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# Invasive axis deer and wild boar in a protected area in Argentina, controlled hunting, and Taylor's law

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## Abstract

*Context.* Spatial and temporal variability in population density tends to increase with an increasing mean density, as widely documented by Taylor's law (TL) of fluctuation scaling. A management program based on local hunters has been used to control invasive wild boar and axis deer in a protected area of north-eastern Argentina since 2006.

*Aim.* We determine the effects of species (boar or deer), hunting shift (diurnal, overnight), human disturbance (by comparing one section open for public use, one not) and time scale (one-versus three-month periods) on the values of the parameters of TL, and consider both its spatial and temporal forms.

*Methods.* Park management collected data on the hunting efforts and harvest of 6104 hunting parties shooting from elevated blinds from 2006 to 2015. The log-transformed sample means and variances of four indices of relative abundance were computed for each period and blind, and analysed through least-squars linear regression and ANCOVA.

*Key results.* Axis deer satisfied the spatial TL by all four indices, but wild boar had a significantly non-linear relationship for crude catch per unit effort (CP–UE) only. In the spatial TL, the slope *b* did not deviate significantly from 1 when using crude or standardised catch per hunting-party session or standardised CPUE, but *b* was substantially >1 for crude CPUE in both species (range, 1.307–1.434). Human disturbance, hunting shift, and time scale did not significantly modify the parameters of the spatial TL, except in two cases. All metrics at identified blinds over consecutive trimesters confirmed the temporal TL. Wild boar crude catch was 43% greater in the restricted zone of greater conservation value, whereas axis deer catch was 60% greater in the public-use zone.

*Conclusions.* With rare exceptions, TL describes well the mean and variance of four metrics of abundance of wild boar and axis deer under sustained hunting pressure. This paper may be the first to demonstrate the connection of TL with any aspect of vertebrate pest control.

*Implications.* TL identifies key zones with a high mean and high variance of ungulate density for targeted control, and can be used to attain fixed-precision estimates of abundance through sequential sampling.

Keywords: invasive exotic species, abundance, wildlife management, population dynamics, ungulates, Taylor's law.

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# Introduction

Invasive alien species play a major role in the current biodiversity crisis (Spear and Chown 2009). Various ungulates feature high in the list of exotic invasive species, including the wild boar (*Sus scrofa*) and several deer species (Lowe *et al.* 2004). Wild boar and chital or Indian spotted deer (*Axis axis*) have been introduced to many parts of the world for venison, sport hunting and other uses (Dolman and Wäber 2008). They have expanded their range and became overabundant because of the convergence of multiple

processes, including lack or loss of top predators, reforestation, intensification of crop production, supplementary feeding, inadequate harvest policies, and declining hunter numbers (Gamelon *et al.* 2012; Massei *et al.* 2015; Vetter *et al.* 2015; Carpio *et al.* 2021). Wild boar populations are increasing in the northern hemisphere (Snow *et al.* 2017) and the southern cone of South America (Pedrosa *et al.* 2015).

Wild boar and axis deer damage plant community structure and dynamics, compete with native grazers and livestock,

transmit several zoonotic pathogens, and cause traffic accidents (Choquenot et al. 1996; Hone 2002; Campbell and Long 2009; Barrios-García and Ballari 2012; Hess et al. 2015). Wild boar prey on native fauna and raid crops. Axis deer impacts have caused concern in Hawaii, California and New Zealand (Gogan et al. 2001; Nugent and Choquenot 2004; Hess et al. 2015). Because of the benefits and losses to different sections of society, the goals of ungulate management programs range from eradication or control to sustainable use (Gogan et al. 2001; Hone 2002; Nugent and Choquenot 2004; Hess et al. 2015). Programs to control wild boar and deer have applied shooting with firearms from the ground, vehicles, or air; hunting with trained dogs; poisoned bait; walk-in baited traps and snares (boar); and fencing (deer; Gogan et al. 2001; Nugent and Choquenot 2004; West et al. 2009; Campbell and Long 2009; Massei et al. 2015; Davis et al. 2016).

Both invasive species occur through much of Argentina and in many protected areas (Chébez and Rodríguez 2014). Wild boar strongly reduced the recruitment of the yatay palm tree (Butia vatav) in El Palmar National Park (hereafter EPNP or the park, located in north-eastern Argentina), which was created to preserve one of the few remaining high-density natural stands of this palm tree. Non-systematic control efforts initiated in 1983 and recommenced in 1995-1996 to cull exotic ungulates proved insufficient and unsustainable. The park then implemented a new management program based on local sport hunters who hunted with dogs and performed controlled shooting from elevated blinds in a closely supervised fashion from 2006 on (Gürtler et al. 2017). These efforts strongly reduced the relative abundance of wild boar within 2 years and stabilised it at levels associated with minimal soil damage (Gürtler et al. 2017). Meanwhile, axis deer numbers continued to increase steadily despite increasing harvesting (Gürtler et al. 2018). The reasons for the strong inverse correlation between boar and deer numbers remain unclear.

Spatial and temporal variability in the population abundance or density of living organisms is a central issue in ecology. One of the most widely documented patterns is that variability in population density tends to increase with an increasing mean density. Taylor's law (TL) relates the sample variance v to the sample mean m of population density or abundance through a linear relationship on log-log coordinates,  $\log_{10} v \approx$  $a + b \times \log_{10} m$  (Taylor 1961), or equivalently, the power law  $v \approx 10^{a} m^{b}$ . In purely random distributions of individuals (Poisson-distributed data), the variance equals the mean (so b = 1 and a = 0). Depending on the mechanism that generates variability in population density, slope values b significantly greater than 1 may indicate aggregation of organisms or heterogeneity within or among samples. If population density follows a gamma distribution with a constant shape parameter and changing scale parameter (including an exponential distribution as a special case), then b = 2. If population density follows a lognormal distribution, then, depending on the relation between the parameters of the log-normal distribution, TL may hold exactly with b = 2 or approximately with b = 4, b = 2 + 2/3, or other values (Tippett and Cohen 2016, their supplementary information). If population density follows a Wald or inverse Gaussian distribution, then, depending on the relation between the parameters, TL may hold exactly with b = 3.

Taylor's law can be generated by many different models. including the Cohen-Lewontin stochastic population growth model, random walks and other processes (e.g. Keeling and Grenfell 1999; Eisler et al. 2008; Cohen 2013; Cohen et al. 2013b; Cohen and Xu 2015). Empirical examples of TL come from various fields and hundreds of organisms (Eisler et al. 2008), ranging from measles (Keeling and Grenfell 1999), parasite aggregations within individual hosts (Morand and Krasnov 2008) and in space (Cohen et al. 2017a), human population densities (Cohen et al. 2013a), forest trees (Cohen et al. 2016), and crop yields (Döring et al. 2015), to the insect vectors of human disease (Cohen et al. 2017b). Recently, Cohen et al. (2016, p. 1342) identified a knowledge gap, as follows: 'How Taylor's power law, and more generally the relationship of variance to mean of population density, varies with the scale of observation, time, disturbance, and other biotic and abiotic environmental characteristics has received insufficient attention'.

Application of TL to fish and wildlife management and population dynamics has been sparse. It was mainly restricted to sample-size determination of elephants and elephant seals (Barnes *et al.* 1997; Walsh *et al.* 2001; McMahon *et al.* 2005); fisheries fixed-precision sampling (Xu *et al.* 2017, 2019), spatial aggregation of kangaroos (Pople *et al.* 2007), hunting of red grouse (Cattadori *et al.* 2003; Willebrand *et al.* 2011) and willow ptarmigan (Kvasnes *et al.* 2015). The only application for wild boar we know involved the frequency of fresh dung pellets per 100 plots, which showed aggregated temporal and spatial patterns with b = 1.7–2.1 (Hone 2012, pp. 31–32).

Taylor's law has a temporal and a spatial form, as well as other forms. In a temporal TL, *n* populations labelled i = 1, ..., n, are followed over time, and the sample mean population size  $m_i$  over time and the sample variance of population size  $v_i$  over time for each population *i* are both computed, then analysed through a least-square linear regression of  $\log_{10} v_i$  as a function of  $\log_{10} m_i$ , with one (temporal mean, temporal variance) pair for each population *i*. In our study, each population will be a fixed hunting blind in which hunters perform controlled shooting with rifles.

In a spatial TL, different populations of a species occurring in specific sites are grouped into distinct categories such as habitats, labelled *h*, or in distinct time periods, labelled *t*, which we pursue here. The mean  $m_h$  and the variance  $v_h$  of population sizes over all sites of habitat *h* or of time *t* are calculated, and  $\log_{10} v_h$  is regressed on  $\log_{10} m_h$ , with one (spatial mean, spatial variance) pair for each habitat *h* or for each time *t*. Here, each category will be a different trimester *t*, in which hunters perform controlled shooting with rifles from elevated blinds. We also considered 1-month periods instead of trimesters, to see whether the scale of observations modified the parameters of TL.

We show here how well TL describes the temporal and spatial mean and variance of various metrics of the relative abundance of wild boar and axis deer under sustained hunting pressure in a protected area of north-eastern Argentina over a decade. In each hunting session, each hunting party's effort and harvest by controlled shooting from an elevated blind with a fixed position were rigorously documented. We determine the effects of ungulate species, hunting shift (diurnal vs overnight), human disturbance (by comparing two park sections differing in whether they were open for public use or not, the latter being of greater conservation value), and time scale of observations

YPI	No. of h	unting	Session du	ration (h)	Hunting	effort (party-h)	Ax	is deer	Wi	ld boar	Undefi	ned species	Total
	Sessions	Parties	Mean	s.d.	Crude	Standardised	Killed	Wounded	Killed	Wounded	Killed	Wounded	
0	50	374	11.7	3.0	4368	1753	110	3	292	7	0	1	413
1	39	575	7.4	3.0	4252	2683	141	3	218	16	1	0	379
2	34	556	5.1	1.6	2820	2281	120	9	108	4	2	7	250
3	30	565	5.0	1.6	2827	2346	161	14	104	5	6	0	290
4	28	569	4.8	1.5	2716	2357	175	30	118	14	0	2	339
5	28	610	4.9	1.1	3003	2685	213	16	185	10	11	4	439
6	15	411	4.9	2.0	2002	1754	180	23	150	12	0	4	369
7	25	712	5.6	2.1	3996	3526	330	17	174	12	0	3	536
8	29	871	5.7	1.5	4974	3755	442	17	168	7	0	7	641
9	26	861	5.5	1.2	4749	3813	516	33	195	14	0	9	767
Total	304	6104	5.8	2.5	35 707	26953	2388	165	1712	101	20	37	4423

 Table 1.
 Annual crude hunting effort and catch of wild boar and axis deer killed or wounded by controlled shooting with rifles from elevated blinds according to year post-intervention (YPI) at El Palmar National Park, 2006–2015

(1-month vs 3-month periods) on the values of the parameters of TL, and consider both the spatial and temporal forms of TL.

