

Occupancy dynamics of South American Sea-Lions in Brazilian Haul-outs

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Abstract

Along the Brazilian coast only two haul-outs of South American sea lions (*Otaria flavescens*) are known: Ilha dos Lobos and Molhe Leste, both located in the southernmost state of Brazil, Rio Grande do Sul. Most sea lions observed in these haul-outs are adult and sub-adult males. It is supposed that the species' presence in these areas is due to food supply and absence of parental assistance by males. This study analysed the use of these haul-outs by *O. flavescens* between 1993 and 2002 based on counting data of observed individuals. Bayesian generalised linear mixed models were used to evaluate differences in abundance between areas, long term trends and seasonal patterns. Results showed that for *O. flavescens* abundance had a long term trend of increased average occupancy over the study period, with seasonal variation reaching the highest within-year value in August (Ilha dos Lobos) and October (Molhe Leste). The novel application of this powerful statistical modelling approach resulted in a useful tool to quantify occupancy dynamic.

Keywords: *Otaria flavescens*, Bayesian generalised linear mixed models, covariates, abundance.

Dinâmica de Ocupação do Leão-Marinheiro-do-Sul no Brasil

Resumo

Ao longo da costa do Brasil apenas duas colônias não-reprodutivas de leões-marinheiros-do-sul (*Otaria flavescens*) são conhecidas: Ilha dos Lobos e Molhe Leste, ambas localizadas no estado do Rio Grande do Sul. A maioria dos leões-marinheiros observados nestas colônias são machos adultos e sub-adultos. Supõe-se que a presença da espécie nestas áreas está relacionada ao forrageamento e a ausência de cuidado parental pelos machos. Este estudo analisou a dinâmica de ocupação e abundância de *O. flavescens* nas colônias não-reprodutivas entre 1993 e 2002, baseado em uma série temporal de dados de contagens de indivíduos. Modelos lineares generalizados mistos Bayesianos foram usados para avaliar diferença na abundância entre áreas, tendência de uso em longo prazo e padrões sazonais. Os resultados mostram que a abundância de *O. flavescens* variou sazonalmente, atingindo pico intra-anual em agosto (Ilha dos Lobos) e outubro (Molhe Leste), acompanhado de um aumento da ocupação média dos refúgios ao longo do período de estudo. A nova aplicação desta poderosa forma de modelagem estatística mostrou-se útil para quantificar a dinâmica de ocupação.

Palavras-chave: *Otaria flavescens*, modelos lineares generalizados mistos Bayesianos, covariáveis, abundância.

1. Introduction

South American sea lions *Otaria flavescens* (Shaw, 1800) are widely distributed, occurring along approximately 10,000 km of the South American Atlantic and Pacific coasts from Torres (Brazil) (Vaz-Ferreira, 1981) to Cape Horn (Chile) and from Cape Horn to Zorritos (Peru) (Cappozzo, 2002). The species' distribution is characterised by features which allow breeding and for-

aging activities, such as solid substrate and food availability (Bowen et al., 2009; Cappozzo and Perrin, 2009).

The pinnipeds' presence far from their usual breeding areas is often the result of erratic movements. However, some species like *O. flavescens*, display frequent and seasonal movements (Rosas et al., 1994; Messias et al., 1994, Simões-Lopes, 1995). They are present in the Argentinean harbours of Mar del Plata (Rodríguez, 1990)

and Puerto Quequén (Westergaard et al., 1979) where they form stable colonies. In these haul-outs, individuals dwell most part of the year with displacement to rookeries only during the breeding season. Mating occurs from December to January with a few births in February. Hence, the breeding dynamics of the species is characterised by an intra-annual peak (Vaz-Ferreira, 1981; Campagna, 1985).

Similar colonies are also found in two haul-outs on the coast of Rio Grande do Sul state (RS), Brazil: Ilha dos Lobos (29°20' S, 52°06' W) and Molhe Leste (32°11' S, 52°04' W) (Vaz-Ferreira, 1981; Rosas et al., 1994) (Figure 1). It is presumed that individuals present in these sites come from the rookeries in Uruguay in search of food (Castello and Pinedo, 1977).

Most sea lions found in both areas are adult and sub-adult males (Rosas et al., 1994; Silva, 2002). Adult males, due to the absence of parental care, move away from the rookeries to return in the next breeding season (Rosas et al., 1994). Females divide their time between foraging trips and fasting periods at nursing, therefore presenting more limited movements (Vaz-Ferreira, 1981; Campagna et al., 2001). Preliminary results of Giardino et al. (2009) in a capture-recapture study suggested that males with high site-fidelity to the haul-out of Puerto Quequén in winter developed a particular strategy of mating that kept them connected with rookeries of Uruguay and Patagonia.

