genus (Carlson & Baremore, 2003; Motta *et al.*, 2005). Therefore, precautionary measures should be taken into account for this overexploited species and special management actions should be undertaken to ensure its sustainability (Rose *et al.*, 2001).

A hypothesis of seasonal enrichment of the continental shelf followed by different predator species with an increase of interspecific and intraspecific competition could lead to the low growth rates of R. acutus (Walker et al., 1998; Ba et al., 2013a). This result is highlighted by the numerous proportion of individuals with empty stomachs throughout the year for this species (Ba et al., 2013b). The decreasing food availability for R. acutus along the Senegalese coast could lead starving individuals to approach and be caught by baited fishing gear; this may have consequences on growth. An opposite trend was exhibited for its congener R. terraenovae in the Gulf of Mexico (Carlson & Baremore, 2003). Low growth coefficients for R. acutus reported in this study could provide indices of fisheries overexploitation and might lead to changes in its ecosystem status in this region (Shepherd & Myers, 2005). Musick (1999) noted that species with a K coefficient close to 0.1 are very vulnerable to overexploitation and the value in this study was not very far from 0.1. Even though differences in vertebral preparation samples and in band counting could produce variation in the growth rates, this study showed a low growth rate for R. acutus in relation to other Rhizoprionodon species, especially tropical species (Carlson & Baremore, 2003). Notwithstanding these previous hypotheses to explain low growth rate, it is difficult to evaluate the relative effects of all the potential causes of growth rate reduction.

Age at maturity is also a critical variable for demographic modelling of elasmobranch populations, and therefore for fisheries management. The size at the first sexual maturity was $92.5 \,\mathrm{cm} \, L_{\mathrm{T}}$ for females and $82 \,\mathrm{cm} \, L_{\mathrm{T}}$ for males along the Senegalese coast (Ba et al., 2013a). These values correspond to individuals of 5.8 and 4.8 years for females and males, not far from their maximum ages. Such information can have important effects on stock assessments and fishery models (Musick, 1999; Campana, 2001). Harry et al. (2010) estimated the age at first sexual maturity using the mean-adjusted age at first band at 1.6 years for females and 1.0 years for males. Surprisingly, these values are very far from the present data (5.8 and 4.8 years for females and males, respectively). For other Rhizoprionodon species, male and female R. taylori matured after only 1.0 years (Simpfendorfer, 1993), Rhizoprionodon lalandii at 3.3 years, Rhizoprionodon porosus at 2.6 years (Lessa et al., 2009) and R. terraenovae at 2.6 years (Loefer & Sedberry, 2003). The previous estimations of age at maturity gave younger ages, and the evolution of this age, which is probably linked with overexploitation, should be checked carefully. Life-history traits, such as slow growth rates, mean that many sharks have a low resilience to fishing mortality, being more susceptible to overfishing

than most teleosts and invertebrate species (Musick *et al.*, 2000; Frisk *et al.*, 2001). In addition, *R. acutus* gathers in schools by sex and reproductive state (Ba *et al.*, 2013*a*), so that fishing may deplete large segments of particular age classes or sexes, including reproductively active segments (Heyman *et al.*, 2001). Management of the *R. acutus* fishery may need to incorporate this new data into fishing management practice, also taking into account differences between sexes.

Consequently, more efforts to increase the knowledge of the population biology of *R. acutus* should be made in order to ensure its sustainable management. This study provides the first detailed estimates of age and growth for *R. acutus* in West African waters, and should be used in further stock evaluations in this region.

