Long-term demographic dynamics of a keystone scavenger disrupted by human-induced shifts in food availability

Pablo Almaraz^a, Félix Martínez^b, Zebensui Morales-Reyes^c, José A. Sánchez-Zapata*c, Guillermo Blanco^d

^aDepartment of Ecology and Coastal Management, Instituto de Ciencias Marinas de Andalucía, ICMAN-CSIC, Campus Río San Pedro, 11510, Puerto Real, Spain.

 $^{\rm b}$ Escuela Internacional de Doctorado, Universidad Rey Juan Carlos (URJC), C/ Tulipán s/n, E-28933 Móstoles, Madrid, Spain.

^cDepartment of Applied Biology, Centro de Investigación e Innovación Agroalimentaria y Agroambiental (CIAGRO-UMH), Miguel Hernández University of Elche, Avda. de la Universidad, s/n, 03202 Elche, Spain.

^dDepartment of Evolutionary Ecology, Museo Nacional de Ciencias Naturales (MNCN-CSIC), José Gutiérrez Abascal 2, 28006 Madrid, Spain.

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^{*}Corresponding Author: José A. Sánchez-Zapata. E-mail: toni@umh.es

Abstract.Scavenging is a key ecological process controlling energy flow in ecosys-1 tems and providing valuable ecosystem services worldwide. As long-lived species, the demographic dynamics of vultures can be disrupted by spatio-temporal fluctuations in food availability, with dramatic impacts on their population viability and the ecosystem services provided. In Europe, the outbreak of Bovine Spongiform Encephalopathy (BSE) in 2001 prompted a restrictive sanitary legislation banning the presence of livestock carcasses in the wild at a continental scale. In long-lived vertebrate species the buffering hypothesis predicts that the demographic traits with the largest contribution to population growth rate should be less temporally variable. The BSE outbreak provides a unique opportunity to test for the impact of demographic buffering in a keystone scavenger suf-10 fering abrupt but transient food shortages. We study the 42-year dynamics (1979-2020) of one of the world's largest breeding colonies of Eurasian griffon vultures (Gyps fulvus). We 12 fitted an inverse Bayesian state-space model with density-dependent demographic rates to the time-series of stage-structured abundances to investigate shifts in vital rates and population dynamics before, during and after the implementation of a restrictive sanitary 15 regulation. Prior to the BSE outbreak the dynamics was mainly driven by adult sur-16 vival: 83% of temporal variance in abundance was explained by variability in this rate. 17 Moreover, during this period the regulation of population size operated through density-18 dependent fecundity and sub-adult survival. However, after the onset of the European 19 ban, a one-month delay in average laying date, a drop in fecundity and a reduction in 20 the number of fledglings induced a transient increase in the impact of fledgling and sub-21 adult recruitment on dynamics. Although adult survival rate remained constantly high, 22 as predicted by the buffering hypothesis, its relative impact on the temporal variance 23 in abundance dropped to 71% during the sanitary legislation and to 54% after the ban 24 was lifted. A significant increase in the relative impact of environmental stochasticity on 25 dynamics was modeled after the BSE outbreak. These results provide empirical evidence 26 on how abrupt environmental deterioration may induce dramatic demographic and dy-27 namic changes in the populations of keystone scavengers, with far-reaching impacts on 28 ecosystem functioning worldwide.

Key words: carrion; inverse demographic modelling; mad cow disease; matrix modelling; state-space modelling; scavenging; vultures.

INTRODUCTION

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Predictable food subsidies from humans are increasingly altering ecosystem structure and functioning in multiple ways, with far-reaching consequences for environmental conserva-34 tion worldwide (Oro et al. 2013). Such food subsidies are known to influence the pop-35 ulation size and trends of different generalist vertebrate species. For example, seabirds 36 usually benefit from fisheries discards (e.g., Bicknell et al. 2013), garden birds from sup-37 plementary feeding in urban areas (e.g. Fuller et al. 2008) or game species including 38 mammals and birds from diversionary and management feeding (e.g. Putman and Staines 39 2004). Scavengers are among these vertebrates that are tightly linked to food subsidies derived from human activities including shepherding, hunting and supplementary feeding 41 (Donazar 1993; Mateo-Tomás and Olea 2010; Blanco 2014; Cortés-Avizanda et al. 2016). Interactions between humans and scavengers have been closely connected since the Late Pliocene when early hominids probably competed for food with other scavengers (Moleón et al. 2014a). Nowadays, even if wild ungulate carcasses from hunting are an impor-45 tant source of food for scavengers, these species mostly rely on livestock for food worldwide (Donazar 1993; Mateo-Tomás et al. 2015; Lambertucci et al. 2009; Lambertucci 47 et al. 2018). Indeed, beyond the ecological function of eliminating wild ungulate carcasses, vertebrate scavengers provide an important ecosystem service by eliminating both 49 domestic ungulate carcasses from agricultural waste, and carcasses derived from hunting (Moleón et al. 2014b; Morales-Reyes et al. 2015) 51