We suggest some practical consequences and potential uses of TL in wildlife management to plan more efficient control measures. The present paper may be the first to demonstrate the connection of TL with any aspect of vertebrate pest control, and in particular with two major invasive alien species under sustained management over a decade.

## Materials and methods

#### Study area

El Palmar National Park (31°55'S, 58°16'W), in Entre Ríos, north-eastern Argentina, covers ~8500 ha of savannas, grasslands, scrublands and gallery forests, including high-density stands of the palm Butia yatay, limited by the Uruguay River in the east and a fast highway (Route 14) in the west (Gürtler et al. 2017). The park is mainly surrounded by dense eucalyptus and pine plantations, sometimes combined with cattle for firebreak maintenance, some cattle ranching (on both park borders by the Uruguay River), and a small patch for crops (on permanent rotation including maize, soybean, sorghum and wheat, adjacent to the southern border). Native scrubland occupies a very small fraction of lands adjacent to the park. Intensified agriculture is restricted to the upper watershed of Arroyo Palmar, some 30 km to the west of the park. A permanent water course (Arroyo del Palmar) divides the park into two zones; another three minor water courses cross the park from west to east. The northern zone is open for public use. The southern zone is a restricted area closed to tourists and considered of greater conservation value. Both zones are covered by palm trees, and are poached for deer, boar and capybaras. The increasing expansion of several exotic trees and shrubs, closely linked to the suppression of spontaneous fires and exclusion of cattle in 1970, led to the current lignification of both zones (Ruiz Selmo et al. 2007; Batista et al. 2018). Prescribed fires conducted irregularly since 2002 have not been able to reduce the coverage of woody plants and restore the savanna landscape.

The mean annual temperature over 2006–2015 was 19°C, and the annual mean rainfall was 1389 mm at the closest weather station in Concordia. Long droughts ( $\geq$ 3 months) occurred from the 2005 spring–summer to the 2006 winter; over the same period in 2008–2009; from summer 2011 to winter 2012; over the 2013 fall–winter, and from summer 2014 to fall 2015.

# Field data

The data come from the ongoing exotic-mammal management program at EPNP initiated in 2006. The program recruited local sport and subsistence hunters to cull wild boar and exotic deer under a regulated framework. The goals were to reduce the negative impacts of wild boar (on ground rooting and the recruitment of yatay palm trees) and to reduce the abundance of axis deer and blackbucks. The details of the program and a map of the study area have been described elsewhere (Gürtler *et al.* 2017).

Wild boar and axis deer were culled by controlled shooting, hunting with dogs, and spotlight hunting with spotlights and firearms from pick-up trucks. Hunting with trained dogs targeted wild boar only (although the axis deer incidentally found were dispatched) mainly over 2006–2007, and was accomplished by hunters using several packs of mixed-breed dogs (Table 1 in Gürtler *et al.* 2018). Spotlight hunting (which had a marginal role over 2006–2015) and dog-hunting are excluded from further consideration here.

Controlled shooting using rifles with designated calibres was conducted from elevated blinds (watchtowers) distributed rather uniformly across the park (see fig. 2b in Gürtler et al. 2017). The number of active blinds (i.e. blinds that returned at least one hunting form over any year, regardless of whether they fired any shot) increased from 29 to 40 between year 0 and 3, then decreased to 36 by year 6, and increased to 46 in year 9; 47 identified blinds ever existed over the decade. Each blind had 5-20 shooting lanes in which the vegetation had been cleared. Each lane was from 200 m to 300 m long. Most blinds remained at a fixed position over the decade; a small fraction of them was relocated in the vicinities of the former position for improved accessibility, safety and hunting success. Typically, each hunting party had one or two authorised shooters, and one person in charge of illuminating the target and assisting in retrieving the quarry. However, during some overnight sessions conducted in the initial years and rarely thereafter, two self-identified groups of hunters occupied the same blind over a given session, although not at the same time; for simplicity, we considered only the first of them. Overall, this occurred in less than 1% of all

active hunting parties. Hunters and individual blinds were not in strict correspondence at all times: some hunters used the same blind for all or most of their affiliation to the program; some changed blinds on a few occasions after an extended trial period; some shifted between two blinds over several years, and some drop-outs were replaced by newcomers as time went by. Hunters were required to bait the surroundings of their assigned blind with corn or ground pet food and blocks of salt once or twice a week before hunting sessions; in practice, the type, amount and frequency of baiting varied widely.

Hunting sessions were generally conducted between 1600 hours and 1800 hours and 1000 hours and 1200 hours (diurnal or short sessions) every 1 or 2 weeks over 10–11 months a year, but the separation between successive sessions varied widely across the decade. Overnight (or long) sessions frequently started in the evening (between 1600 hours and 1900 hours up to April 2013, and from 2200 hours on since September 2014) and extended up to 0600-0800 hours. All sessions conducted between January 2006 and end of March 2007 were overnight, and thereafter occurred 0-2 times a year. The mean duration of hunting sessions decreased from 11.7 h in 2006 (overnight) to  $\sim$ 5 h (diurnal) from 2008 onward (Table 1), depending on park decisions, weather conditions, and the choice of individual hunting parties. No catch quotas or limits to the number of shots allowed during a hunting session were set. Each hunter was allowed to take back home up to one annual trophy to minimise selective hunting.

Every hunting session had a park ranger who supervised the activities *in situ* or at the central operating post (since 2008) through a VHF radio, with which he or she communicated with all hunting parties and other assisting park staff to announce the start and termination of a hunting session and grant permission for ground movement. Hunters communicated to the supervisor any relevant event, reported every shot, and requested permission to descend to finish off or retrieve the quarry or search for wounded specimens within 200 m from the hunting blind. The radio operator recorded in a dedicated logbook the timing and number of shots that each self-identified hunting party made, whether they killed a deer or boar, and any other relevant events (e.g. poaching).

The culled specimens were brought to the central operating post and measured. For each hunting party, park staff filled in a numbered form regardless of whether any wild boar or deer was dispatched, and the reporting hunter signed it. Each form included date, hunters' names, hunting method, initiation and termination times of the session, firearms and calibres, catch time, whether any boar or deer was wounded and escaped, each specimen's morphometric measurements, and other details.

## Definitions and metrics of relative abundance

For present purposes, we adopt the following definitions: a 'hunting party' represents a group of hunters hunting together (using the same blind) on a particular occasion (session), regardless of whether any exotic ungulate was dispatched or any shot was fired. For the few cases in which two parties successively occupied the same blind on a particular session (which occurred in less than 1% of all hunting parties), for simplicity, we only considered the first of them. A 'hunting session' is the activity of a hunting party on a given day over one continuous

time interval. If the same hunting party hunts on the same session running overnight, we count this as one hunting session. Active hunting parties are those who participated in a hunting session, regardless of whether they were effective in culling or not and of how long they hunted.

Catch ( $C_j$ , or harvest) is the number of specimens culled by a hunting party (*j*) during a fraction or the whole of a hunting session using a given hunting method.  $C_j$  includes a few culled specimens in a poor body condition which were discarded and not measured. For each ungulate species separately (and for both pooled) culled by a defined method, aggregate catch ( $C_t$ ) is the sum, over all hunting parties *j*, of the catch of hunting party *j* at time *t* (i.e. session, trimester or month, and year). Wounded specimens that escaped (as reported by hunters) were considered separately as putative crippling losses; these data were widely variable over time and thus were considered less reliable. Therefore, we alternatively excluded them or pooled them with killed boar, deer and undefined ungulate species (i.e. prey).

For a given hunting party, session and method, hunting effort  $(E_j)$  is the sum of hunting-party hours regardless of whether the party caught any boar or deer, and regardless of whether there was one shooter or there were two shooters on a hunting blind in a given session.  $E_j$  was computed as the effective duration of individual sessions ( $\pi_{j,t}$ , the difference between termination and initiation times of each hunting party at time *t*). Standardised hunting effort is constrained to hunting activity executed over 17–23 h (i.e. the most typical time band and duration across the decade) to allow for hypothetically greater ungulate activity over this period and the much longer duration of sessions during 2006 and 2007 (see above). Aggregate hunting effort ( $E_t$ ) was the total number of party hours invested by all hunting parties in a given hunting session or time period *t* by using a defined method.

The CPUE index (*C*/*E*, catch per unit effort) is linearly related to density *D* (abundance *N* relative to area *A*, *N*/*A*) through the catchability coefficient (*q*) under the assumptions of equal *D* across the area, units of effort (here hunting parties) acting independently, low catch probabilities, and constant *q* and *A* (Clark 1985; Hilborn and Walters 1992; Skalski *et al.* 2005). Explicitly, C/E = qD.

We used four partially related metrics to represent the relative abundance of wild boar and axis deer, and of both species combined (including killed and wounded specimens of any type, i.e. prey, as if hunters were not able to distinguish between exotic ungulate species) over a given time period (Table 1). The underlying rationale was to allow for the limitations of crude indices as surrogates of population size, and because to the best of our knowledge, whether crude or standardised measures of CPUE are better metrics for wildlife has not been addressed before. For crude catch per hunting-party session (C, metric 1), the time unit is the entire hunting session regardless of its duration. Standardised catch per hunting party (metric 2) is the sum of specimens culled over 17-23 h by each hunting party, regardless of the total duration of each session. Crude CPUE (metric 3) is C/E (crude catch per crude effort). Standardised CPUE (metric 4) restricted both the catch and hunting effort to a fixed time window (17-23 h), because hunting sessions differed in total duration and not all hunting parties spent the same amount of time on a given session. For

each metric, we calculated the mean and variance across active hunting parties for each trimester or month (spatial TL) or for each hunting blind (temporal TL) across the decade.

#### Datasets

The management program kept hunting records and compiled a digital database for interim assessment and decision-making. Our current analysis is based on a recent complete review of the digital database against the original hunting forms (in paper format), with the goals of verifying data, filling in missing information, and using revised criteria (e.g. alternative exclusion of wounded specimens). These revisions led to slight changes in catch and hunting effort numbers relative to the data used in previous analyses (Gürtler et al. 2017, 2018). The revised database for controlled shooting includes 6104 records, 1712 wild boar and 2388 deer specimens culled with exact date and time of session, hunting method and hunting party form. Two relevant variables had missing data, namely, hunting blind identity (28 of 6104 records, 0.5%), and time of catch (668 of 4100 guarry, 16.3%). Specimens that were reportedly wounded and escaped were considered separately from killed specimens. For 158 specimens with complete morphometric data (excluding tail length and weight) but lacking species identification, we assigned them to species on the basis of a discriminant analysis of log<sub>10</sub>-transformed measurements as described in Gürtler et al. (2017). Here, we used the available hunting sessions' logbooks to trace back the reported species identity of specimens with missing identification in the hunting forms; the concordance between morphometric assignments and logbook records among 55 specimens with data was 100%. Data for the 10-year period (Table S1) and the code used for analysis (Appendix S1) are included as supplementary files.