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References

- Ba, A., Ba, C. T., Diouf, K., Ndiaye, P. I. & Panfili, J. (2013a). Reproductive biology of the milk shark *Rhizoprionodon acutus* (Carcharhinidae) off the coast of Senegal. *African Journal of Marine Science* **35**, 223–232.
- Ba, A., Diop, M. S., Diatta, Y., Dossa, J. & Ba, C. T. (2013b). Diet of the milk shark, *Rhizopri-onodon acutus* (Chondrichthyes: Carcharhinidae), from the Senegalese coast. *Journal of Applied Ichthyology* 1, 1–7.
- Barker, M. J., Gruber, S. H., Newman, S. P. & Schluessel, V. (2005). Spatial and ontogenetic variation in growth of nursery-bound juvenile lemon sharks, *Negaprion brevirostris*: a comparison of two age-assigning techniques. *Environmental Biology of Fishes* **72**, 343–355.
- Bass, A. J., D'Aubrey, J. D. & Kistnasamy, N. (1975). Sharks of the east coast of southern Africa. III. The families Carcharhinidae (excluding *Mustelus* and *Carcharhinus*) and Sphyrnidae. *Oceanographic Research Institute Investigational Report* 33, 1–100.
- Baum, J. K., Myers, R. A., Kehler, D. G., Worm, B., Harley, S. J. & Doherty, P. A. (2003). Collapse and conservation of shark populations in the northwest Atlantic. *Science* 299, 389–392.
- Beamish, R. J. & Fournier, D. A. (1981). A method for comparing the precision of a set of age determinations. *Canadian Journal of Fisheries and Aquatic Sciences* **38**, 982–983. doi: 10.1139/f81-132
- von Bertalanffy, L. (1938). A quantitative theory of organic growth (inquiries on growth laws II.). *Human Biology* **10**, 181–213.
- Braccini, J. M., Gillanders, B. M., Walker, T. I. & Tovar-Avila, J. (2007). Comparison of deterministic growth models fitted to length-at-age data of the piked spurdog (*Squalus megalops*) in south-eastern Australia. *Marine and Freshwater Research* **58**, 24–33.
- Branstetter, S. (1987a). Age and growth validation of new-born sharks held in laboratory aquaria, with comments on the life history of the Atlantic sharpnose shark, *Rhizoprionodon terraenovae*. *Copeia* **1987a**, 291–300.
- Branstetter, S. (1987b). Age, growth and reproductive biology of the silky shark, *Carcharhinus falciformis*, and the scalloped hammerhead, *Sphyrna lewini*, from the northwestern Gulf of Mexico. *Environmental Biology of Fishes* **19**, 161–173.
- Branstetter, S. (1990). Early life-history implications of selected carcharhinoid and lamnoid sharks of the northwest Atlantic. In *Elasmobranchs as Living Resources: Advances in the Biology, Ecology, Systematics, and the Status of the Fisheries* (Pratt, H. L. Jr., Gruber, S. H. & Taniuchi, T., eds), pp. 17–28. *NOAA Technical Report* **NMFS 90.**
- Burnham, K. P. & Anderson, D. R. (2002). *Model Selection and Multimodel Inference: A Practical Information-Theoretic Approach*. New York, NY: Springer.
- Cailliet, G. & Goldman, K. (2004). Age determination and validation in chondrichthyan fishes. In *Biology of Sharks and their Relatives* (Carrier, J. C., Musick, J. A. & Heithaus, M. R., eds), pp. 399–447. San Diego, CA: CRC Press.