Among vertebrates, vultures are one of the most threatened scavengers worldwide (Buechley and Şekercioğlu 2016). The main extrinsic threats for vultures include dietary pollutants (i.e. poisoning and veterinary drugs such as diclofenac and antibiotics), col-

lision and electrocution with electric infrastructures, direct persecution and reduction in food availability (Ogada, Keesing, and Virani 2012; Blanco et al. 2016; Buechley and Şek-57 ercioğlu 2016). Carcass availability might be subject to changes related to socioeconomic shifts in livestock production and management (e.g. the abandonment of traditional farm-59 ing practices; Olea and Mateo-Tomás 2009), rewilding processes (e.g. increase of wild ungulate populations; Cortés-Avizanda, Donázar, and Pereira 2015) or sanitary regulations 61 (Margalida et al. 2010; Blanco 2014). Thus, these changes might affect not only resource availability but also their predictability and quality (Donázar, Margalida, and Campión 63 2009; Cortés-Avizanda et al. 2012; Blanco 2014; Blanco, Junza, and Barrón 2017; Blanco et al. 2019). These factors can have a profound influence on population dynamics by 65 driving key demographic parameters ultimately determining population dynamics, age structure and reproductive performance. The life-history strategies of long-lived species 67 in fluctuating environments are predicted to evolve to reduce the temporal variability of population growth rate (Roff 2002; Doak et al. 2005). In particular, the demographic 69 buffering hypothesis (Pfister 1998) predicts that those demographic rates with a larger contribution to growth rate, and hence with a larger impact on extinction probability, 71 should be less variable across time. Since population growth rate is particularly sensitive to adult survival in long-lived species (Saether and Bakke 2000; Gaillard and Yoccoz 2003), canalized adult survival is predicted to stabilize long term dynamics (e.g., Rotella et al. 2012). In unpredictable environments, in particular in the presence of abrupt per-75 turbations on demographic rates induced by human activities, the ability of populations 76 of long-lived species to buffer those rates more strongly affecting population growth will determine their extinction probability. Nevertheless, there is little evidence on the magnitude of the effects of changes in food availability on the long-term demography and population dynamics of long-lived scavenger species (Margalida, Colomer, and Oro 2014). 80

In Europe, the outbreak of Bovine Spongiform Encephalopathy (BSE) led to the subsequent application of a restrictive sanitary legislation that critically limited the use of animal by-products not intended for human consumption (Regulation EC 1774/2002).

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This legislation banned the disposal of livestock carcasses in the field, which originated a major conservation problem, namely a new source of greenhouse gases emissions associated 86 with the destruction of carcasses in authorized plants (Morales-Reyes et al. 2015), and further affected the ecosystem services provided by scavengers (Margalida and Colomer 88 2012; Blanco 2014; Moleón et al. 2014b; Morales-Reyes et al. 2015). This conflict between sanitary and environmental policies led to an intense debate about the European disposi-90 tions that regulated the use of animal by-products and their implications for conservation of necrophagous birds (Tella 2001; Donázar, Margalida, and Campión 2009; Margalida et 92 al. 2010). In particular, virtual population modelling predicts that food shortage derived from sanitary regulations may induce rapid population declines in the Eurasian griffon vulture (Gyps fulvus) (Margalida and Colomer 2012). This colonial species is by far the most abundant and widespread obligate scavenger in Europe, and is also considered the dominant species in scavenger guilds (Cortés-Avizanda et al. 2012). Roughly, 30000-37000 pairs currently breed in Spain (> 95\% of the EU population (Del Moral and Molina 98 2018). However, there is a lack of knowledge on the long-term breeding biology and on the demographic consequences of sanitary regulation on vultures' population dynamics 100 and conservation. 101

Here, we take advantage of the long-term monitoring of one of the largest colonies of 103 Eurasian griffon vultures in Europe to analyse multi-decadal changes in the demographic dynamics of this species. The different trophic scenarios that arose after the dramatic 105 change in food availability derived from the European sanitary regulations provides an 106 excellent opportunity to conduct a natural experiment to study aspects of demography and population dynamics directly dependent on food availability that can be hardly reproduced experimentally. The main goal of this analysis will be to model the long-term dynamics of vital rates before, during and after the implementation of the European 110 sanitary regulation EC 1774/2002, in order to explore potential temporal shifts in the demographic structure induced by abrupt environmental deterioration. According to the 112 buffering hypothesis (Pfister 1998) we predict that adult survival of vultures remained 113

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constant in spite of the abrupt food shortage. In particular, within this scenario we
evaluated how a large shift in food availability derived from the application of sanitary
regulations affected, i) the population size, ii) stage structure, iii) laying dates and iv)
breeding success of Eurasian griffon vultures, and how these traits impacted upon longterm population dynamics.

