

**Condition analysis of the Brazilian sharpnose shark**

*Rhizoprionodon lalandii*: evidence of maternal investment for initial post-natal life

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### 1 INTRODUCTION

Analysing the degree of well-being (condition) of an animal is strategic in evaluating vital processes such as migration, feeding and reproduction. Condition reflects the accumulation of energy for future use to alleviate the effects of scarcity of food resources and the costs of reproductive and migratory activities (Gallagher et al., 2014; Irschick & Hammerschlag, 2014). Despite its importance, studies about the condition of elasmobranchs are scarce compared with teleosts, mostly due to the relative smaller participation of sharks and rays in total fishery production, lack of systematic monitoring of fisheries landings, difficulty in accessing whole specimens and the migratory habits of many species (Castro et al., 1999; Hussey et al., 2009).

Fish condition indicators are traditionally calculated based on the length–mass relationship (Bolger & Connolly, 1989; Froese, 2006; Jakob et al., 1996) as the condition factor and metrics that relate mass of organs with total mass (Hussey et al., 2009), such as liver (hepato-somatic index) and gonads (gonado-somatic index). Recently, biochemical analysis (e.g. fatty acids) and the relationship between circumference and body length measures have also been used for large sharks (Gallagher et al., 2014; Irschick & Hammerschlag, 2014). In elasmobranchs, condition metric fluctuations have usually been related with energetic costs during the reproductive cycle (Allen & Cliff, 2000; Capapé & Reynaud, 2011; Estalles et al., 2008; Fermin & Bashirullah 1984; Grijalba-Bendeck et al., 2008; Hoffmayer et al., 2006; Machado et al., 2001; Peres & Vooren, 1991), migration (King, 1984), responses to variations in food availability (Oddone & Amorim, 2007) and other factors that influence the well-being of organisms (Braccini & Chiaramonte, 2002; Hoffmayer et al., 2006; Parsons & Hoffmayer, 2005).

Placental viviparity is usually regarded as the most efficient reproductive strategy among elasmobranchs due to the close relationship of nutrient demand and support between the embryos and the mother (Parsons et al., 2007). According to Hussey et al. (2010), maternal investment extends to postnatal life of two shark species with biennial reproductive cycles, the dusky shark *Carcharhinus obscurus* (LeSueur 1818) and spinner shark *Carcharhinus brevipinna* (Valenciennes 1839), which have enlarged livers in the new-born. This condition would assist with the beginning of foraging activities and increase survival probability during the first weeks and months of life.

The sharpnose sharks, genus *Rhizoprionodon* Whitley 1929 are represented worldwide by at least seven small coastal placental species (up to 1,780 mm total length; \(L_T\)) with annual reproductive cycles and concurrent ovarian and gestation cycles (Compagno, 1984). The
Brazilian sharpnose shark *Rhizoprionodon lalandii* (Valenciennes 1839) is the most abundant coastal shark species in south-eastern Brazil, comprising approximately 60% of total shark landings in artisanal fisheries (Motta et al., 2005). In the same area, the species attains a maximum size of 800 mm TL and sexual maturity at 590–650 mm TL (Motta et al. 2007). Moreover, three seasonal occurrence patterns related to size classes are recognized: August–September neonates are most frequent; November–March, juveniles are most common; April–July, adults are the most numerically abundant (Motta et al. 2007). Data on condition of coastal placental sharks with annual reproductive cycles. somatic index) by length classes and seasons as models for small artisanal fishery in south-eastern Brazil. The aim of the present study is the total body mass.

In this context, the present work reports an analysis of condition of *R. lalandii* from a long-term data set from the sharks caught by the artisanal fishery in south-eastern Brazil. The aim of the present study was to assess two condition metrics (condition factor and hepato-somatic index) by length classes and seasons as models for small coastal placental sharks with annual reproductive cycles.

## 2 | MATERIALS AND METHODS

### 2.1 | Sampling

Specimens of *R. lalandii* (*n* = 8,268) were obtained weekly from artisanal fisheries operating along the south-central coast of São Paulo State, south-eastern Brazil and landed at Pescadores Beach, Itanhaém municipality (24° 11′ S; 46° 48′ W; Figure 1), between January 1997 and December 2003. The fishing fleet consisted of small motorized boats (4–10 m long) using 1,500 m long monofilament gillnets with stretched-mesh sizes of 7, 12 and 14 cm. The nets were set c. 5–20 km from the shore in waters between 5 and 35 m deep. All landed sharks were sexed, measured (TL, mm) and weighed (total body mass, *MT*, g). After evisceration by fishermen still on the beach for commercial purposes, the livers were weighed in g (*n* = 2,071).