According to Rosas et al. (1994), young individuals have less swimming skills and strength than older animals, which limit their movements to shorter distances

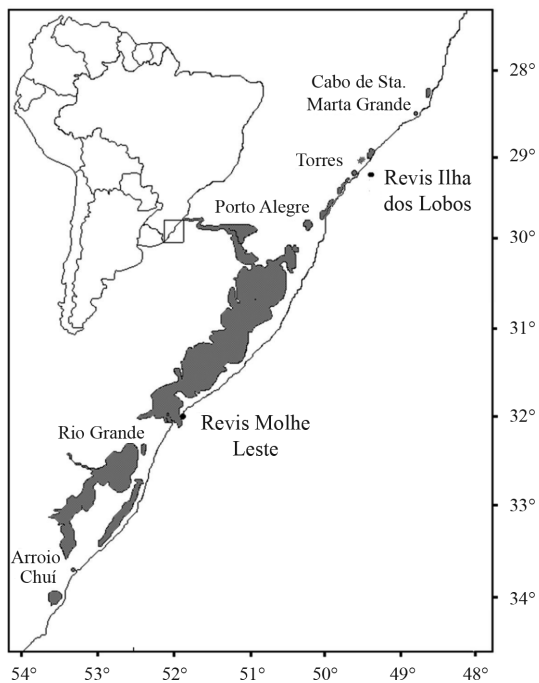


Figure 1 - Study area: the coastline of Rio Grande do Sul, Brazil, between Torres and Arroio Chuí highlighting the Ilha dos Lobos REVIS and Molhe Leste REVIS.

and explain the small number of young animals (≤ 3 years of age) found in southern Brazil during winter months.

After heavy exploitation of sea lions in the first half of the twentieth century, protection measures aiming at population recovery were established. Meanwhile, the abundance of populations in Argentina is increasing (Dans et al., 2004). However, population abundances in the Malvinas Islands and in Uruguay seem to be decreasing (Thompson et al., 2005; Páez, 2005). Therefore, a better understanding on population dynamics is important to design effective conservation strategies.

Based on the possible Uruguayan origin of the sea lions occurring in southern Brazil and the availability of a considerable database of animal counting accumulated over a decade, we analysed the occupancy dynamics and abundance trends of *O. flavescens* in both haul-outs of Rio Grande do Sul, Brazil.

2. Materials and Methods

2.1. Study area and data collection

The coast of Rio Grande do Sul comprises a 618 km long stretch of sandy beach that is interjected only by the Tramandaí River mouth, on the north coast, and by the Patos Lagoon Estuary mouth, on the south coast (Figure 1).

The Patos Lagoon Estuary has a 20 km long entrance channel which is fixed by a 4 km long granite dike into the ocean (Seeliger et al., 1997), known as “Molhe Leste (ML)”. The structure which was built to enable a safe navigation also became a suitable resting place for South American sea lions (Rosas et al., 1994). The other site, named “Ilha dos Lobos (IL)”, is the only coastal island of Rio Grande do Sul deriving from the Mesozoic era basaltic spill. This island is located 2 km from the beach at Torres and has an area of 1,700 km² comprising a levelled ground (Leinz and Amaral, 1966, Vieira, 1984). Both areas are conservation units named “Refúgio da Vida Silvestre” (REVIS) and legally protected by Brazilian federal laws.

Between January 1993 and December 2002, 188 surveys were conducted comprising both locations. Monitoring activities consisted of one to four visits per month. Whenever possible, both sites were surveyed the same day and hour. Due to limitations in logistic and financial support, from ten years of monthly data collection, visits to ML and IL were not possible in 12 and 39 months, respectively.

In most pinniped populations, one cannot assume that all individuals are available at the time of local count (Buckland and York, 2002). If ignored, this fact will result in underestimation due to availability bias. However, since at ML and IL the maximum number of individuals occurs in the early morning hours, with a decrease during the day to increase again at dawn (Rosas et al., 1994; Sanfelice et al., 1996), all field counts were set for 8:00 am.

Visits to both sites were made by boat, with the exception at ML on days with strong winds from the southern quadrant when the animals rested at the dike's central interspace, less visible from boats. On these occasions visits were made by walking the dike from land. We assumed same sighting efficiency for both approaches and used the data without distinction. Animals were counted by naked eye in one path transect with approximate duration of 15 min, simultaneously by two observers to ensure that animals were neither missed nor double counted. The animals in the water and near to the boat were counted as well.

2.2. Data analysis

We defined as the response variable $y_{[i]} = \text{"maximum number individuals observed at a given REVIS in month } i\text{"}$, ($i = 0, 1, 2, \dots, n$), and considered it to follow a Poisson distribution with mean $\lambda_{[i]}$.