#### Fitting and testing Taylor's law

We fitted TL and tested its adequacy as a description of the data in three steps, following the detailed account in Cohen et al. (2017b). First, for each ungulate species separately and for both combined, we computed the sample mean  $(m_h)$  and the sample variance  $(v_h)$  of the catch and CPUE achieved by each active hunting party at a given blind and session over each trimester. For example, for each trimester, we pooled the numbers of boar killed by a hunting party in a hunting blind at a given hunting session (variable X, either catch or CPUE), then computed the mean and variance of X across active hunting parties in this trimester (regardless of the number of hunting sessions or active parties during the 3-month period), then log<sub>10</sub>-transformed both the mean and the variance of X, and performed an ordinary leastsquares linear regression of  $\log_{10} v_h$  on  $\log_{10} m_h$ , t = 1, ..., 38, across all trimesters with data. We repeated these procedures for each ungulate species culled (taken separately and combined), hunting shift (diurnal versus overnight), and human disturbance (public vs restricted zones) using Stata 15.1 (Stata Corp 2018). We also included a least-squares linear regression for the combined data for ungulate species, including the number of killed or wounded specimens regardless of whether they had been identified to species level (i.e. prey). Analyses always complied with the requirements that the mean abundance was greater than 0, that at least 15 observations (here hunting parties for each

3-month or 1-month period) were used to calculate each mean and variance, and that the linear regression (here, for a given species, zone, and hunting shift) should include at least five paired data of  $v_h$  and  $m_h$  (Taylor *et al.* 1988, p. 721).

For this and all subsequent statistical tests of a null hypothesis, the critical *P*-value was set at 0.01. We did not attempt to correct the critical value of these tests for simultaneous statistical inference (Miller 1966) because of the complex dependence of the data used in these tests. For example, the widely used Bonferroni correction assumes that all tests are statistically independent, which is not true here because the four different metrics of abundance are based on overlapping data. Hence, we rely on a combination of statistical hypothesis testing and visual judgments.

Second, we tested for curvature in the relation of  $\log_{10} v_h$  to  $\log_{10} m_h$ , by fitting a quadratic regression  $\log_{10} v_h = a + b \times$  $\log_{10} m_h + c \times (\log_{10} m_h)^2$  by least-squares (Taylor *et al.* 1978, p. 388, their eqn 14). We refer to this quadratic generalisation of TL as QTL. If the confidence interval of the coefficient c did not include 0, the data rejected TL. We examined the residuals of the linear regression models for heteroskedasticity, normality, skewness and kurtosis by using the commands swilk, estat hottest and estat imtest using Stata 15.1. The results of these tests were not always consistent, even when these tests were supposed to evaluate the same characteristic of the residuals. In such cases, we examined the data graphically and used our judgment to decide on the overall pattern of the data, rather than relying on contradictory evaluations from different tests of statistical significance. Each residual of the TL linear regression measured the stability of population abundance following Döring et al. (2015).

Third, when the analyses in steps 1 and 2 did not reject TL, we used analysis of covariance (ANCOVA, implemented in the anova command of Stata 15.1) to test for differences in the slope and intercept of the species-specific TLs fitted to different subsets of the data (i.e. ungulate species, zones, and hunting shifts). For example, one ANCOVA treated 'species' as a categorical variable and asked whether 'species' or the interaction term 'species  $\times \log$  sample mean' significantly influenced log sample variance. If the interaction term influenced log sample variance, then 'species' affected the slope. If both 'species' and the interaction term influenced log sample variance, then both the intercept and the slope of TL depended on the ungulate species. We used Welch's t-test for two quantities with unequal variances to compare slope estimates under two conditions. We present back-transformed means and 95% confidence intervals of indices of ungulate abundance for convenience of interpretation.

Finally, we asked whether the choice of the time scale over which relative abundances were computed would affect the parameters of Taylor's law. Therefore, we repeated steps 1-3 described above for 1-month periods over the 10-year study.

#### Results

# Spatial Taylor's law Wild boar

The spatial TL described controlled shooting of wild boar reasonably well (Fig. 1, Table 2). TL, which corresponds to the



**Fig. 1.** The spatial TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the relative abundance of *wild boar*, under conditions of controlled shooting from a fixed hunting blind. Each point represents the (*a*) spatial mean and spatial variance of crude catch per hunting party-session, (*b*) standardised catch per hunting party-session, (*c*) crude catch per unit effort, and (*d*) standardised catch per unit effort across El Palmar National Park at one trimester over 2006–2015. The solid straight lines are fitted by least-squares regression to the data from each trimester separately. The dashed belt above and below each solid regression line represents a 95% confidence interval for individual residuals. Numbers next to the circles represent trimesters over the 10-year period.

null hypothesis that c = 0 in the QTL, was rejected (P < 0.01) only for crude CPUE. The low-variance outlier in trimester 2 and the high-variance outlier in trimester 22 (Fig. 1*c*) probably contributed to the rejection of TL for crude CPUE. We discuss outliers below. For the remaining three of the four metrics, the slope coefficients *b* of TL were insignificantly different from 1 (Table 2, Fig. 1*a*, *b*, *d*). However, the intercept *a* of TL was significantly positive for standardised catch and significantly negative for standardised CPUE, rejecting the Poisson model of random catches in these cases.

The extreme data point labelled 2 (corresponding to the second trimester in 2006) in three of the metrics differed from other trimesters in having few hunting parties on each session, long overnight sessions, and mostly comprising blinds located in the restricted zone, and hunting procedures being in the adjustment stage (Fig. 1a-c). Trimester 1 suffered the same problems. Trimester 22 had fewer hunting sessions than did preceding and subsequent trimesters, mostly because of management-related issues. When we re-ran the analyses in Table 2 after deleting trimesters 2 and 22, of the four metrics for each species, none rejected TL in favour of QTL.

The Shapiro–Wilk test showed significant deviations from normality for crude CPUE only in both TL and QTL, but the other tests showed significant deviations of residuals from normality, homoskedasticity and normal kurtosis for crude catch and crude CPUE.

# Axis deer

The data for axis deer were consistent with (i.e. failed to reject) the spatial TL for the four metrics, with no significant curvature. The log mean crude catch was highly significantly correlated with the log variance of crude catch across all active hunting blinds over both zones of the park (adj.  $R^2 = 0.779$ ); the slope coefficient did not differ significantly from 1 (b = 1.175 point estimate  $\pm 0.103$  standard error) but the intercept differed highly significantly from 0 ( $a = 0.250 \pm 0.051$ ; Fig. 2a, Table 2). The first and second trimesters appeared repeatedly as outliers (Fig. 2a, b, d), along with trimesters 7 and 17 sporadically. The slope coefficient b was not significantly different from 1 for standardised catch ( $1.000 \pm 0.076$ , adj.  $R^2 = 0.749$ ) and standardised CPUE ( $b = 1.307 \pm 0.120$ , adj.  $R^2 = 0.760$ ; Fig. 2b-d, Table 2).

Table 2. Lines	ar regression esti	imates (by ord	inary least-squ	ares) of the para	neters of th	e spatia	I Taylor	's law (J	(L) $\log_{10} \nu$ =	= a + b	< log <sub>10</sub> n	i, wher	m = san	nple mean	v = sam	ple varia	nce of
various metrics	of the relative ab	undance of wil	d boar, axis dee	er and both specie	s combined (	includi	ng killed	and wou	unded speci	mens, or	excludi	no mon	inded prey	) compute	ed for 3-m	onth peri	odsat
				H	l Palmar N	ational	Park, 20	06-2015									
Minimum log me	ean, minimum ove	er all hunting bli	inds with at leas	t one prey of the lo	g10 of the san	nple mea	un m <sub>t</sub> cato	ch or CPI	JE in time t.	Maximu	m log m	ean, ma	iximum ov	er all hunti	ing blinds	with at lea	astone
prey of the $\log_{10}$	of the sample mea	an $m_t$ catch or C	PUE. Range log	g mean, maximum	log mean mir	nus mini	mum log	g mean, to	o indicate th	e range o	f the abs	cissa in	the scatter	rplot of TL	. Intercept	_a, least-	square
estimate of the ir.	f h Curvature c 1	g-log form of T	aylor's law (TL) imate of the cur	$\log_{10} v = a + b \times v_{\text{struct}}$	log <sub>10</sub> m. s.e 4ratic Tavlot	a, stand	ard error	of the est	timate of a. S	Slope_b, ftheeti	least-squ	lare est $\sim_{adi} R^{i}$	imate of th	e slope b of R <sup>2</sup> (adinste	f TL. s.e] ed for the i	b, standar	d error f fitted
narameters) d f	,	learnes of freed	alound (d f.) aduale	number of observ	ations minus	the num	har of fi		matars Tas		to onlow	the tec	, uujuuuu t of the nul	ll hundthae	is that the	intercent	0-0
Test_c_0, P-vali	ue of the test of th	regrees of freed	sis that the coef	ficient c of the que	adratic term	equals 0	. A P-va	lue of <	0.01 rejects	the adec $\frac{D}{M}$	luacy of	TL bed	ause the r	elationship	of log <sub>10</sub>	v to log <sub>10</sub>	m = 0.
Heteroskedastici hypothesis	ity, P-value of the sfor heteroskedasi	test of the null h ticity, skewness	ypothesis that these of the	the residuals from the Catch, catch per hi	le linear regr unting sessio	ession al ession al n; Stanc	re homos e homos l., standa	kedastic, rdised; C	that is, all h PUE, catch	ave the s ave the s	ame vari effort; H	ance. Ir let., he	uformation teroskedas	matrix, <i>P</i> -ticity; IM,	value of th informati	le test of t on matrix	he null
Species	Metric	Min log mean	Max log mean	Range log mean	Intercept_a	s.ea	Slope_b	s.eb	Curvature_c	s.ec	Adj. $R^2$	d.f.	rest_a_0	Test_c_0	Swilk	Het.	IM
Wild boar	1. Crude catch	-0.930	0.000	0.930	0.073	0.044	0.949	0.072	-0.675	0.259	0.823	36	0.105	0.013	0.309	0.013	0.002
	2. Stand. catch	-1.572	-0.121	1.451	0.170	0.041	1.096	0.052	-0.243	0.104	0.922	36	< 0.001	0.025	0.107	0.033	0.026
	3. Crude CPUE	-1.658	-0.740	0.918	0.002	0.279	1.434	0.215	0	I	0.540	36	0.993	I	<0.001 <	<0.001	0.006
	3. Crude CPUE	-1.658	-0.740	0.918	3.306	1.008	7.064	1.676	2.335	0.691	0.643	35	0.002	< 0.001	0.004	0.074	0.204
	4. Stand. CPUE	-1.857	-0.857	1.000	-0.429	0.129	1.050	0.094	-0.050	0.302	0.771	36	0.002	0.025	0.165	0.039	0.041
Axis deer	1. Crude catch	-0.848	-0.057	0.791	0.250	0.051	1.175	0.103	0.664	0.450	0.779	36	< 0.001	0.149	0.004	0.683	0.073
	2. Stand. catch	-1.155	-0.196	0.958	0.171	0.053	1.000	0.076	-0.099	0.305	0.749	36	< 0.001	0.756	< 0.001	0.921	0.137
	3. Crude CPUE	-1.914	-0.885	1.030	-0.133	0.146	1.307	0.120	-0.654	0.355	0.760	36	0.367	0.074	0.034	0.595	0.318
	4. Stand. CPUE	-1.851	-0.895	0.957	-0.352	0.138	1.068	0.106	-0.346	0.426	0.731	36	0.015	0.422	< 0.001	0.199	0.004
Both spp., including	1. Crude catch	-0.388	0.248	0.637	0.129	0.019	1.188	0.094	1.213	0.453	0.810	36	< 0.001	0.011	0.289	0.012 <	<0.001
wounded																	
	2. Crude CPUE	-1.129	-0.504	0.626	0.048	0.204	1.669	0.231	2.545	1.175	0.580	36	0.814	0.037	0.017	0.341 <	<0.001
Both spp.,	1. Crude CPUE	-1.149	-0.527	0.622	0.068	0.219	1.655	0.239	2.767	1.200	0.559	36	0.759	0.027	0.078	0.182 <	<0.001
excluding wounded																	