### 2.2 | Relative growth of liver and condition metrics

The relative growth of the liver and hepato-somatic index (*I*s) were studied for 893 females and 1,178 males. Relative growth of the liver was analysed using the allometric equation *M*ₜ = *a L*ₜ⁻*b*, where *M*ₜ is the total liver mass, *L*ₜ total length and *a* and *b* are parameters estimated by least-squares fitting method after log-transformation of *M*ₜ and *L*ₜ data. The hepato-somatic index (*I*s) was calculated by the expression *I*s = 100*M*ₜ*MT*⁻¹, where *M*ₜ is the total liver mass and *MT* is the total body mass.

Fulton’s condition Factor (*K*) was calculated for 3,962 females and 4,306 males by the expression *K* = *M*ₜ(*L*ₜ³)⁻¹ (Fulton, 1904), where *M*ₜ is the total body mass, *L*ₜ total length and *b* is the regression coefficient of the length–mass relationship *M*ₜ = *a L*ₜ⁻*b*. In the present study values for *b* of 3.22 and 3.13 were used for females and males, respectively (Motta et al., 2014). Linear regressions of *I*s and *L*ₜ were calculated separately for females and males and then compared by analysis of co-variance (ANCOVA).

Time-series of *K* and *I*s by quarter of the year and length classes (50 mm) were generated for females and males using generalized linear models (GLM). Like other predictive models, GLMs allow removal of unsystematic variations caused by some components (i.e. variables) and identify systematic trends or signals in the data. The basic GLM structure used in this study was: *μ* = *Xβ*, where *X* is the matrix of explanatory variables, *β* is the vector of explanatory variables and *μ* is the expected values for response variables.

The GLMs fitted to both response variables (*K* and *I*s) in both time-series scenarios (quarter and quarter of the year) used year, quarter of the year, season and length class (50 mm range) as independent class variables and the *L*ₜ as an independent continuous variable. Exclusively for *I*s, we considered in the modelling process the inclusion of *K* as a continuous independent variable. We chose not to use *I*s, while modelling *K*, to avoid a significant reduction in the number of observations of this variable (significantly higher than *I*s in this study: 8268 vs. 2071). Models were then refined by interactively selecting the most appropriated combination of explanatory variables according to their statistical significance.

## 3 | RESULTS

### 3.1 | Relative liver growth

The analysis of the relationship between *M*ₜ and *L*ₜ (Figure 2) showed a tendency for continuous growth until the sizes at first maturity, after which a great dispersion of data was observed. The co-variance analysis revealed significant differences between *M*ₜ and *L*ₜ relationships of females and males for *R. lalandii* (between the regression coefficients *F*₁,₂₀₇₀ = 4.154, *P* < 0.001, between the intercepts *F*₁,₂₀₇₀ = 4.054, *P* < 0.001).

### 3.2 | Condition metrics analysis

Significant differences between the length classes for *I*s (Figure 3 and Tables S1 and S2) and *K* (Figure 4 and Tables S3 and S4) were detected. Both metrics showed high values for length classes of neonates (300–350 mm *L*ₜ), decreasing to the classes between 400 and 500 mm *L*ₜ. From these length classes, a gradual increase of *I*s and *K* were observed, especially for the females (Figures 3 and 4).

The variables selected to build the model structure that best explains each variable response of this study (*K* and *I*s), as well as the estimated coefficients and their respective indices of statistical significance are available in the Supporting Information (Tables S1–S5). In general, the best model for both *K* and *I*s comprised only quarter of the year and length class (Tables S1–S4). However, considering that in the analysis of *I*s the variable *K* was also tested through GLM analysis, we found the same significance during the modelling process as shown in Tables S1 and S2.

Due to high co-linearity, our models did not converge when seasons (summer, autumn, winter and spring) and quarter of the year (1997 summer, 1997 autumn et seq.) were placed together. Hence, we additionally performed a sensitivity analysis of *K* and *I*s in relation to
season only (Figure 5 and Table S5). In these additional analyses, there were significant seasonal differences for $I_{he}$ (Figure 5 and Table S5). Low $I_{he}$ values were observed during the summer for both sexes. In other seasons, high equivalent values were recorded in the winter–spring period for females while the highest value for males was observed in the autumn. Regarding $K$, (Figure 5 and Table S5), low values for both sexes were found in the spring while high equivalent values were observed in the other seasons, except in the summer for females.