- Cailliet, G. M., Smith, W. D., Mollet, H. F. & Goldman, K. J. (2006). Age and growth studies of chondrichthyan fishes: the need for consistency in terminology, verification, validation, and growth function fitting. *Environmental Biology of Fishes* 77, 211–228.
- Campana, S. E. (2001). Accuracy, precision and quality control in age determination, including a review of the use and abuse of age validation methods. *Journal of Fish Biology* **59**, 197–242. doi: 10.1577/1548-8659
- Capapé, C., Diop, M. & N'dao, M. (1994). Observations sur la biologie de la reproduction de dix-sept espèces de Sélaciens d'intérêt économique capturés dans la région marine de Dakar-Ouakam (Sénégal, Atlantique orientale tropicale). *Bulletin Institut Fondamental d'Afrique Noire* 47, 87–102.
- Capapé, C., Diatta, Y., Diop, M., Guélorget, O., Vergne, Y. & Quignard, J. P. (2006). Reproduction in the milk shark, *Rhizoprionodon acutus* (Rüppel, 1837) (Chondrichthyes: Carcharhinidae) from the coast of Senegal (eastern tropical Atlantic). *Acta Adriatica* 47, 111–126.
- Carlson, J. K. & Baremore, I. E. (2003). Changes in biological parameters of Atlantic sharpnose shark *Rhizoprionodon terraenovae* in the Gulf of Mexico: evidence for density dependent growth and maturity. *Marine and Freshwater Research* **54**, 227–234.
- Carlson, J. K. & Sulikowski, J. R. (2006). Do differences in life history exist for blacktip sharks, *Carcharhinus limbatus*, from the United States South Atlantic Bight and Eastern Gulf of Mexico? *Environmental Biology of Fishes* 77, 279–292.
- Castillo-Geniz, J. L., Marques-Farias, J. F., Rodriguez De La Cruz, M. C., Cortés, C. & Cid Del Prado, A. (1998). The Mexican artisanal shark fishery in the Gulf of Mexico: towards a regulated fishery. *Marine and Freshwater Research* **49**, 611–620.
- Chang, W. Y. B. (1982). A statistical method for evaluating the reproducibility of age determination. *Canadian Journal of Fisheries and Aquatic Sciences* **39**, 1208–1210.
- Compagno, L. J. V. (1984). FAO Species Catalogue. 4. Requins du monde: un catalogue annoté et illustré des espèces de requins connues à ce jour. Partie 2. Carcharhiniformes. *FAO Fisheries Synopsis* **125**, 251–655.
- Cortés, E. (2000). Life history patterns and correlations in sharks. *Reviews in Fisheries Science* **8.** 299–344.
- Diouf, K., Guilhaumon, F., Aliaume, C., Ndiaye, P., Do Chi, T. & Panfili, J. (2009). Effects of the environment on fish juvenile growth in West African stressful estuaries. *Estuarine, Coastal and Shelf Science* **83,** 115–125.
- Dulvy, N. K., Fowler, S. L., Musick, J. A., Cavanagh, R. D., Peter, M., Kyne, L. R. H., Carlson, J. K., Davidson, L. N., Fordham, S. V., Francis, M. P., Pollock, C. M., Simpfendorfer, C. A., Burgess, G. H., Carpenter, K. E., Compagno, L. J., David, A., Ebert, C. G., Heupel, M. R., Livingstone, S. R., Sanciangco, J. C., Stevens, J. D., Valenti, S. & White, W. T. (2014). Extinction risk and conservation of the world's sharks and rays. *eLife* 3, 1–34.
- Ferretti, F., Worm, B., Britten, G. L., Heithaus, M. R. & Lotze, H. K. (2010). Patterns and ecosystem consequences of shark declines in the ocean. *Ecology Letters* **13**, 1071.
- Frisk, G. F., Miller, T. J. & Fogarty, M. J. (2001). Estimation and analysis of biological parameters in elasmobranch fishes: a comparative life history study. *Canadian Journal of Fisheries and Aquatic Sciences* **58**, 969–981.
- García, V. B., Lucifora, L. O. & Myers, R. A. (2008). The importance of habitat and life history to extinction risk in sharks, skates, rays and chimaeras. *Proceedings of the Royal Society B* **275**, 83–89.
- Goldman, K. J., Cailliet, G. M., Andrews, A. H. & Natanson, L. J. (2012). Assessing the age and growth of chondrichthyan fishes. In *Biology of Sharks and their Relatives*, 2nd edn (Carrier, J. C., Musick, J. A. & Heithause, M. R., eds), pp. 423–451. Boca Raton, FL: CRC Press.
- Harry, A. V., Simpfendorfer, C. A. & Tobin, A. J. (2010). Improving age, growth, and maturity estimates for aseasonally reproducing chondrichthyans. *Fisheries Research* **106**, 393–403.
- Heithaus, M., Wirsing, A. J. & Dill, L. M. (2012). The ecological importance of intact top-predator populations: a synthesis of 15 years of research in a seagrass ecosystem. *Marine and Freshwater Research* **63**, 1039–1050.