We further assumed a linear relation between $\log(\lambda_{[i]})$ and the explanatory variables 'Site', 'Month', 'Month²' and 'Year'. The only categorical variable 'Site' was specified as:

$$x_{0[i]} = 1, \text{ for "Molhe Leste" (ML)}$$

$$x_{0[i]} = 0, \text{ for "Ilha dos Lobos" (IL)}$$

The other covariates were numerical and, for computational convenience, rescaled to have mean zero and standard deviation equal to one.

The hierarchical structure of the model was as follows:

$$y[i] \sim \text{Poisson}(\lambda_{[i]})$$

$$\log(\lambda_{[i]}) \sim \text{Normal}(\mu_{[i]}, \sigma)$$

$$\mu_{[i]} = \beta_0[1]x_{0[i]} + \beta_0[2](1 - x_{0[i]}) + \beta_1x_1 + \beta_2x_2 + \dots + \beta_nx_n$$

where β_0 is the intercept and $\beta_1, \beta_2, \beta_n$ are the coefficients of the explanatory variables.

Since data $y_{[i]}$ may show extra-variation characterised as an over-dispersed Poisson, we added a random effect with standard deviation σ to deal with this situation. When $\sigma = 0$ we fall back to a standard General Linear Model (GLM) or Poisson regression model. For $\sigma > 0$ we have a General Linear Mixed Models (GLMM) (Faraway, 2006). Over-dispersion is often addressed by considering only fixed effects ($\sigma = 0$) but using a negative binomial distribution for $y_{[i]}$ instead (Kinas et al., 2005). However, the choice taken here is better suited to explore the intended hierarchical model structure (Royle and Dorazio, 2008).

Twelve models with different combinations of explanatory variables, including interaction terms between 'Month' or 'Month²' with 'Site', and with or without random effect were fitted to the data. Model parameters were estimated through Bayesian analysis (Gelman et al., 1995) using Markov chain Monte Carlo (MCMC) algorithms to obtain the posterior distribution (McCarthy,

2007). We used *OpenBUGS* in combination with *R software* through libraries *R2WinBUGS* and *BRugs* (Sturtz et al., 2005; Thomas et al., 2006) to run the MCMC. Priors were chosen to be weakly informative. For each β_j this prior had a normal distribution with mean zero and variance equal to 10^3 . The prior for the over-dispersion parameter σ was a uniform distribution between 0 and 10.

Models were compared by the Deviance Information Criterion (DIC) (Spiegelhalter et al., 2002).

$$DIC = \hat{D} + 2p_D,$$

where \hat{D} is the expected posterior deviance and p_D the effective number of estimated parameters. In hierarchical models the effective number of parameters can be much smaller than the actual number of parameters due to the build-in dependence structure among parameters. Furthermore, models for which differences in DIC and the DIC of the best fit (ΔDIC) are up to about 10 cannot be ranked with confidence (Spiegelhalter et al., 2002). Therefore, to choose among models within this range of ΔDIC , considerations about the descriptive capacity of ecological important features should be taken into account as well. We used this approach here. A Bayesian residual analysis was also performed on the chosen model, based on the Pearson residuals (Dobson, 2002).

$$r_{[i]} = \frac{y_{[i]} - e^{\lambda_{[i]}}}{\sqrt{e^{\lambda_{[i]}}}}$$

where $y_{[i]}$ is the individual's counts and $e^{\lambda_{[i]}}$ is the mean predict abundance.

Within the Bayesian context we can obtain the posterior distribution for each Pearson residual and calculate the corresponding 95% posterior probability intervals. Only intervals which exclude zero indicate a bad fit. This objective residual check is unique to a Bayesian residual analysis and facilitates the identification of potential outliers.

Occupation probability $\phi_{[i]}$ is defined as the probability that at least one individual is present at month i (i.e. $y_{[i]} > 0$) and is therefore a function of the predict abundance $\lambda_{[i]}$:

$$\phi_{[i]} = P(y_{[i]} > 0) = 1 - P(y_{[i]} = 0)$$

$$\phi_{[i]} = 1 - e^{-\lambda_{[i]}}$$

Using this formula and the posterior distributions of each $\lambda_{[i]}$ over the period 1993 to 2002, posterior distributions and 95% probability intervals for occupation probabilities were also obtained.

3. Results

The number of surveys per season is summarised in Table 1. The total number of counted individuals per survey varied from 0 to 121.

All posterior distributions were obtained with a Markov chain of 31,000 values, a burn-in period of 10,000 values and thinning of 2, resulting in a simulated

Table 1 - Number of surveys per season to Molhe Leste (ML) and Ilha dos Lobos (IL), southern Brazil, from January 1993 to December 2002.