Materials and methods

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Study area and fieldwork

We collected demographic data in the gorges of the Riaza River (41°31'N, 3°36'W), north 121 of Segovia Province, Central Spain. The area includes a complex of cliffs and canyons 122 where a large population of about 716 pairs of Eurasian griffon vulture were breeding 123 in 2020. We censused the colony every two or three years from 1984 to 1994 and then 124 annually until 2020; further information was obtained from the national census of 1979 and 125 other sources (http://www.naturalicante.com/mochila/Montejo/Hojas-e-Informes-censo. 126 htm; Fernández y Fernández-Arroyo 1996). We conducted five complete surveys each 127 breeding season in order to detect all pairs (Martinez, Rodriguez, and Blanco 1997). We 128 examined both partners of each breeding pair to assess whether they were morphologi-129 cally adults or sub-adults; individuals were categorized as sub-adults when they had not 130 acquired full adult appearance (at 5-6 years old) on the basis of their general body colour, 131 bill colour and, especially the colour, length and shape of the ruff feathers (Elosegui 1989; Blanco and Martinez 1996; Duriez, Eliotout, and Sarrazin 2011). The age structure of 133 pairs was recorded in three possible combinations: adult-adult, adult-sub-adult and sub-134 adult-sub-adult (Blanco and Martinez 1996; Blanco, Martinez, and Traverso 1997). To 135 control for errors in the stage-classification of the monitored individuals due to uncertain 136 ageing, we specified a state-space approach (King et al. 2010) linking the demographic 137 process to the data through an observation model (see subsection **Observation model**). 138

Breeding Eurasian griffon vultures are year-round residents in the study area. Egg

laying began in late December and the last clutches were laid by mid-March (Martínez, 141 Blanco, and Rodríguez-Martínez 1998). We conducted regular and intensive monitoring 142 throughout the breeding season to determine whether the pairs laid (breeding pairs) or do not laid (non-breeding pairs) despite they showed typical pair-bonding behavior, including 144 close contact, nest building and defense, and copulation. We observed at a distance 145 the presence of eggs in the nests, recorded the start of incubation and calculated the 146 date of hatching based on nestling age (Elosegui 1989). Taking into account all these criteria and an incubation period of 55 days we determined laying dates within 10-day 148 periods from 10 December onward (Martínez, Blanco, and Rodríguez-Martínez 1998). Nests were regularly checked to verify breeding failure or the success of each pair in 150 fledgling young; young fledged from June-August (Fargallo et al. 2018). All observations 151 were made by telescope at distances that avoided disturbing the birds in the colony. 152 Besides the long-term monitoring of breeding pairs in the colony, we focused on the behavior of 456 individuals ringed as nestlings since 1990. Of these individuals, only 154 9 bred outside the focal colony and most in colonies <50 Kms. away from there. In 155 contrast, 136 birds bred or attempted to bred in the focal colony, some for more than 20 156 years. These data point to the strong philopatry of the Eurasian griffon vulture to the 157 study area (F. Martínez and G. Blanco in prep.). These results agree with the dispersal 158 behavior of the studied species in other colonies (Zuberogoitia et al. 2013). Overall, we 159 considered breeders in this population as highly philopatric to their natal colony 160

Stage-structured density-dependent population dynamics model

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Demographic population modelling is usually conducted through the use of population projection matrices (Caswell 2001). This direct approach uses empirical estimates of vital rates, such as age-dependent fecundity and survival, to project the rate of increase of a population. In the presence of age- or stage-structured population time series, an alternative approach is to use inverse demographic modelling (Wood 1997; Caswell 2001). In this approach, vital rates are estimated through the fitting of a set of difference equations describing the life-history of a species to age- or stage-structured time

series data (Wood 1994; Caswell 2001; Gross, Craig, and Hutchinson 2002; Wielgus et 169 al. 2008). In this study, we fit an inverse Bayesian stage-structured stochastic density-170 dependent demographic model to data assembled from distance observations of multiple 171 cohorts of individuals during a 42-year period. As a non-invasive demographic approach 172 (Wielgus et al. 2008), this method allows for the estimation of demographic vital rates 173 from population-abundance data, with no need of capturing-recapturing individuals across 174 time. Although this approach has been used previously (Gross, Craig, and Hutchinson 2002; Wielgus et al. 2008), these implementations were deterministic. In contrast, here we 176 propose a fully stochastic strategy taking into account demographic and environmental stochasticity, as well as sampling error and missing data. Our Bayesian approach thus 178 allows for the efficient propagation and classification of uncertainty from the data to vital 179 rates and population growth rates. 180