Invasive axis deer, wild boar and Taylor's law



**Fig. 2.** The spatial TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the relative abundance of *axis deer*, under conditions of controlled shooting from a fixed hunting blind. Each point represents the (*a*) spatial mean and spatial variance of crude catch per hunting party-session, (*b*) standardised catch per hunting party-session, (*c*) crude catch per unit effort and (*d*) standardised catch per unit effort across El Palmar National Park at one trimester over 2006–2015. Key as in Fig. 1.

Intercepts were significantly different from 0 for crude and standardised catch only. Residuals were not normally distributed in most cases, according to the Shapiro–Wilk test, and showed significant deviations from homoskedasticity and normal kurtosis only for standardised CPUE.

ANCOVA showed no significant effects of species on coefficients *a* and *b* of TL for crude catch, standardised catch and standardised CPUE, suggesting that if species affected the slope at all, it did so weakly for deer crude catch only. We did not compare crude CPUE between species by ANCOVA because QTL described crude CPUE better for boar, and TL described CPUE better for deer.

# Pooled catch

For both species combined, the crude catch of prey (including or excluding killed or wounded boar and deer) and crude CPUE were adequately described by TL. However, the quadratic coefficient c of QTL was positive but not significantly positive, suggesting a convex relation between log variance and log mean when both ungulate species were combined.

## Human disturbance

When we separated the data from the sites in the public-use zone from those from the restricted zone, TL was not rejected for any metric or either species (Table 3). Hence, it was valid to compare these zones using ANCOVA. ANCOVA yielded no significant differences in the parameters of the spatial TL between the publicuse zone and the restricted zone for any of the four metrics for wild boar and axis deer (Table 3, Fig. 3), indicating no significant effects of anthropic disturbances and other undefined landscape variation (i.e. forest structure and adjacent crops) on TL parameters. The adj.  $R^2$  was greater than 0.75 in 12 of 16 cases (16 cases = 4 metrics × 2 species × 2 zones; Table 3).

For wild boar, the slopes of TL for crude catch per huntingparty session varied little, from 0.972 to 1.016 in the restricted and public-use zones respectively (Table 3). Very slight differences between zones were recorded in the slopes for standardised catch (1.112 and 1.101), standardised CPUE (1.065 and 1.198), and crude CPUE (1.343 and 1.380). Intercepts were significantly different from 0 for standardised catch and standardised CPUE in the public-use zone only.

For axis deer, the slopes of TL for the public-use zone had a wider range (1.072–1.260) than did those for the restricted zone (1.202–1.341), but for each metric the difference between zones was slight except for standardised CPUE (b = 1.072 and 1.283; Table 3). When intercept estimates differed strongly from 0, they tended to do it in both zones. Residuals for deer metrics frequently departed from normality and homoscedasticity.

					. 201	06–2015, ac	cording	to hunti	ing zone									
	T	Key as m 1	able 2. Catch,	catch per huntin	g-session; Stand	., standardis	ed; CPU	E, catch	per unit	ettort; Het.,	heterosk	edasticit	ty; IM,	informati	ion matrix			
Species	Metric	Level	Min log mean	Max log mean	Range log mean	Intercept_ a	s.ea	$Slope_b$	s.eb	Curvature_c	SE_c	Adj. $R^2$	d.f.	Fest_a_0	Test_c_0	Swilk	Het.	IM
Wild boar	1. Crude catch	Public use	-1.200	0.477	1.677	0.111	0.046	1.016	0.067	-0.246	0.172	0.865	35	0.021	0.161	0.664	0.975	0.408
	1. Crude catch	Restricted	-0.869	-0.012	0.857	0.064	0.048	0.972	0.096	-0.510	0.409	0.737	35	0.189	0.220	0.726	0.028	0.062
	2. Stand. catch	Public use	-1.447	0.477	1.924	0.186	0.041	1.112	0.049	-0.148	0.093	0.938	35	< 0.001	0.121	0.308	0.819	0.590
	2. Stand. catch	Restricted	-1.183	-0.097	1.086	0.143	0.053	1.101	0.081	-0.513	0.238	0.842	34	0.011	0.039	0.265	0.082	0.053
	3. Crude CPUE	Public use	-1.930	-0.620	1.310	-0.071	0.239	1.343	0.177	0.865	0.486	0.612	35	0.769	0.084	< 0.001	< 0.001	< 0.001
	3. Crude CPUE	Restricted	-1.640	-0.693	0.947	-0.179	0.193	1.380	0.160	0.409	0.528	0.671	35	0.360	0.444	0.001	0.880	0.529
	4. Stand. CPUE	Public use	-1.920	-0.176	1.744	-0.403	0.142	1.065	0.099	-0.418	0.245	0.762	35	0.008	0.096	0.162	0.921	0.257
	4. Stand. CPUE	Restricted	-1.901	-0.699	1.202	-0.298	0.115	1.198	0.091	-0.379	0.239	0.833	34	0.014	0.122	0.472	0.383	0.137
Axis deer	1. Crude catch	Public use	-1.107	0.301	1.408	0.263	0.040	1.260	0.082	0.543	0.205	0.868	35	< 0.001	0.012	< 0.001	0.006	< 0.001
	1. Crude catch	Restricted	-1.290	-0.067	1.223	0.306	0.055	1.284	0.082	0.168	0.229	0.870	36	< 0.001	0.467	0.001	0.002	0.003
	2. Stand. catch	Public use	-1.447	0.097	1.544	0.219	0.043	1.118	0.061	0.326	0.142	0.904	35	< 0.001	0.028	<0.001	0.114	0.003
	2. Stand. catch	Restricted	-1.362	-0.163	1.199	0.281	0.063	1.202	0.073	-0.281	0.236	0.880	36	< 0.001	0.241	< 0.001	< 0.001	0.001
	3. Crude CPUE	Public use	-1.989	-0.767	1.222	-0.239	0.148	1.208	0.128	-0.389	0.321	0.710	35	0.114	0.234	0.006	0.569	0.190
	3. Crude CPUE	Restricted	-2.088	-0.824	1.264	-0.158	0.139	1.341	0.098	0.612	0.251	0.835	36	0.262	0.020	0.015	0.572	0.007
	4. Stand. CPUE	Public use	-1.972	-0.681	1.291	-0.356	0.105	1.072	0.081	0.077	0.260	0.829	35	0.002	0.769	<0.001	0.621	0.053
	4. Stand. CPUE	Restricted	-1.964	-0.855	1.108	-0.117	0.134	1.283	0.090	0.132	0.302	0.846	36	0.389	0.665	0.002	< 0.001	< 0.001

Table 3. Linear regression estimates (by ordinary least-squares) of the parameters of the spatial Taylor's law (TL)  $\log_{10} v = a + b \times \log_{10} m$ , with m = sample mean, v = sample variance ofvarious metrics of the relative abundance of wild boar, axis deer and both species combined (including killed and wounded specimens, or excluding wounded prey) at El Palmar National Park,

On average, across 36–38 trimesters with data (depending on zone and metric), the back-transformed mean crude catch (per hunting-party session) and crude CPUE of wild boar in the restricted zone were 43% (t = 2.5, 66.5 d.f., P = 0.015) and 34% (t = 2.3, 73.9 d.f., P = 0.023) greater than those in the public-use zone respectively (Fig. 4*a*). Conversely, the mean crude catch and crude CPUE of axis deer were 60% (t = 3.4, 75.9 df, P = 0.001) and 87% (t = 4.6, 74.0 df, P < 0.001) greater in the public-use zone respectively (Fig. 4*b*). Standardised catch and standardised CPUE displayed similar patterns and narrower differences between zones for each ungulate species taken separately.

# Hunting shift

When we separated the data from the diurnal hunting shifts from those from the overnight hunting shifts, TL was not rejected for any metric or either species, except crude CPUE of axis deer in diurnal shifts (Table 4), which demonstrated statistically significant convexity. Hence, it was valid to compare these shifts using ANCOVA in all cases except crude CPUE for axis deer. ANCOVA found no significant differences in the slopes and intercepts of the spatial TL between hunting shifts (diurnal versus overnight sessions) for crude or standardised catch and standardised CPUE of both ungulates taken separately (Table 4, Fig. 5). For wild boar, the slopes of TL slightly varied between diurnal and overnight shifts for most metrics (e.g. for crude catch, from 1.080 to 1.086 respectively; Table 4, Fig. 5a, c). The hunting shift weakly modified the slope of crude CPUE (P = 0.018) for wild boar. For axis deer, standardised catch and standardised CPUE coefficients for overnight shifts tended to exceed those for diurnal shifts (Table 4, Fig. 5b, d). Nearly all intercept estimates for crude or standardised catch of axis deer were significantly different from 0.

Taylor's law closely fitted the data in 11 of 16 cases (adj.  $R^2 > 0.75$ ), with looser fits for standardised CPUE of deer in overnight shifts (adj.  $R^2 = 0.557$ ). As mentioned above, TL was rejected for crude CPUE of deer in diurnal shifts, and the adj.  $R^2 = 0.449$  of TL was notably lower than was the adj.  $R^2 = 0.574$  of QTL, as expected (Table 4, Fig. 5). As for wild boar, residuals for half of the models of axis deer showed significant deviations from normality according to the Shapiro–Wilk test, and rarely deviated significantly from homoskedasticity and normal kurtosis.

On average, across the 10-year period, the back-transformed mean crude catch of wild boar in overnight shifts was 39% greater than that in diurnal shifts (which were much shorter), but the difference was not statistically significant (t = -1.7, 25.05 d.f., P = 0.11; Fig. 6a). Similarly, the mean crude catch of axis deer per hunting-party session was 36% greater in overnight shifts, but again the difference between shifts was not significant (t = 1.7, 22.6 d.f., P = 0.10; Fig. 6b). However, in terms of crude CPUE, the mean indices were marginally greater in diurnal shifts both for wild boar (by 38%; t = 1.7, 29.16 d.f., P = 0.098) and axis deer (by 39%; t = 1.9, 22.17 d.f., P = 0.074). For deer, both standardised catch and standardised CPUE weakly favoured diurnal shifts (P = 0.025).