According to GLM-derived time series of seasonal mean values of $K$, the spring presented the lowest value in four of the 7 years: 1998, 1999, 2001 and 2003, in both sexes (Figure 6). The significant differences detected between analysed factors indicated high interannual variability in the condition of the species (Tables S3 and S4). The GLM-derived time series of seasonal mean values of $I_{he}$ (Figure 7) did not present a clear trend, possibly due to low sample size, mainly from the year 2001. Nevertheless, significant differences were also detected (Tables S1 and S2) in a small fraction of the analysed factors when compared with $K$ analyses.

4 | DISCUSSION

Given the difficulties in obtaining data on commercially caught sharks, the present study represents one of the largest and most comprehensive (8,268 individuals) on the condition metrics of a shark species. The discontinuities demonstrated in development of the liver after attaining sexual maturity may be explained by the presence of individuals at different stages of the reproductive cycle in the sample, indicating the participation of the liver in the reproduction of the species. The performance of liver reserves in energy-consuming activities, such as reproduction, has already been documented for other elasmobranchs (Hoffmayer et al., 2006; Hussey et al., 2009; Lteif et al., 2016; Peres & Vooren, 1991; Pethybridge et al., 2014). Some authors have emphasized the importance of the reserves for vitellogenesis and gestation of females (Gallagher et al., 2014; Peres & Vooren, 1991; Rochowski et al., 2015), including $R. lalandii$ in northern Brazil, where Lessa (1988) recorded a doubling of liver mass in females at the end of gestation. In the present study, discontinuity of liver growth after...
sexual maturity was observed, with both sexes showing significant differences, although they were more prominent for females.

The high condition observed for neonates, detected in both K and I, was associated with the possible transfer of energy reserves from the mother to the offspring; here, the I reaches 6–8% for females and males, respectively. Evidence of maternal investment for postnatal life has also been reported for two viviparous shark species with biennial reproductive cycles, C. obscurus and C. brevipinna (Hussey et al., 2010). According to these authors, this strategy maximizes the chances of survival in the first few months of life, during the development of foraging skills of the individuals. In R. lalandii, the investment is relevant because of the small birth size and the shared use of the same nursery area with at least four larger shark species (Gadig et al., 2002; Motta et al., 2005).

The liver reserve depleted in 400–500 mm length classes of juveniles, reproductive cycles, C. obscurus and C. brevipinna (Hussey et al., 2010). According to these authors, this strategy maximizes the chances of survival in the first few months of life, during the development of foraging skills of the individuals. In R. lalandii, the investment is relevant because of the small birth size and the shared use of the same nursery area with at least four larger shark species (Gadig et al., 2002; Motta et al., 2005). The liver reserve depleted in 400–500 mm length classes of juveniles,

**FIGURE 2** Relationship between liver mass (M_H) and total length (L_T) of (a) female (y = 0.0000002x^{3.0}, R^2 = 0.85, P < 0.001, n = 893) and (b) male (y = 0.0000003x^{2.89}, R^2 = 0.85, P < 0.001, n = 1,178) *Rhizoprionodon lalandii*

**FIGURE 3** Trends in generalized linear model standardized mean (±95% C.I.) hepato-somatic indices (I_H) for total-length (L_T) classes of (a) female and (b) male *Rhizoprionodon lalandii*
where the $I_H$ had the lowest value in the life cycle (c. 4% for females and 6% for males). Coincidentally, vertebrae sectioned from specimens of these size classes show additional band pairs after the birthmark, which may indicate physiological stress at this life stage (F. S. Motta, pers. comm.). In Hawaii, natural mortality of juvenile scalloped hammerhead shark *Sphyrna lewini* (Griffith & Smith 1834) was associated with poor condition during the first months of life (Duncan & Holland, 2006). The hypothesis of maternal investment for postnatal life may have been little studied due to the difficulty of access to neonates in previous condition investigations. Most studies do not cover this age group (Andrade et al., 2008; Hoffmayer et al., 2006; Peres & Vooren, 1991), either because sampling did not occur in nursery areas or because of the selectivity of the fishing gear.