Surveys/seasons	Summer	Autumn	Winter	Spring
ML	37	43	47	45
IL	29	31	32	42

sample of 10,500 values. Statistical diagnostic tools showed good convergence properties of the chains in all models and were omitted. Posterior means for the parameters in all proposed models are given in Table 2.

By adding covariates the DIC decreased indicating a better fit. However, strong downward changes in DIC values occurred consistently whenever, for a given model, the over-dispersion parameter σ was added, as can be seen when comparing models 1, 3, 4, 6 and 7 with 1d, 3d, 4d, 6d and 7d, respectively. This is a strong evidence that data are over-dispersed and that a mixed model is appropriate. According to DIC, the best fit (model 6d) included covariates ‘Month’, ‘Year’, ‘Month²’ and over-dispersion σ . However, model 7d with month-site interaction includes important ecological features which are helpful to distinguish the abundance oscillation between sites. For being acceptably close to the best model (i.e: Δ DIC = 11) we chose model 7d for the remaining analysis.

The high posterior probability $P(\beta_0[1] > \beta_0[2]) = 0.996$ is strong evidence that the site-effect favours larger average numbers at IL in comparison to ML (Bayes fac-

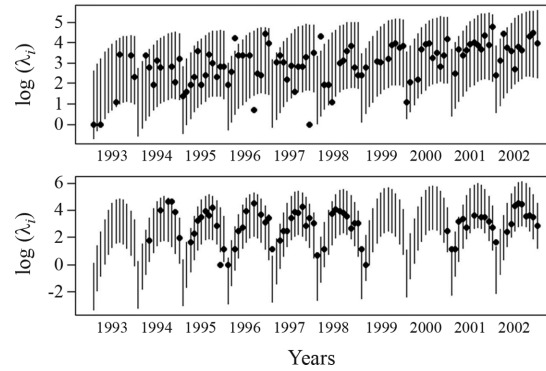


Figure 2 - Monthly 95% probability intervals for the logarithm of expected abundance of *Otaria flavescens* (vertical lines) and the logarithm of observed counts (points) at Molhe Leste REVIS (above) and Ilha dos Lobos REVIS (below). Model 7d.

tor (BF) = 249 indicating that this hypothesis is 249 times more probable than its alternative). In absolute values, this difference represents about 18 individuals (95% probability interval 5.5 to 32.6).

Since the quadratic term (Month²) (Table 2) is smaller than zero (the posterior 95% probability intervals do not include zero) there is statistical evidence that abundance increases from January up to a maximum. This maximum is reached at different times in both sites, in August (winter) and October (early spring) for IL and ML, respectively. After reaching its intra-year maxima, abundance at both sites behave differently. While at IL it

Table 2 - Models with different parameters combinations β_0 ([1] = ML or [2] = IL), β_1 (Month), β_2 (Year), β_3 (Month²) and σ (over-dispersion) ranked in decreasing order of DIC (exception 7(d)). Values are posterior means.

Models	Parameters						DIC
	β_0		β_1	β_2	β_3	σ	
	ML	IL	Month	Year	Month ²	Over-dispersion	
1	3.282	3.242	-	-	-	-	5035
2	3.247	3.208	-	0.263	-	-	4701
3	3.211	3.166	0.386	-	-	-	4341
4	3.175	3.133	0.385	0.261	-	-	4015
5	3.546	3.526	0.534	-	-0.435	-	3848
6	3.499	3.485	0.529	0.252	-0.422	-	3552
7	3.241	3.841	0.371 (ML) 1.010 (IL)	0.255	-0.061 (ML) -1.217 (IL)	-	2936
1d	2.777	2.523	-	-	-	1.334	1210
3d	2.767	2.501	0.697	-	-	1.136	1209
4d	2.762	2.519	0.694	0.424	-	1.059	1209
6d	3.382	3.157	0.728	0.397	-0.642	0.921	1208
7d(*)	3.045	3.652	0.539(ML) 1.012(IL)	0.392	-0.240(ML) -1.111(IL)	0.802	1219

decreases, at ML high abundance is maintained until December.

The posterior 95% probability intervals (Pr.I.) of $\log(\lambda_{[i]})$ obtained for model 7d and the corresponding observed log-counting $\log(\lambda_{[i]})$ are further shown for ML and IL (Figure 2). The quadratic relation to abundance (Figure 3) and the marked seasonal trend (Figures 2), which is different for each site, are expressions of the Month-Site interaction.

A positive coefficient for 'Year' (Table 1, Figure 2) indicates a long-term increasing trend of abundance for both sites.