# Effects of time scale on Taylor's law parameters

Tables S2 and S3 provide the same statistics as Tables 2–4, but computed for monthly periods rather than trimesters. The spatial



**Fig. 3.** The spatial TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the crude catch per unit effort of exotic mammals, under conditions of controlled shooting from a fixed hunting blind located in the public-use zone (triangles, red regression line) or restricted zone (diamonds, green regression line) of the park. Each point represents the spatial mean and spatial variance of crude catch per unit effort for (*a*) wild boar and (*b*) axis deer across El Palmar National Park at one trimester over 2006–2015. Key as in Fig. 1.



**Fig. 4.** Back-transformed spatial mean and 95% confidence intervals of the relative abundance of (*a*) wild boar and (*b*) axis deer by four metrics ( $\times$ 100) according to hunting zone (public use versus restricted use) across El Palmar National Park at one trimester over 2006–2015.

TL was not rejected for either species and for any metric except wild boar using standardised catch or crude CPUE for all data pooled (Table S2), and for wild boar crude CPUE in the diurnal shift only (Table S3).

In general, the choice of time scale affected little the slope coefficients for each metric, but estimates based on monthly periods were nearly always greater for axis deer and for the restricted zone (Table S2). Standard errors were smaller, and the fit of the models improved slightly in most cases.

The choice of a monthly time scale modified the effects of hunting shift and zone on some spatial TL parameters. ANCOVA detected highly significant differences in the slope, but not in the intercept, of TL between hunting shifts for crude CPUE of wild boar, and in the slope and intercept of standardised catch for axis deer only. Regarding human disturbance effects, ANCOVA yielded no significant differences in the parameters of the spatial TL between public- and restricted-use zones for any of the four indices of wild boar. However, for axis deer, both zone and the interaction term zone  $\times$  log mean weakly (0.01 < P < 0.05) increased log sample variance of crude and standardised catch only.

All indices of relative abundance computed for 1-month periods differed more strongly between hunting zones and shifts than did those computed for 3-month periods and maintained the direction of differences between categorical levels. Wild boar was highly significantly more abundant in the restricted-use (southern) zone by the four metrics, whereas axis deer was captured more frequently in the public-use zone according to all metrics, with differences being highly significant in crude CPUE only (P < 0.001). Regarding hunting shift effects, the crude (P = 0.002) and standardised catch (P = 0.017) of wild boar and its standardised CPUE (P = 0.002) in overnight sessions significantly exceeded the corresponding indices recorded in diurnal sessions. For deer, all indices were significantly (0.001 < P < 0.002) greater in diurnal sessions, except crude catch, which was greater in overnight sessions, although not significantly so.

					20	06-2015, ac	cording	to hunti	ing shift									
	1	Key as in T	able 2. Catch, (	catch per huntin	g-session; Stand	., standardis	ed; CPU	JE, catch	per unit	effort; Het.,	heterosk	edasticit	y; IM,	informati	ion matrix			
Species	Metric	Level	Min log mean	Max log mean	Range log mean	Intercept_ a	s.e_a	Slope_b	s.eb	Curvature_c	s.ec	Adj. $R^2$	d.f.	Test_a_0	Test_c_0	Swilk	Het.	IM
Wild boar	1. Crude catch	Diurnal	-1.230	-0.234	0.996	0.156	0.047	1.080	0.071	-0.302	0.197	0.874	32	0.002	0.136	0.212	0.990	0.562
	1. Crude catch	Overnight	-1.322	0.000	1.322	0.120	0.065	1.086	0.109	-0.145	0.267	0.845	17	0.082	0.595	0.197	0.089	0.426
	2. Stand. catch	Diurnal	-1.591	-0.378	1.213	0.239	0.040	1.172	0.047	0.007	0.114	0.950	32	< 0.001	0.951	0.007	0.340	0.701
	2. Stand. catch	Overnight	-1.562	-0.121	1.441	0.178	0.091	1.151	0.114	-0.097	0.266	0.885	12	0.076	0.723	0.005	0.112	0.142
	3. Crude CPUE	Diurnal	-1.937	-0.740	1.198	0.197	0.173	1.539	0.132	0.735	0.294	0.803	32	0.263	0.018	0.005	< 0.001	< 0.001
	3. Crude CPUE	Overnight	-2.284	-1.028	1.256	-0.605	0.247	1.151	0.169	-0.244	0.422	0.716	17	0.026	0.571	0.001	0.411	0.448
	4. Stand. CPUE	Diurnal	-2.244	-0.921	1.323	-0.113	0.090	1.276	0.061	0.213	0.132	0.929	32	0.217	0.116	0.049	0.689	0.410
	4. Stand. CPUE	Overnight	-1.811	-0.857	0.954	-0.413	0.270	1.076	0.202	-0.525	0.791	0.677	12	0.152	0.520	0.025	0.467	0.131
Axis deer	1. Crude catch	Diurnal	-0.961	-0.186	0.775	0.211	0.048	1.152	0.098	0.216	0.389	0.805	32	< 0.001	0.583	0.073	0.737	0.475
	1. Crude catch	Overnight	-0.848	0.142	0.990	0.212	0.062	1.101	0.139	0.036	0.648	0.785	16	0.003	0.956	0.055	0.962	0.441
	2. Stand. catch	Diurnal	-1.535	-0.196	1.338	0.138	0.043	0.978	0.063	-0.004	0.151	0.881	32	0.003	0.978	0.005	0.142	0.210
	2. Stand. catch	Overnight	-1.591	-0.385	1.206	0.322	0.135	1.209	0.144	-0.035	0.429	0.833	13	0.019	0.936	0.100	0.052	0.094
	3. Crude CPUE	Diurnal	-1.489	-0.838	0.651	-0.400	0.224	1.047	0.198	0	I	0.449	32	0.083	I	0.452	0.455	< 0.001
	3. Crude CPUE	Diurnal	-1.489	-0.838	0.651	3.369	1.187	7.774	2.096	2.947	0.915	0.574	31	0.008	0.003	< 0.001	0.156	0.334
	3. Crude CPUE	Overnight	-1.914	-0.761	1.153	-0.679	0.176	1.036	0.136	-0.557	0.384	0.771	16	0.001	0.167	0.448	0.886	0.707
	4. Stand. CPUE	Diurnal	-2.228	-0.895	1.333	-0.409	0.132	1.038	0.102	-0.108	0.246	0.755	32	0.004	0.664	0.003	0.641	0.033
	4. Stand. CPUE	Overnight	-1.825	-1.026	0.799	-0.260	0.385	1.126	0.261	0.033	1.167	0.557	13	0.511	0.978	0.007	0.480	0.757

Table 4. Linear regression estimates (by ordinary least-squares) of the parameters of the spatial Taylor's law (TL) log<sub>10</sub> v = a + b × log<sub>10</sub> m, with m = sample mean, v = sample variance of

various metrics of the relative abundance of wild boar, axis deer and both species combined (including killed and wounded specimens, or excluding wounded prey) at El Palmar National Park,

#### Temporal Taylor's law

All metrics for axis deer and wild boar culled at identified blinds over a variable number of trimesters were consistent with (failed to reject) TL (Table 5). Adjusted  $R^2$  ranged from 0.744 to 0.909. Three of the eight slope coefficients (8 = 2 species × 4 metrics) were significantly greater than 1 (for standardised CPUE of deer and boar and crude CPUE of boar), whereas all eight intercepts but one differed significantly from 0 (Figs 7, 8, Table 5). ANCOVA of TL parameters yielded no significant differences between ungulate species for any of the four metrics (P > 0.2).

The hunting blinds showing maximum performance (i.e. greater mean crude catch or crude CPUE among those participating in more than 80 sessions) for a given species were mostly invariant of the individual metric considered, but the exact blinds and park sections greatly differed between wild boar (Fig. 7) and deer (Fig. 8). For wild boar crude catch, the three top hunting blinds (3S, 1S, and 1N) and some immediate followers (0S and 4S) were concentrated at the park's western extreme close to a permanent water course (Arroyo El Palmar; Fig. 7). For deer crude catch, five of the top six blinds (10N, 8N, 21N, 25N and 18N) were concentrated in a stretch of 1 or 2 km of the public-use zone covered by a dense forest of exotic trees or shrubs adjacent to the river coastline, on the eastern extreme of the park (Fig. 8).

## Discussion

#### Principal findings

We found that the spatial and temporal Taylor's law (TL) adequately described the relationship of the log sample variance to the log sample mean of four indices of the relative abundance of wild boar and axis deer at multiple fixed hunting sites of a protected area over a decade, except for 4 of 89 cases tested for curvilinearity of the relation. In most cases where TL held for both species and the slopes *b* could be validly compared, they did not differ significantly between wild boar and axis deer in both the spatial TL and the temporal TL, and  $\sim 1 < b < 2$ , as widely verified (e.g. Taylor *et al.* 1978, 1988).

If captures of boar and deer had been Poisson distributed across hunting blinds and hunting sessions in each trimester (or month), with different mean values in different trimesters (or months), then the slope of the spatial TL would have been expected not to differ significantly from b = 1 and the intercept of the spatial TL would have been expected not to differ significantly from a = 0, because the Poisson distribution (a commonly used standard for pure 'randomness') has a variance equal to its mean, for every value of the mean.

In our data, the slopes *b* of TL were not significantly different from 1 by three of our four metrics, but one of the most frequently used metrics (crude CPUE) was consistently greater than 1 in all cases we examined (e.g. ranges 1.307-1.669 for the spatial TL in Table 2, 1.127-1.323 for the temporal TL in Table 5), regardless of whether each ungulate species was taken separately or both were combined, and regardless of whether killed and reportedly wounded specimens were pooled (prey). Thus, the slopes of TL depended on the specific metric considered.

When TL was not rejected, tests of the null hypothesis that a = 0 rejected the null hypothesis with P < 0.01 in 5 of the 11



**Fig. 5.** The spatial TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the relative abundance of exotic mammals, under conditions of controlled shooting from a fixed hunting blind in diurnal (triangles) and overnight (diamonds) hunting shifts. Each point represents (*a*, *c*) the spatial mean and spatial variance of crude catch per hunting-party session and (*b*, *d*) crude catch per unit effort for (*a*, *b*) wild boar and (*c*, *d*) axis deer across El Palmar National Park at one trimester over 2006–2015. Key as in Fig. 1.