- Henderson, A. C., McIlwain, J. L., Al-Oufi, H. S. & Ambu-Ali, A. (2006). "Reproductive biology of the milk shark *Rhizoprionodon acutus* and the bigeye houndshark *Iago omanensis* in the coastal waters of Oman". *Journal of Fish Biology* **68**, 1662–1678.
- Heyman, W. D., Graham, R. T., Kjerfve, B. & Johannes, R. E. (2001). Whale sharks, *Rhincodon typus*, aggregate to feed on fish spawn in Belize. *Marine Ecology Progress Series* 215, 275–282.
- Huveneers, C., Stead, J., Bennett, M. B., Lee, A. K. & Harcourt, R. G. (2013). Age and growth determination of three sympatric wobbegong sharks: how reliable is growth band periodicity in Orectolobidae. *Fisheries Research* **147**, 413–425.
- Joung, S. J., Liao, Y. Y. & Chen, C. T. (2004). Age and growth of sandbar shark, *Carcharhinus plumbeus*, in northeastern Taiwan waters. *Fisheries Research* **70**, 83–96.
- Katsanevakis, S. (2006). Modelling fish growth: model selection multi-model inference and model selection uncertainty. *Fisheries Research* **81**, 229–235.
- Kvalseth, T. O. (1985). Cautionary note about R². American Statistician **39**, 279–285.
- Lessa, R., Santana, F. M. & Almeida, Z. S. (2009). Age and growth of the Brazilian sharpnose shark, *Rhizoprionodon lalandii* and Caribbean sharpnose shark, *R. porosus* (Elasmobranchii, Carcharhinidae) on the northern coast of Brazil (Maranhão). *Pan-American Journal of Aquatic Sciences* **4,** 532–544.
- Liu, K., Chen, Č., Liao, T. & Joung, S. (1999). Age, growth, and reproduction of the pelagic thresher shark, *Alopias pelagicus* in the northwestern Pacific. *Copeia* **1999**, 68–74.
- Loefer, J. K. & Sedberry, G. R. (2003). Life history of the Atlantic sharpnose shark (*Rhizoprionodon terraenovae*) (Richardson, 1836) off the southeastern United States. *Fishery Bulletin* **101**, 75–88.
- Lombardi-Carlson, L. A., Cortés, E., Parsons, G. R. & Mainire, C. A. (2003). Latitudinal variation in life-history traits of bonnethead sharks, *Sphyrna tiburo* (Carcharhiniformes: Sphyrnidae) from the eastern Gulf of Mexico. *Marine and Freshwater Research* **54**, 875–883.
- McAuley, R. B., Simpfendorfer, C. A., Hyndes, G. A., Allison, R. R., Chidlow, J. A., Newman, S. J. & Lenanton, R. C. J. (2006). Validated age and growth of the sandbar shark, *Carcharhinus plumbeus* (Nardo 1827) in the waters off Western Australia. *Environmental Biology of Fishes* 77, 385–400.
- Mollet, H. F., Ezcurra, J. M. & O'Sullivan, J. B. (2002). Captive biology of the pelagic stingray, *Dasyatis violacea* (Bonaparte, 1832). *Marine and Freshwater Research* **53**, 531–541.
- Motta, F. S., Gadig, O. B. F., Namora, R. C. & Braga, F. M. S. (2005). Size and sex compositions, length–mass relationship, and occurrence of the Brazilian sharpnose shark, *Rhizoprionodon lalandii*, caught by artisanal fishery from southeastern Brazil. *Fisheries Research* **74**, 116–126.
- Moulton, P. M., Walker, T. I. & Saddlier, S. R. (1992). Age and growth studies of gummy shark, *Mustelus antarcticus* Günther, and school shark, *Galeorhinus galeus* (Linnaeus), from southern Australian waters. *Australian Journal of Marine and Freshwater Research* **43**, 1241–1267.
- Musick, J. A. (1999). Ecology and conservation of long-lived marine animals. In *Life in the Slow lane: Ecology and Conservation of Long-Lived Marine Animals* (Musick, J. A., ed.), pp. 1–10. *American Fisheries Society Symposium* **23.**
- Musick, J. A., Burgess, G., Vailliet, G., Camhi, M. & Fordham, S. (2000). Management of sharks and their relatives (Elasmobranchii). *Fisheries* **25**, 9–13.
- Natanson, L. J., Casey, J. G. & Kohler, N. E. (1995). Age and growth estimates for the dusky shark, *Carcharhinus obscurus*, in the western North Atlantic Ocean. *Fishery Bulletin* **193**, 116–126.
- Olsen, E. M., Heino, M., Lilly, G. R., Morgan, M. J., Brattey, J., Ernande, B. & Dieckmann, U. (2004). Maturation trends indicative of rapid evolution preceded the collapse of northern cod. *Nature Communications* **428**, 932–935.
- Pardo, S. A., Cooper, A. B. & Nicholas, K. D. (2013). Avoiding fishy growth curves. *Methods in Ecology and Evolution* **4,** 353–360.
- Parsons, G. R. (1985). Growth and age estimation of the Atlantic sharpnose shark, *Rhizoprion-odon terraenovae*: a comparison of techniques. *Copeia* **1985**, 80–85.
- Ricker, W. E. (1979). Growth rates and models. In *Fish Physiology*, Vol. VIII (Hoar, W. S., Randall, D. J. & Brett, J. R., eds), pp. 677–743. New York, NY: Academic Press.