The residual analysis for model 7d is shown in Figure 4. Points indicate the residuals and vertical lines are the 95% Pr.I. Intervals above the line indicate months with

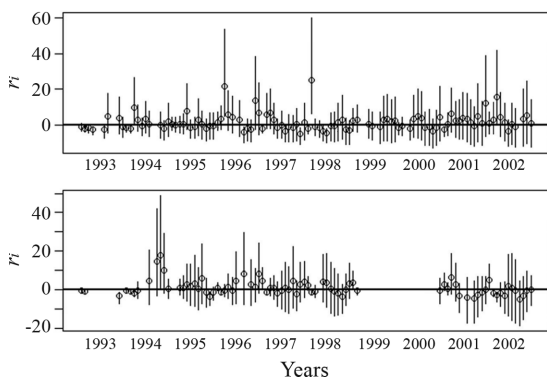


Figure 4 - Posterior standard residuals means (points) with 95% probability intervals (vertical lines) for Molhe Leste REVIS (above) and Ilha dos Lobos REVIS (below). The horizontal lines represent posterior mean equal to zero. Model 7d.

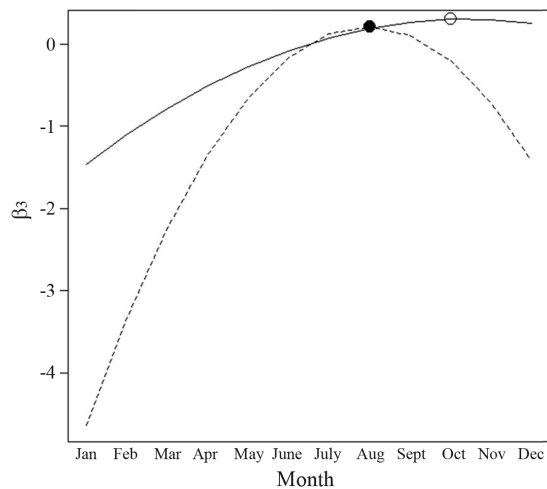


Figure 3 - Quadratic effect of covariate Month on the abundance of *Otaria flavescens* as represented by the coefficient β_3 ; continuous line represents Molhe Leste (ML) and dotted line Ilha dos Lobos (IL); maxima are represented by the open dot for ML and the solid dot for IL.

model underestimates, while intervals below the line indicate model overestimates. All remaining points are within acceptable error range. Hence, for 86.1% of the data the model provides a good description of abundance, with overestimation in 11.1% and underestimation in the remaining 2.8% of the data.

Finally, the estimated occupancy probabilities for model 7d (Figure 5) define periods of the year in which area use was certain (probability 1). But, the increase over the years, in the smallest occupancy probability is a very clear indication that these sites are becoming more intensely occupied year after year.

4. Discussion

4.1. Covariates

The success of data analysis and inference depends of the choice of a statistical model that best approximates the data structure and the underlying ecological process. The principle of parsimony states that a good model should thrive for the smallest number of parameters able to represent the relevant features of data (Burnham and Anderson, 2002). The choice among covariates to incorporate is, therefore, an important part in model selection and DIC a useful criterion to choose from competing candidates.

As with other penalised likelihood criteria (e.g., Akaike Information Criterion or Bayesian Information Criterion), we caution that DIC is not intended for identification of the "correct" model, but rather designed to merely compare a collection of alternative formulations (Carlin, 2000). In ecology there is often a great deal of ambiguity about the process involved in determining the true value of the response. If a number of possible model structures fit the observed data similarly, then there will be considerable uncertainty about which model is in fact the best one. If a plausible alternative model structure results in predictions that are different from those of the chosen best model, then there is some risk involved in bravely ignoring the alternative (Wintle et al., 2003).

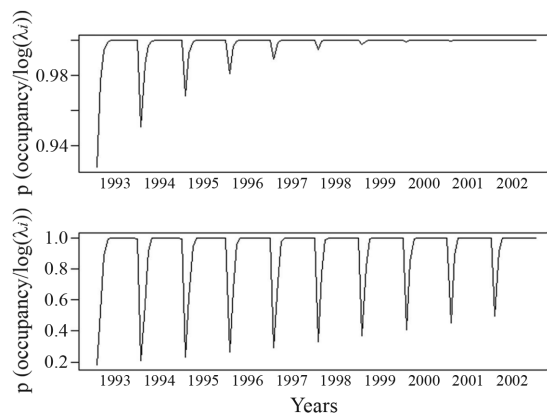


Figure 5 - Monthly occupancy probability of *Otaria flavescens* (posterior mean) estimated for model 7d at Molhe Leste REVIS (above) and Ilha dos Lobos REVIS (below).

In our analysis all selected covariates contributed to the decrease in DIC indicating that they help to explain the abundance variation. But, only the inclusion of the over-dispersion parameter was able to cause a decisive reduction in DIC. Similar preference for an over-dispersed model fit was achieved by Kinas et al. (2005) using a negative binomial distribution of the response variable in a conventional generalized linear model (GLM).