Motta et al. (2007) showed that juvenile *R. lalandii* predominate between November and March, with the transition between neonates and juveniles occurring in October. Thus, the low value of $K$ during spring may be evidence of the directing of energy obtained from feeding for corporal growth of young individuals that do not yet invest in reproduction. During summer, $K$ values are high once more, although $I_H$ shows the lowest value in this season, possibly reflecting an investment in gonad maturation from the accumulated energy in the liver. According to Motta et al. (2007), mating in the study area takes place between April and June for first-maturing females and between July and September for post partum females (southern hemisphere autumn and winter respectively), where both condition metrics show high values, indicating the level of energetic demand required for this critical phase of the life cycle. A similar pattern was recorded for the milk shark *Rhizoprionodon acutus* (Rüppell 1837) off the Senegal coast (Ba et al., 2013). On the other hand, Machado et al. (2001) associated the highest $I_H$ values of the Caribbean sharpnose shark *Rhizoprionodon porosus* (Poey 1861) in north-east Brazil with the period before reproduction, which is the opposite of that observed in the present study.

Regarding $K$ analyses for the 7 years sampled, the great magnitude of variation is highlighted in 1998, with one of the highest and the lowest values within the whole set of samples, in autumn and spring respectively. The 1997–1998 the El Niño phenomenon was one of the strongest events ever recorded in Peruvian waters (Garcia & Vieira, 2001), with worldwide effects. In south-eastern Brazil, low salinity in estuaries, increased rainfall and increased water surface temperature occurred between July 1997 and September 1998 (Garcia & Vieira, 2001). In this same region, significant changes in species composition and patterns of diversity were found in the years of intense El Niño (1983–1984 and 1997–1998), with marine fish visiting estuaries less often (Garcia & Vieira, 2001; Paes & Moraes, 2007).

The influence of climatic phenomena on reproduction and feeding, prolonging the spawning period and causing weight loss, has already been observed for teleost species (Dioses et al., 2002; Martinez et al. 1985; Santander & Zuzunaga, 1984), but studies of effects on elasmobranchs are still limited. One study on the effect of El Niño on the condition of Peruvian hake *Merluccius gayi peruanus* (Guichenot 1848) showed that during the phenomenon condition values were low, mainly for the year 1998 (Ballón et al., 2008). Otherwise, in Greenland, the condition of the Atlantic cod *Gadus morhua* L. 1.758 was positively related to the increase in water temperature (Lloret & Rätz, 2000). For the Atlantic sharpnose shark *Rhizoprionodon terraenovae* (Richardson 1836), although not during an El Niño, decrease of condition related to increase of water temperature and consequent increase in oxygen consumption (Hoffmayer et al., 2006; Parsons & Hoffmayer, 2005) was recorded. The authors suggested that the sharks should increase their feeding intensity or mobilize energy reserves to supplement the low
FIGURE 5  Trends in generalized linear model standardized mean (±95% C.I.) (a), (b) hepato-somatic index ($I_h$) and (c), (d) Fulton’s condition factor ($K$) by seasons for (a), (c) female and (b), (d) male of *Rhizoprionodon lalandii*. 

FIGURE 6  Trends in generalized linear model standardized mean (±95% C.I.) Fulton’s condition factor ($K$) between January 1997 and December 2003 for (a) female and (b) male *Rhizoprionodon lalandii*. 
FIGURE 7  Trends in generalized linear model standardized mean (±95% C.I.) hepato-somatic index ($I_{HS}$) between January 1997 and December 2003 for (a) female and (b) male Rhizoprionodon lalandii

classification. In the present study, summer, autumn and winter seasons of 1998 correspond to a period of increasing water temperature (Garcia & Vieira, 2001) and R. lalandii exhibit high condition values, which may indicate that the species was able to maintain its well-being even under such circumstances. On the other hand, after the El Niño effects in the spring of 1998, its condition decreased drastically. In fact, more detailed studies are necessary to establish the relationship between the El Niño phenomenon and elasmobranch condition.

It was possible to describe seasonal variations in the condition of R. lalandii, despite the annual reproductive cycle with concurrent vitellogenesis and gestation. In some periods the interpretation of the results was difficult because of the occurrence of specimens at different life-cycle stages. The study also highlights the relevance of the liver in reproduction, accumulating energy for the beginning of postnatal life in neonates and for gonad maturation in adults.

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AUTHOR CONTRIBUTIONS

J.T.C. contributed with ideas, data analysis and manuscript preparation; R.R.P.B. contributed with the data analysis improvement using generalized linear model (GLM) and manuscript revision; F.S.M. contributed with delineating, data collection and data analysis and manuscript preparation.

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