- Rose, K. A., Cowan, J. H., Winemiller, K. O. & Myers, R. A. (2001). Blackwell Science Ltd Compensatory density dependence in fish populations: importance, controversy, understanding and prognosis. *Fish and Fisheries* 2, 293–327.
- Shepherd, T. D. & Myers, R. A. (2005). Direct and indirect fishery effects on small coastal elasmobranchs in the northern, Gulf of Mexico. *Ecology Letters* **8**, 1095–1104.
- Simpfendorfer, C. A. (1993). Age and growth of the Australian sharpnose shark, *Rhizoprionodon taylori*, from north Queensland, Australia. *Environmental Biology of Fishes* **36**, 233–241.
- Simpfendorfer, C. A., Chidlow, J., McAuley, R. & Unsworth, P. (2000). Age and growth of the whiskery shark, *Furgaleus macki*, from southwestern Australia. *Environmental Biology of Fishes* **58**, 335–343.
- Simpfendorfer, C. A., McAuley, R. B., Chidlow, J. & Unsworth, P. (2002). Validated age and growth of the dusky shark, *Carcharhinus obscurus*, from Western Australian waters. *Marine and Freshwater Research* **53**, 567–573.
- Stevens, J. D. & McLoughlin, K. J. (1991). Distribution, size and sex composition, reproductive biology and diet of sharks from northern Australia. *Australian Journal of Marine and Freshwater Research* **42**, 151–199.
- Stevens, J. D., Bonfil, R., Dulvy, N. K. & Walker, P. A. (2000). The effects of fishing on sharks, rays, and chimaeras (Chondrichthyans), and the implications for marine ecosystems. *ICES Journal of Marine Science* **57**, 476–494.
- Sulikowski, J., Kneebone, J., Elzey, S., Jurek, J., Danley, P., Howell, W. & Tsang, P. (2005). Age and growth estimates of the thorny skate (Amblyrajaradiata) in the western Gulf of Maine. *Fishery Bulletin* **103**, 161–168.
- Svedäng, H. & Hornborg, S. (2014). Selective fishing induces density-dependent growth. *Nature Communications* **5**, 1–6.
- Valadou, B., Brêthes, J. C. & Inejih, C. A. O. (2006). Observations biologiques sur cinq espèces d'Élasmobranches du Parc national du Banc d'Arguin (Mauritanie). *Cybium* **30**, 313–322.
- Walker, T. I., Taylor, B. L., Hudson, R. J. & Cottier, J. P. (1998). The phenomenon of apparent change of growth rate in gummy shark (*Mustelus antarcticus*) harvested off southern Australia. *Fisheries Research* **39**, 139–163. doi: 10.1016/S0165-7836(98)00180-5
- Walmsley-Hart, S. A., Sauer, W. H. H. & Buxton, C. D. (1999). The biology of the skates *Raja* wallacei and *R. pullopunctata* (Batoidea: Rajidae) on the Agulhas Bank. South Africa. *South African Journal of Marine Science* **21**, 165–179.
- Wintner, S. B., Cliff, G., Kistnasamy, N. & Everett, B. (2002). Age and growth estimates for the Zambezi shark, *Carcharhinus leucas*, from the east coast of South Africa. *Marine and Freshwater Research* **53**, 557–566.

Electronic Reference

Simpfendorfer, C. A. (2003). *Rhizoprionodon acutus*. The IUCN Red List of Threatened Species. Available at http://www.iucnredlist.org/details/41850/0