**Fig. 6.** Back-transformed spatial mean and 95% confidence intervals of the relative abundance of (a) wild boar and (b) axis deer by four metrics (×100) according to hunting shift (diurnal versus overnight) across El Palmar National Park at one trimester over 2006–2015.

cases in Table 2 for the spatial TL and 8 of the 11 values in Table 5 for the temporal TL. In general, the null hypothesis that a = 0 was *not* rejected in precisely those cases where the slope

b > 1 substantially, as if the TL regressions (both spatial and temporal) had either zero intercept and steep positive slope or positive intercept and lower slope close to 1. These results were

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Key as in Table 2, but the	minimum, maxir	mum and range	log mean and lc catch <sub>j</sub>	og variance are h per unit effort; H	ere calculate let., heterosk	ad over tu cedasticit	me at eau ty; IM, ir	ch given iformati	hunting blin on matrix	d. Catch	ı, catch <u></u>	er hur	tting-sessi	on; Stand.	, standarc	lised; C	PUE,
Species	Metric	Min log mean	Max log mean	Range log mean	Intercept_a	s.ea	Slope_b	s.eb	Curvature_c	s.ec	Adj. $R^2$	df	Fest_a_0	Test_c_0	Swilk	Het.	M
Wild boar	1. Crude catch	-1.362	-0.182	1.180	0.163	0.035	1.085	0.052	-0.244	0.149	0.909	43	<0.001	0.110	0.223	0.475	0.080
	2. Stand. catch	-1.342	-0.255	1.088	0.178	0.046	1.090	0.057	-0.242	0.169	0.893	43	< 0.001	0.161	0.578	0.885	0.339
	3. Crude CPUE	-2.081	-0.856	1.225	-0.109	0.162	1.323	0.117	0.105	0.317	0.744	43	0.504	0.741	< 0.001	0.002	0.060
	4. Stand. CPUE	-2.041	-0.885	1.156	-0.299	0.102	1.149	0.071	-0.236	0.198	0.857	43	0.005	0.241	0.692	0.575	0.643
Axis deer	1. Crude catch	-1.090	-0.046	1.044	0.174	0.033	1.058	0.060	-0.082	0.206	0.867	46	< 0.001	0.694	0.091	0.111	0.265
	2. Stand. catch	-1.481	-0.058	1.423	0.158	0.039	1.040	0.057	-0.116	0.141	0.876	46	< 0.001	0.414	0.322	0.745	0.362
	3. Crude CPUE	-1.940	-0.729	1.211	-0.387	0.083	1.127	0.067	-0.144	0.184	0.857	46	< 0.001	0.439	0.208	0.571	0.456
	4. Stand. CPUE	-2.153	-0.733	1.420	-0.280	0.080	1.161	0.061	0.028	0.143	0.886	46	< 0.001	0.847	0.026	0.100	0.161
Both sp. including wounded	1. Crude catch	-0.673	0.110	0.783	0.119	0.023	1.114	0.100	0.087	0.401	0.725	46	<0.001	0.829	0.003	0.647	0.064
	2. Crude CPUE	-1.410	-0.557	0.854	-0.291	0.154	1.297	0.167	0.175	0.612	0.557	46	0.064	0.776	<0.001	0.400	0.454
Both sp. excluding wounded	1. Crude CPUE	-1.526	-0.570	0.956	-0.258	0.145	1.310	0.151	0.106	0.479	0.611	46	0.082	0.825	< 0.001	0.162	0.430

Table 5. Linear regression estimates (by ordinary least-squares) of the parameters of the temporal Taylor's law (TL) log<sub>10</sub>  $\nu = a + b \times \log_{10} m$ , with m = sample mean,  $\nu =$  sample variance of various metrics of the relative abundance of wild boar, axis deer and both species combined (including killed and wounded specimens, or excluding wounded prey) at El Palmar National Park,

robust to the choice of time scale (trimesters or monthly periods).

Either combination of TL parameters (zero intercept and slope greater than 1 or positive intercept and slope indistinguishable from 1) rejected Poisson distributions as a sufficient model for the variability in harvests. Although many factors could cause deviations from pure randomness, two mechanisms that have received much attention among ecologists are behavioural and environmental. Individual deer and boar are likely to be attracted or repelled by other individuals of the same or other species, and these attractions or repulsions are likely to change over time, for example, seasonally, resulting in spatially and temporally varying aggregations. Only foxes in the park may affect the offspring of either exotic ungulate, which lack an apex predator in the region. Axis and other deer species may escape from the immediate presence of wild boar (Tolleson et al. 1995; Latham 1999; Ferretti et al. 2008). Moreover, spatial and temporal environmental variations (e.g. availability of water, hunting pressure, food resources, and weather) are likely to induce or inhibit aggregations of deer and boar around the hunting blinds where controlled shooting was implemented.

We used four metrics to describe the relative abundance of wild boar and axis deer because crude catch or crude CPUE were sometimes considered poor indices of population density, especially in some fisheries (e.g. Walters 2003; Maunder and Piner 2015), but there is no broad consensus on this issue (Pauly et al. 2013). However, CPUE sometimes provided a valid index of wildlife density for management of deer and game birds (e.g. Novak et al. 1991; Lancia et al. 1996; Cattadori et al. 2003; Rist et al. 2010). The relative density of wild boar sighted correlated positively and significantly with other indices such as the fraction of plots with fresh feral pig dung (Hone 2012, p. 35). Here, all four metrics for log mean abundance (temporal TL) were highly significantly (P < 0.001) correlated for wild boar (range of r, 0.9090–0.988) and deer (range of r, 0.964–0.983) across all active hunting blinds. However, crude CPUE differed from all other metrics for boar and deer in that the slope coefficient was significantly greater than 1 in nearly every case, and tended to be much greater when pooled boar and deer or prey were considered in the spatial TL (b = 1.655 and 1.669). This was less obvious for the temporal TL (b = 1.114 and 1.310). These results imply that the variance in crude CPUE increased faster than in proportion to the mean CPUE. One possible interpretation is that when the local density of wild boar or deer increased or more specimens were attracted to the baits (or more bait was deployed regularly), more skilful hunters or those with better gear (or investment in bait) may have taken advantage of these opportunities, whereas less experienced hunters equipped with poorer gear (or with poorer baiting habits) continued to cull quarry at roughly the same rate. Similarly, the slopes b tended to depart more from 1 when both ungulate species were combined or wounded prey were included, again suggesting that hunters facing better chances (i.e. either species) would tend to depart more from their average performance. Standardised CPUE and standardised catch would be less affected by better conditions or chances because the restricted time window in which hunting efforts and outcomes were considered put a cap on both, and allowed fewer opportunities for improved performance; thus, as for



Fig. 7. The temporal TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the relative abundance of *wild boar*, under conditions of controlled shooting from a fixed hunting blind. Each point represents the temporal mean and temporal variance of (*a*) crude and (*b*) standardised catch per hunting party-session, (*c*) crude catch per unit effort and (*d*) standardised catch per unit effort across El Palmar National Park at one hunting blind over 2006–2015. Key as in Fig. 1.

other metrics, b was consistently around 1 or marginally different from 1 in both spatial and temporal TL.

Neither human disturbance (i.e. zone) nor the hunting shift significantly modified the parameters of the spatial TL, except in one case. However, Fig. 4 shows that, on average, across the 10-year period, the mean relative abundance of wild boar was consistently and significantly greater in the restricted zone, whereas that of axis deer was significantly greater in the public-use zone, and more specifically on the stretch of exotic trees or shrubs adjacent to the Uruguay River. Similarly, axis deer occupancy peaked in proximity to permanent water courses in Australia's dry landscapes (Forsyth et al. 2019), as wild boar did in the lower Mississippi valley, USA (e.g. Paolini et al. 2019). Both zones in the park were embedded in forest plantations but differed in the intensity of disturbance (e.g. tourismrelated vehicle use) and other features. Only the restricted zone had permanent crop fields adjacent to the border, and these were in proximity to better, more abundant refuge afforded by vegetation type, more permanent water courses and wetlands within the park. The between-zone differences in ungulate abundance were unexpected and have implications for dedicated research efforts and management (see below).

The effect of hunting shift on abundance depended on the metric of abundance. Mean crude catches of boar and deer were substantially greater in overnight shifts (which were much longer than diurnal shifts), but differences in mean crude CPUE were marginally significant and slightly favoured diurnal shifts. According to local hunters and rangers, both ungulate species apparently displayed a peak of activity right before sunset and at dawn. The available evidence does not allow us to conclude whether diurnal or overnight shifts are better for culling boar or deer, in part because in these unplanned comparisons, the exact timing of overnight shifts was quite variable over time, as was the duration of hunting sessions. These factors are likely to affect both the chance of hunter-prey encounter at the blinds and hunter performance over prolonged sessions (i.e. fatigue). Costeffectiveness and safety issues may need to be factored in, and these depend on hunters' motivations and setting. Notwithstanding these concerns, deer crude catch was the only metric that favoured overnight shifts substantially over the remaining



**Fig. 8.** The temporal TL described the relationship between  $y = \log_{10} v$  and  $x = \log_{10} m$  of the relative abundance of *axis deer*, under conditions of controlled shooting from a fixed hunting blind. Each point represents the temporal mean and temporal variance of (*a*) crude and (*b*) standardised catch per hunting party-session, (*c*) crude catch per unit effort and (*d*) standardised catch per unit effort across El Palmar National Park at one hunting blind over 2006–2015. Key as in Fig. 1.

metrics for deer (Fig. 6b). Camera-trap surveys conducted in the public-use section of the park over 2017–2018 unexpectedly showed that axis deer displayed major activity periods around 0100 hours and 0700 hours over late winter and early summer (Nicosia *et al.* 2019). On the basis of these results, we implemented a collaborative intervention trial with park management and program-affiliated hunters, in which a series of four or five diurnal hunting sessions was followed by another series of five overnight sessions by using standard hunting procedures. The trial confirmed that deer harvest measured both by crude catch and crude CPUE was three to four times greater in overnight sessions (Gürtler *et al.*, unpubl. report to Administración de Parques Nacionales 2019).

The hunting blinds with top mean crude catch or CPUE for a given species over a decade exceeded the rest by at least an order of magnitude, and differed between wild boar (Fig. 7) and axis deer (Fig. 8). Both the top blinds and their immediate followers were concentrated in exactly opposite stretches of the park: axis deer closer to the river coastline on the eastern extreme, and wild boar on the western extreme by the entry of a permanent water

course (believed to be a corridor for boar from the upper watershed) with a dense riverine forest in proximity to highdensity stands of palm trees. The latter shed an enormous amount of fruit from mid-summer to mid-fall, which is readily consumed by wild boar (Ballari *et al.* 2015). These substantial differences are even more noteworthy given the 10-year-long period of observations, during which the holders of local blinds experienced variable rates of turnover, improvement of hunting gear, and long-term landscape change (i.e. lignification). As explained above, various sources of heterogeneity in the local environment may explain the differences in relative abundance between zones, some of them of anthropic origin.