Despite the fact that DIC indicated 6d as the best fit, we opted for choosing the sub-optimal model 7d to make our inferences because this model describes important aspects of the distinctions between both sites. Although slightly over-parameterised, we believe that the ecological features that emphasise such distinction between the haul-outs is a fair price to pay.

The hypothesis formulated by Silva (2002) that sea lions, when leaving Uruguay after the reproductive season first arrive at ML, divide themselves between the two sites in a later time and return to Uruguay after a final “pit-stop” at ML, is corroborated by the distinct monthly occupancy patterns in both haul-outs.

The residual analysis was presented in the form of probability intervals. This is a unique feature of the Bayesian approach to residual analysis and a helpful tool to easily identify unacceptable discrepancies. Since in a Poisson regression variances are proportional to means, different residuals have to be analysed distinctively. This explains why some small residuals are considered unacceptable (short probability interval) in comparison to others that are much larger but associated to wide probability intervals.

4.2. Seasonality

Observed counts are useful as relative indexes of population size and indicators of abundance patterns (Márquez et al., 2006). Due to a possible availability bias, the resulting abundance estimates should be treated as “minimal average abundance”. But, provided that the number of animals absent from the haul-out at time of counting represents an approximately stable percentage of total abundance, seasonal variation and long-term trends should be analysed as if they were bias-free.

The seasonal movements observed in our data may be partially attributed to changes in the availability of food resources near the rookeries (Rosas et al., 1994; Stern, 2002), since food distribution in the ocean may be heavily influenced by physical properties like water temperature and nutrient availability (Bowen et al., 2002). The location of biologically productive sites and associated physical fronts may result in high abundance of prey that compound the diet of pinnipeds (Bowen et al., 2002).

The south Brazilian coast suffers influence of the Subtropical Convergence consisting of the confluence of tropical waters provided by the Brazil Current (BC) and sub-Antarctic waters provided by the Malvinas Current (MC). Wind-driven circulation can cause latitudinal displacement of this mixture zone (Palma et al., 2004). These waters are also influenced by the continental water discharges of the La Plata and Patos Lagoon estuaries

which support high biological productivity (Castro and Miranda, 1998). All these factors are important features that impact fish abundance on the continental shelf, mainly demersal teleosts (family *Sciaenidae*) (Haimovici et al., 1997), the preferred prey of *O. flavescens*.

Another important productivity source, ensuring high fish catches in the region, is the intrusion of nutrient through the South Atlantic Central Water along the bottom of the shelf reaching sub-superficial layers. This upwelling occurs mostly in the border region between the states Santa Catarina and Rio Grande do Sul, specifically at Torres where IL is located (Brandini, 1988).

Furthermore, stochastic events like *El Niño Southern Oscillation (ENSO)* may carry out prey to the adjacent area of the Patos Lagoon estuary (García et al., 2003), causing the South American sea lions displacement in order to feed elsewhere. Thus, strong discharges caused by the *ENSO* may change the foraging behaviour including longer trip period and distances and increase of foraging effort (Bowen et al., 2002). This hypothesis corroborates the observations of Silva (2002) who described low abundance for *O. flavescens* in 1998 at ML, a year with strong *El Niño*. However, none of the covariates related to *ENSO*, like *Southern Oscillation Index*, *Oceanic Niño Index* and *Multivariate ENSO Index* had significant effect and were therefore omitted from the text.

Further influence on the foraging of pinnipeds is water temperature, which seems to condition the arrival of sea lions at the Rio Grande do Sul shelf. According to Pinedo (1990), the displacement of *O. flavescens* is supported by the cold waters of the MC inner branch. The mean latitude at which the BC meets the MC depends on season (Reid et al., 1977). The winter average latitude is 25.8° S and has been reported to reach the minimal latitude of 25.2° S in August (Souza and Robinson, 2004). The Southern sea lions also reached the highest abundance at RS in winter and beginning of spring and specifically in August at IL which also agrees with the hypothesis put forward by Pinedo (1990).

This study allows the conclusion that abundance of *O. flavescens* on haul-outs of Rio Grande do Sul varied seasonally reaching maximum numbers in winter and early spring with peaks in August and October for Molhe Leste and Ilha dos Lobos, respectively. This cyclic pattern is combined with a long-term linear increasing trend in abundance. This trend is further supported by the long-term increase observed for minimum intra-year occupation probability by *O. flavescens*. Finally, the hierarchical Bayesian inference of an over-dispersed Poisson model was able to provide a good fit to data and to contribute to better understand the interplay of various covariates in abundance and occupancy patterns.