We tested TL by using a large amount of relative abundance data that have some limitations. Several hunting-related factors (e.g. bait, night-vision gear, hunters' experience, weather) are expected to affect every relative abundance metric via their influence on hunting effectiveness. Hence, slight deviations from linearity in TL should not be given too much weight. Unlike in another multi-site study in which the data analysed for TL had been collected systematically according to defined sampling protocols (e.g. Cohen *et al.* 2017*b*), here, how each hunting party operated varied widely in several key aspects. Wild boar was hunted with dogs over the first 2 years after program onset, and this may have affected (positively or negatively) the effectiveness of controlled shooting. Virtually all outlier trimesters corresponded to fall and summer, and none occurred in winter. Fog is more frequent over the fall season, mostly affecting wetland sections of the park and permanent streams, and preventing hunters from sighting the prey; this would explain widely variable catches at some locations. Summer trimesters had reduced or nil coverage of hunting sessions from year 7 to 9.

#### Implications for management of invasive species

Taylor's law identifies key zones or sections with a high mean and high variance of boar or deer relative density. We are not aware that TL has been used to show the most productive hunting spots of any wildlife species. If the goal is wildlife control or elimination, these 'hot spots' and associated habitats should be preferentially targeted with adequate methods. Locations or habitats with exceptionally low mean density and low variance (i.e. in the lower extreme of the prediction interval) entail stable, low-density local populations, whereas spots of exceptionally high mean density and high variance may represent either sources or sinks that demand adequate prioritisation. Even for locations that lie within the prediction interval of the log-log regression, high mean catches or CPUE with a low or high variance over time may signal different processes, such as focal immigration of exotic ungulates to the protected area, inconsistent baiting practices or poaching (i.e. local depletion of the target species by unauthorised third parties). For example, local hunters and rangers recurrently blamed poachers for preemptive culling of wild boar at the western extreme of the restricted zone, which translated into large swings in the harvest achieved by program hunters. According to these sources, poaching activity has decreased over time in direct correspondence to the decline in wild boar abundance caused by program activities, increased with recurrent economic crises, and likely shifted to the increasingly abundant axis deer and capybaras. In the past, poachers used dogs, horses and knives to hunt wild boar when it was abundant. Alternatively, successful program hunters may locally deplete the target species so that a period of time is required for local repopulation.

The greater abundance of wild boar in the restricted zone (with greater conservation value) entails greater risks of predation of yatay palm tree saplings and would strongly reduce its recruitment, one of the main goals of the management program and main reasons why it was created and has been sustained. Identifying the functional relationship between wild boar abundance and palm tree recruitment at local levels would provide valuable information and improved metrics for performance monitoring of the program, especially in the restricted zone.

Taylor's law can be used to design sequential sampling to attain a fixed-precision estimate of wildlife abundance, as was recently undertaken for fish assemblages in a lake in Africa (Xu *et al.* 2019). This framework and TL can be used to sample wild boar or deer more efficiently. To apply this method to the boar and deer data, the mean and the variance have to be calculated cumulatively, starting with the first 5 or 10 observations, and

then incrementing with each additional observation. Here an 'observation' could mean a single hunting session, or a trimester (including all the hunting sessions in that trimester), for example. A cumulative sample size curve is plotted, where x = the cumulative number of observations, and y = the cumulative number of boar (or deer) harvested by the *x* observations. When a certain line constructed from the parameters of the cumulative TL and from the desired fixed precision (see eqn 2 in Xu *et al.* 2019) intersects the cumulative sample size curve, the desired fixed precision has been attained.

In addition to fixed-precision sequential sampling, TL can be used to establish an upper and a lower 'economic injury limit' (EIL) that determines whether to start control measures (above the upper EIL), stop control measures and sampling (below the lower EIL), or continue sampling to improve the estimated abundance (between the lower EIL and the upper EIL). For wild boar, the maximum extent of ground rooting (one of the typical damages) allowed may represent the EIL. Given the positive relationship between two indices of wild boar relative abundance and ground rooting (e.g. Hone 2002, 2012, pp. 35, 59), the threshold CPUE at which control measures should be started is relevant and needs to be identified with the required precision.

## Questions for future research

As with hunting blind locations, TL may also help identify key habitats where intensified control measures may be more costeffective. The underlying drivers for heterogeneities in wild boar and deer abundance remain to be identified and have implications for improved management.

The exotic ungulate management program was based on the hidden assumption that the catches of wild boar and axis deer are independent, in the sense that shooting specimens from one species would not affect the chance of sighting, shooting and dispatching specimens from the second species. However, shooting is known to affect the behaviour of ungulates, and more specifically, wild boar displays elaborate evasive tactics when confronting hunting (Ohashi *et al.* 2013; Thurfjell *et al.* 2013), as do various deer species when confronting wild boar (Latham 1999). Whether the actual catch of hunting parties is spatially dependent cannot be inferred from our current analyses. These issues may be relevant for TL and improved management.

# **Conflicts of interest**

The authors declare no conflicts of interest.

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#### References

Ballari, S. A., Cuevas, M. F., Ojeda, R. A., and Navarro, J. L. (2015). Diet of wild boar (*Sus scrofa*) in a protected area of Argentina: the importance of baiting. *Mammal Research* 60, 81–87. doi:10.1007/s13364-014-0202-0

- Barnes, R. F., Beardsley, K., Michelmore, F., Barnes, K. L., Alers, M. P., and Blom, A. (1997). Estimating forest elephant numbers with dung counts and a geographic information system. *The Journal of Wildlife Management* 61, 1384–1393. doi:10.2307/3802142
- Barrios-García, M. N., and Ballari, S. A. (2012). Impact of wild boar (Sus scrofa) in its introduced and native range: a review. Biological Invasions 14, 2283–2300. doi:10.1007/s10530-012-0229-6
- Batista, W. B., Mochi, L. S., and Biganzoli, F. (2018). Cattle decrease plant species diversity in protected humid temperate savanna. *Phytocoenolo*gia 48, 283–295. doi:10.1127/phyto/2018/0244
- Campbell, T. A., and Long, D. B. (2009). Feral swine damage and damage management in forested ecosystems. *Forest Ecology and Management* 257, 2319–2326. doi:10.1016/j.foreco.2009.03.036
- Carpio, A. J., Apollonio, M., and Acevedo, P. (2021). Wild ungulate overabundance in Europe: contexts, causes, monitoring and management recommendations. *Mammal Review* **51**, 95–108. doi:10.1111/mam.12221
- Cattadori, I. M., Haydon, D. T., Thirgood, S. J., and Hudson, P. J. (2003). Are indirect measures of abundance a useful index of population density? The case of red grouse harvesting. *Oikos* **100**, 439–446. doi:10.1034/j. 1600-0706.2003.12072.x
- Chébez, J. C., and Rodríguez, G. (2014). La fauna gringa: especies introducidas en la Argentina. (Fundación de Historia Natural Félix de Azara: Buenos Aires, Argentina.)
- Choquenot, D., McIlroy, J., and Korn, T. (1996). Managing vertebrate pests: feral pigs. (Bureau of Resource Sciences, Australian Government Publishing Service: Canberra, ACT, Australia.)
- Clark, C. W. (1985). Bioeconomic modelling and fisheries management. (Wiley: New York, NY, USA.)
- Cohen, J. E. (2013). Taylor's power law of fluctuation scaling and the growth-rate theorem. *Theoretical Population Biology* 88, 94–100. doi:10.1016/j.tpb.2013.04.002
- Cohen, J. E., and Xu, M. (2015). Random sampling of skewed distributions implies Taylor's power law of fluctuation scaling. *Proceedings of the National Academy of Sciences of the United States of America* 112, 7749–7754. doi:10.1073/pnas.1503824112
- Cohen, J. E., Xu, M., and Brunborg, H. (2013a). Taylor's law applies to spatial variation in a human population. *Genus* 69, 25–60.
- Cohen, J. E., Xu, M., and Schuster, W. S. (2013b). Stochastic multiplicative population growth predicts and interprets Taylor's power law of fluctuation scaling. *Proceedings of the Royal Society B. Biological Sciences* 280, 20122955. doi:10.1098/rspb.2012.2955
- Cohen, J. E., Lai, J., Coomes, D. A., and Allen, R. B. (2016). Taylor's law and related allometric power laws in New Zealand mountain beech forests: the roles of space, time and environment. *Oikos* 125, 1342–1357. doi:10.1111/oik.02622
- Cohen, J. E., Poulin, R., and Lagrue, C. (2017a). Linking parasite populations in hosts to parasite populations in space through Taylor's law and the negative binomial distribution. *Proceedings of the National Academy* of Sciences of the United States of America 114, E47–E56. doi:10.1073/ pnas.1618803114
- Cohen, J. E., Rodríguez-Planes, L. I., Gaspe, M. S., Cecere, M. C., Cardinal, M. V., and Gürtler, R. E. (2017b). Chagas disease vector control and Taylor's law. *PLoS Neglected Tropical Diseases* 11, e0006092. doi:10.1371/journal.pntd.0006092
- Davis, N. E., Bennett, A., Forsyth, D. M., Bowman, D. M. J. S., Lefroy, E. C., Wood, S. W., Woolnough, A. P., West, P., Hampton, J. O., and Johnson, C. N. (2016). A systematic review of the impacts and management of introduced deer (family Cervidae) in Australia. *Wildlife Research* 43, 515–532. doi:10.1071/WR16148
- Dolman, P. M., and Wäber, K. (2008). Ecosystem and competition impacts of introduced deer. Wildlife Research 35, 202–214. doi:10.1071/WR07114
- Döring, T. F., Knapp, S., and Cohen, J. E. (2015). Taylor's power law and the stability of crop yields. *Field Crops Research* 183, 294–302. doi:10.1016/j.fcr.2015.08.005