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References

- BOWEN, WD., BECK, CA. and AUSTIN, DA., 2009. Pinniped Ecology. In PERRIN, WF., WURSIG, B. and THEWISSEN, JGM. (Eds.). *Encyclopedia of Marine Mammals*. San Diego: Academic Press. p. 852-860.
- BRANDINI, FP., 1988. Hydrography, phytoplankton biomass and photosynthesis in shelf and oceanic waters off south-eastern Brazil during autumn (May/June, 1983). *Braz. J. Oceanogr.*, vol. 36, no. 1-2, p. 63-72.
- BUCKLAND, ST. and YORK, AE., 2009. Abundance Estimation. In PERRIN, WF., WURSIG, B. and THEWISSEN, JGM. (Eds.). *Encyclopedia of Marine Mammals*. San Diego: Academic Press. p. 1-6.
- BURNHAM, KP. and ANDERSON, DR., 2002. *Model Selection and Multimodel Inference: A Practical Information – Theoretic Approach*. New York: Springer Science+Business Media. 488 p.
- CAMPAGNA, C., 1985. The breeding cycle of the southern sea lion, *Otaria byronia*. *Mar. Mam. Sci.*, vol. 1, no. 3, p. 210-218.
- CAMPAGNA, C., WERNER, R., KARESH, W., MARÍN, MR., KOONTZ, F. and COOK, R. and KOONTZ, C., 2001. Movements and location at sea of South American sea lions (*Otaria flavescens*). *J. Zool.*, vol. 255, no. 2, p. 205-220.
- CASTELLO, JP. and PINEDO, MC., 1977. Os visitantes ocasionais do nosso litoral. *Natureza em Revista*, vol. 2, p. 40-46.
- CASTRO, BM. and MIRANDA, LB., 1998. Physical oceanography of the western Atlantic continental shelf located between 4°N and 34°S. In ROBINSON, AR. and BRINK, KH. (Eds.). *The Sea*. Oxford: John Wiley & Sons. p. 209-251.
- DANS, SL., CRESPO, EA., PEDRAZA, SN. and ALONSO, MK., 2004. Recovery of the South American sea lion (*Otaria flavescens*) population in northern Patagonia. *Can. J. Fish. Aquatic Sci.*, vol. 61, no. 9, p.1681-1690.
- DOBSON, AJ., 2002. *An Introduction to Generalized Linear Models*. Florida: Chapman and Hall; CRC. 221 p.
- FARAWAY, JJ., 2006. *Extending the Linear Model with R*. Florida: Chapman and Hall; CRC. 332 p.
- GARCIA, AM., VIEIRA, JP. and WINEMILLER, KO., 2003. Effects of 1997-1998 El Niño on the dynamics of the shallow-water fish assemblage of the Patos Lagoon Estuary (Brazil). *Est., Coast. Shelf Sci.*, vol. 57, no. 3, p. 489-500.
- GELMAN, A., CARLIN, JB., STERN, HS. and RUBIN, DB., 1995. *Bayesian Data Analysis*. New York: Chapman and Hall; CRC. 668 p.
- GIARDINO, G., MANDIOLA, MA., BASTIDA, R. and RODRÍGUEZ, D., 2009. Movimientos estivales de machos de *Otaria flavescens* entre Puerto Quequén (Argentina), Uruguay y Patagonia. In *Proceedings of the workshop Estado de situación del lobo marino común en su área de distribución*. p. 35.
- HAIMOVICI, M., CASTELLO, JP. and VOOREN, CM., 1997. Fisheries. In SEELIGER U., ODEBRECHT C. and CASTELLO JP. (Eds.). *Subtropical Convergence Environments: the Coast and Sea in the Southwestern Atlantic*. New York: Springer-Verlag. p.1-17.
- KINAS, PG., SILVA, KG., ESTIMA, SC. and MONTEIRO, DS., 2005. Generalized linear models applied to stranding data of South American sea lions (*Otaria flavescens*) and South American fur seals (*Arctocephalus australis*) in southern Brazil. *Lajam.*, vol. 4, no. 1, p. 7-14.
- LEINZ, V. and AMARAL, SE., 1966. *Geologia Geral*. São Paulo: Nacional. 512 p.
- MÁRQUEZ, J., YATACO, G., ALZA, L. and VAN OORDT, F., 2006. Changes in population size and distribution of the fur seal *Arctocephalus australis*, and sea lion *Otaria flavescens*, off Peru, from 1971-2006. In *Proceedings of the Humboldt Current System International Conference: Climate, ocean dynamics, ecosystem processes, and fisheries*. p. 193.
- MCCARTHY, MA., 2007. *Bayesian Methods for Ecology*. New York: Cambridge University Press. 296 p.
- PÁEZ, E., 1996. Simulaciones estocásticas en la población de *Otaria flavescens* en Uruguay. In *Proceedings of the VII Reunión de Trabajo de Expertos en Mamíferos Acuáticos de América del Sur*. p. 116.
- PALMA, ED., MATANO, RP., PIOLA, AR. and SITZ, LE., 2004. A comparison of the circulation patterns over the Southwestern Atlantic Shelf driven by different wind stress climatology. *Geophys. Res. Lett.*, vol. 31, no. L24303, p. 1-5.
- PINEDO, MC., 1990. Ocorrência de pinípedes na costa brasileira. *Garcia de Orta, série de Zoologia*, vol. 15, no. 2, p. 37-38.
- REID, JL., NOWLIN, WDJR. and PATZERD, WC., 1977. On the characteristics and circulation of the Southwestern Atlantic Ocean. *J. Phys. Oceanogr.*, vol. 7, no. 1, p. 62-91.
- RODRÍGUEZ, DH., 1990. *Aspectos biológicos, ecológicos e históricos de la colonia de lobos marinos de un pelo, Otaria flavescens (Shaw, 1800), el puerto Mar del Plata*. Mar del Plata: Universidad Nacional de Mar del Plata. 168 p. Tesis de Doctorado en Ciencias.
- ROSAS, FCW., PINEDO MC., MARMONTEL M. and HAIMOVICI M., 1994. Seasonal movements of South American sea lion (*Otaria flavescens*, Shaw) off the Rio Grande do Sul coast, Brazil. *Mammalia*, vol. 58, no. 1, p. 51-59.
- ROYLE, JA. and DORAZIO, RM., 2008. *Hierarchical Modeling and Inference in Ecology*. San Diego: Academic Press. 444 p.
- SANFÉLICE, D., VASQUES, VC. and CRESPO, EA., 1996. Ocupação sazonal por *Otariidae* (Mammalia-Pinnipedia) da Reserva Ecológica da Ilha dos Lobos, Município de Torres, Rio Grande do Sul, Brasil. In *Proceedings of the VII Reunión de Trabajo de Expertos en Mamíferos Acuáticos de América del Sur*. p. 46.
- SEELIGER, U., ODEBRECHT, C. and CASTELO, JP., 1997. *Subtropical Convergence Environments: The coast and the Sea in the Southwestern Atlantic*. New York: Springer-Verlag. 398 p.
- SILVA, KG., 2004. *Os Pinípedes no Brasil. Ocorrências, Estimativas Populacionais e Conservação*. Rio Grande: Universidade Federal do Rio Grande. 242 p. Tese de doutorado em Oceanografia Biológica.