- Eisler, Z., Bartos, I., and Kertész, J. (2008). Fluctuation scaling in complex systems: Taylor's law and beyond. *Advances in Physics* 57, 89–142. doi:10.1080/00018730801893043
- Ferretti, F., Sforzi, A., and Lovari, S. (2008). Intolerance amongst deer species at feeding: roe deer are uneasy banqueters. *Behavioural Processes* 78, 487–491. doi:10.1016/j.beproc.2008.02.008
- Forsyth, D. M., Pople, A., Woodford, L., Brennan, M., Amos, M., Moloney, P. D., Fanson, B., and Story, G. (2019). Landscape scale effects of homesteads, water, and dingoes on invading chital deer in Australia's dry tropics. *Journal of Mammalogy* **100**, 1954–1965. doi:10.1093/jmammal/ gyz139
- Gamelon, M., Gaillard, J. M., Servanty, S., Gimenez, O., Torgo, C., Baubet, E., Klein, F., and Lebreton, J. D. (2012). Making use of harvest information to examine alternative management scenarios: a body weight structured model for wild boar. *Journal of Applied Ecology* 49, 833–841. doi:10.1111/j.1365-2664.2012.02160.x
- Gogan, P. J., Barrett, R. H., Shook, W. W., and Kucera, T. E. (2001). Control of ungulate numbers in a protected area. *Wildlife Society Bulletin* 29, 1075–1088.
- Gürtler, R. E., Izquierdo, V. M., Gil, G., Cavicchia, M., and Maranta, A. (2017). Coping with wild boar in a conservation area: impacts of a 10year management program of *Sus scrofa* in north-eastern Argentina. *Biological Invasions* 19, 11–24. doi:10.1007/s10530-016-1256-5
- Gürtler, R. E., Rodríguez-Planes, L. I., Gil, G., Izquierdo, V. M., Cavicchia, M., and Maranta, A. (2018). Differential long-term impacts of a management control program of axis deer and wild boar in a protected area of north-eastern Argentina. *Biological Invasions* 20, 1431–1447. doi:10.1007/s10530-017-1635-6
- Hess, S. C., Muise, J., and Schipper, J. (2015). Anatomy of an eradication effort. Removing Hawaii's illegally introduced deer. *The Wildlife Professional* 9, 26–29.
- Hilborn, R., and Walters, C. J. (1992). Quantitative fisheries stock assessment. Choice, dynamics and uncertainty. (Chapman and Hall: New York.)
- Hone, J. (2002). Feral pigs in Namadgi National Park, Australia: dynamics, impacts and management. *Biological Conservation* **105**, 231–242. doi:10.1016/S0006-3207(01)00185-9
- Hone, J. (2012). Applied population and community ecology: the case of feral pigs in Australia. (Wiley-Blackwell: West Sussex, UK.)
- Keeling, M. J., and Grenfell, B. T. (1999). Stochastic dynamics and a power law for measles variability. *Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences* **354**, 769–776. doi:10.1098/rstb.1999.0429
- Kvasnes, M. A., Pedersen, H. C., Solvang, H., Storaas, T., and Nilsen, E. B. (2015). Spatial distribution and settlement strategies in willow ptarmigan. *Population Ecology* 57, 151–161. doi:10.1007/s10144-014-0454-1
- Lancia, R. A., Bishir, J. W., Conner, M. C., and Rosenberry, C. S. (1996). Use of catch-effort to estimate population size. *Wildlife Society Bulletin* 24, 731–737.
- Latham, J. (1999). Interspecific interactions of ungulates in European forests: an overview. *Forest Ecology and Management* **120**, 13–21. doi:10.1016/S0378-1127(98)00539-8
- Lowe, S., Browne, M., Boudjelas, S., and De Poorter, M. (2004). 100 of the World's Worst Invasive Alien Species: a Selection from the Global Invasive Species Database. (Invasive Species Specialist Group: Auckland, New Zealand.)
- Massei, G., Kindberg, J., Licoppe, A., Gacic, G., Šprem, N., Kamler, J., Baubet, E., Hohmann, U., Monaco, A., Ozoli, J., Cellina, S., Podgórski, T., Fonseca, C., Markov, N., Pokorny, B., Rosell, C., and Náhlik, A. (2015). Wild boar populations up, numbers of hunters down? A review of trends and implications for Europe. *Pest Management Science* **71**, 492– 500. doi:10.1002/ps.3965
- Maunder, M. N., and Piner, K. R. (2015). Contemporary fisheries stock assessment: many issues still remain. *ICES Journal of Marine Science* 72, 7–18. doi:10.1093/icesjms/fsu015

- McMahon, C. R., Bester, M. N., Burton, H. R., Hindell, M. A., and Bradshaw, C. J. (2005). Population status, trends and a re-examination of the hypotheses explaining the recent declines of the southern elephant seal *Mirounga leonina. Mammal Review* 35, 82–100. doi:10.1111/j. 1365-2907.2005.00055.x
- Miller, R. G. (1966). Simultaneous statistical inference. (McGraw-Hill: New York, NY, USA.)
- Morand, S., and Krasnov, B. R. (2008). Why apply ecological laws to epidemiology? *Trends in Parasitology* 24, 304–309. doi:10.1016/j.pt. 2008.04.003
- Nicosia, G., Rodríguez-Planes, L., and Gürtler, R. E. (2019). Patrones de actividad de ungulados exóticos del Parque Nacional El Palmar sometidos a un intenso control: implicancias para el manejo. In 'XXXII Jornadas Argentinas de Mastozoología'. Libro de Resúmenes, p. 224. (Sociedad Argentina para el Estudio de los Mamíferos (SAREM): Puerto Madryn, Argentina.)
- Novak, J. M., Scribner, K. T., Dupont, W. D., and Smith, M. H. (1991). Catch-effort estimation of white-tailed deer population size. *The Journal* of Wildlife Management 55, 31–38. doi:10.2307/3809238
- Nugent, G., and Choquenot, D. (2004). Comparing cost-effectiveness of commercial harvesting, state-funded culling, and recreational deer hunting in New Zealand. *Wildlife Society Bulletin* **32**, 481–492. doi:10.2193/0091-7648(2004)32[481:CCOCHS]2.0.CO;2
- Ohashi, H., Saito, M., Horie, R., Tsunoda, H., Noba, H., Ishii, H., Kuwabara, T., Hiroshige, Y., Koike, S., Hoshino, Y., Toda, H., and Kaji, K. (2013). Differences in the activity pattern of the wild boar *Sus scrofa* related to human disturbance. *European Journal of Wildlife Research* 59, 167– 177. doi:10.1007/s10344-012-0661-z
- Paolini, K. E., Strickland, B. K., Tegt, J. L., VerCauteren, K. C., and Street, G. M. (2019). The habitat functional response links seasonal third-order selection to second-order landscape characteristics. *Ecology and Evolution* 9, 4683–4691. doi:10.1002/ece3.5072
- Pauly, D., Hilborn, R., and Branch, T. A. (2013). Fisheries: does catch reflect abundance? *Nature* 494, 303–306. doi:10.1038/494303a
- Pedrosa, F., Salerno, R., Padilha, F. V. B., and Galetti, M. (2015). Current distribution of invasive feral pigs in Brazil: economic impacts and ecological uncertainty. *Nature Conservation* 13, 84–87.
- Pople, A. R., Phinn, S. R., Menke, N., Grigg, G. C., Possingham, H. P., and McAlpine, C. L. (2007). Spatial patterns of kangaroo density across the South Australian pastoral zone over 26 years: aggregation during drought and suggestions of long distance movement. *Journal of Applied Ecology* 44, 1068–1079. doi:10.1111/j.1365-2664.2007.01344.x
- Rist, J., Milner-Gulland, E. J., Cowlishaw, G. U., and Rowcliffe, M. (2010). Hunter reporting of catch per unit effort as a monitoring tool in a bushmeat-harvesting system. *Conservation Biology* 24, 489–499. doi:10.1111/j.1523-1739.2010.01470.x
- Ruiz Selmo, R., Minotti, P. G., Scopel, A., and Parimbelli, M. (2007). Análisis de la heterogeneidad fisonómico-funcional de la vegetación del Parque Nacional El Palmar y su relación con la invasión por leñosas exóticas. In 'Teledetección–Hacia un mejor entendimiento de la dinámica global y regional'. pp. 257–263. (Ed. Martin: Buenos Aires, Argentina.)
- Skalski, J. R., Ryding, K. E., and Millspaugh, J. J. (2005). Wildlife demography. Analysis of sex, age, and count data. (Elsevier Academic Press: Boston, MA, USA.)

- Snow, N. P., Jarzyna, M. A., and VerCauteren, K. C. (2017). Interpreting and predicting the spread of invasive wild pigs. *Journal of Applied Ecology* 54, 2022–2032. doi:10.1111/1365-2664.12866
- Spear, D., and Chown, S. L. (2009). Nonindigenous ungulates as a threat to biodiversity. *Journal of Zoology* 279, 1–17. doi:10.1111/j.1469-7998. 2009.00604.x
- Stata Corp (2018) Stata Statistical Software: Release 15.1. (Stata Corporation: College Station, TX, USA.)
- Taylor, L. R. (1961). Aggregation, variance and the mean. *Nature* 189, 732– 735. doi:10.1038/189732a0
- Taylor, L. R., Woiwod, I. P., and Perry, J. N. (1978). The density dependence of spatial behaviour and rarity of randomness. *Journal of Animal Ecology* 47, 383–406. doi:10.2307/3790
- Taylor, L. R., Perry, J. N., Woiwod, I. P., and Taylor, R. A. J. (1988). Specificity of the spatial power-law exponent in ecology and agriculture. *Nature* 332, 721–722. doi:10.1038/332721a0
- Thurfjell, H., Spong, G., and Ericsson, G. (2013). Effects of hunting on wild boar Sus scrofa behavior. Wildlife Biology 19, 87–93. doi:10.2981/12-027
- Tippett, M. K., and Cohen, J. E. (2016). Tornado outbreak variability follows Taylor's power law of fluctuation scaling and increases dramatically with severity. *Nature Communications* 7, 10668. doi:10.1038/ ncomms10668
- Tolleson, D. R., Pinchak, W. E., Rollins, D., and Hunt, L. J. (1995). Feral hogs in the rolling plains of Texas: perspectives, problems, and potential. In 'Proceedings of the twelfth Great Plains wildlife damage control workshop'. (Eds R. E. Masters, and J. G. Huggins) pp. 124–128. (Noble Foundation: Ardmore.)
- Vetter, S. G., Ruf, T., Bieber, C., and Arnold, W. (2015). What is a mild winter? Regional differences in within-species responses to climate change. *PLoS One* 10, e0132178. doi:10.1371/journal.pone.0132178
- Walsh, P. D., White, L. J., Mbina, C., Idiata, D., Mihindou, Y., Maisels, F., and Thibault, M. (2001). Estimates of forest elephant abundance: projecting the relationship between precision and effort. *Journal of Applied Ecology* 38, 217–228. doi:10.1046/j.1365-2664.2001.00578.x
- Walters, C. (2003). Folly and fantasy in the analysis of spatial catch rate data. Canadian Journal of Fisheries and Aquatic Sciences 60, 1433–1436. doi:10.1139/f03-152
- West, B. C., Cooper, A. L., and Armstrong, J. B. (2009). Managing wild pigs: a technical guide. *Human-Wildlife Interactions Monographs* 1, 1–55.
- Willebrand, T., Hörnell-Willebrand, M., and Asmyhr, L. (2011). Willow grouse bag size is more sensitive to variation in hunter effort than to variation in willow grouse density. *Oikos* **120**, 1667–1673. doi:10.1111/ j.1600-0706.2011.19204.x
- Xu, M., Kolding, J., and Cohen, J. E. (2017). Taylor's power law and fixed precision sampling: application to abundance of fish sampled by gillnets in an African lake. *Canadian Journal of Fisheries and Aquatic Sciences* 74, 87–100. doi:10.1139/cjfas-2016-0009
- Xu, M., Kolding, J., and Cohen, J. E. (2019). Sequential analysis and design of fixed-precision sampling of Lake Kariba fishes using Taylor's power law. *Canadian Journal of Fisheries and Aquatic Sciences* 76, 904–917. doi:10.1139/cjfas-2018-0091

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