- SOUZA, RB. and ROBINSON, IS., 2004. Lagrangian and Satellite Observations of the Brazilian Coastal Current. *Cont. Shelf Res.*, vol. 24, no. 2, p. 241-262.
- SPIEGELHALTER, DJ., BEST, NJ., CARLIN, BP. and VANDER LINDE, A., 2002. Bayesian measure of model complexity and fit. *Jour. Royal Stat.*, vol. 64, no. 4, p. 583-639.
- STERN, SJ., 2009. Migration and Movements Patterns. In PERRIN, WF., WURSIG, B. and THEWISSEN, JGM. (Eds.). *Encyclopedia of Marine Mammals*. San Diego: Academic Press. p. 742-748.
- STURTZ, S., LIGGES, UE. and GELMAN, A., 2005. R2WinBUGS: A Package for Running WinBUGS from R. *Journal of Statistical Software*, vol. 12, no. 3, p. 1-16.
- THOMAS, A., O'HARA, B., LIGGES, U. and STURTZ, S., 2006. Making BUGS Open. *R news*, vol. 6, no. 1, p 12-17.
- THOMPSON, D., STRANGE, I., RIDDY, M. and DUCK, CD., 2005. The size and status of the population of southern sea lions *Otaria flavescens* in the Falkland Islands. *Biol. Cons.*, vol. 121, no. 3, p. 357-367.
- VAZ-FERREIRA, R., 1981. South American sea Lion *Otaria flavescens* (Shaw). In RIDGWAY S. and HARRISON R. (Eds.). *Handbook of Marine Mammals*. New York: Academic Press. p. 39-66.
- VIEIRA, EF., 1984. *Rio Grande do Sul: geografia física e vegetação*. Porto Alegre: Sagra. 184 p.
- WESTERGAARD, G., RODRIGUEZ, D., RIVERO, L. and BASTIDA, R., 1999. Dinámica y estructura de edades del apostadero de lobos marinos de un pelo (*Otaria flavescens*) de Puerto Quequén, Argentina. In *Proceedings of the VIII Congreso Latinoamericano de Ciencias del Mar*. p. 265-266.
- WINTLE, BA, MCCARTHY, MA., VOLINSKI, CT. and KAVANAGH, RP., 2003. The use of Bayesian Model Averaged to Better Represent Uncertainty in Ecological Models. *Cons. Biol.*, vol. 17, no. 6, p. 1579-1590.