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The demographic model for the Eurasian griffon vulture is based on the standard avian life cycle (Bennett and Owens 2002). This model considers three separate demographic 183 stages: fledglings, sub-adults and adults (Appendix S1: Fig. S1). Three transitions among 184 life stages and two survival probabilities define the time-relationships between stages, so a 185 set of three difference equations, including environmental and demographic stochasticity, 186 can be derived to model the life cycle. A nonlinear density-dependent function of the 187 Beverton-Holt type was included for each vital rate. This specification is suitable because 188 it is a discrete-time analogous of the continuous-time logistic equation (Bohner and Warth 189 2007), and considers an asymmetric resource partitioning among individuals in a scenario 190 of contest competition typical of colonial species (the results, however, are insensitive to 191 alternative functional specifications of density-dependence). 192

$Model\ construction$

Process model

We let $n_{f,t}$, $n_{s,t}$ and $n_{a,t}$ represent the number of fledglings, sub-adults and adults in the breeding population at time t, respectively. We stress here that the adult stage also

includes a variable proportion of sub-adults individuals that eventually paired with an 197 adult or another sub-adult; the breeding success of these mixed-aged and sub-adult pairs 198 is invariably lower that adult breeding pairs. Some adult pairs may not breed in a given year, but they are regarded as adults. We let F represent the average fecundity (mean 200 number of fledglings produced per adult per time step); be G_f (fledgling recruitment) 201 and G_s (sub-adult recruitment) the average probabilities that a fledgling and a sub-adult 202 recruits to the next stage, respectively (recruitment probabilities); and we let S_s (sub-203 adult survival) and S_a (adult survival) represent the average probabilities that a sub-adult 204 and an adult survives (remains in the same stage) to the next time step, respectively. We let β_i represent the parameters encapsulating the strength of density-dependence 206 modulating each vital rate i, and N_{t-1} the population size at time t-1 summed across 207 stages $(N_{t-1} = n_{f,t-1} + n_{s,t-1} + n_{a,t-1})$. Note that, if the parameter β_i of a given vital rate 208 is estimated to be 0, that vital rate becomes density-independent. The density-dependent demographic model can then be written as: 210

$$n_{f,t} = \frac{F}{1+\beta_1 N_{t-1}} n_{a,t-1} + \varepsilon_{f,t}$$

$$n_{s,t} = \frac{G_f}{1+\beta_2 N_{t-1}} n_{f,t-1} + \frac{S_s}{1+\beta_3 N_{t-1}} n_{s,t-1} + \varepsilon_{s,t}$$

$$n_{a,t} = \frac{G_s}{1+\beta_4 N_{t-1}} n_{s,t-1} + \frac{S_a}{1+\beta_5 N_{t-1}} n_{a,t-1} + \varepsilon_{a,t}$$
(1)

where $\varepsilon_{.,t}$ denotes environmental and demographic stochasticity impacting on each life stage (see below). The set of density-dependent difference equations in Eqn. 1 can be written in compact form as:

$$\mathbf{N}_t = \mathbf{L}\mathbf{N}_{t-1} + \boldsymbol{\varepsilon_t} \tag{2}$$

where $\mathbf{N_t} = (n_{f,t}, n_{s,t}, n_{a,t})^T$ is the vector of stage-structured population sizes and \mathbf{L} is the Lefkovitch matrix (Lefkovitch 1965) including the density-dependent vital rates for each stage:

$$\mathbf{L} = \begin{pmatrix} 0 & 0 & \frac{F}{1+\beta_1 N_{t-1}} \\ \frac{G_f}{1+\beta_2 N_{t-1}} & \frac{S_s}{1+\beta_3 N_{t-1}} & 0 \\ 0 & \frac{G_s}{1+\beta_4 N_{t-1}} & \frac{S_a}{1+\beta_5 N_{t-1}} \end{pmatrix}$$
(3)

Finally, in Eqn. 2, ε_t is a vector of sequentially independent random shocks distributed 217 according to a multivariate normal distribution with mean 0, $\varepsilon_t \sim MVN(0, \Sigma_t)$. The 218 variance-covariance matrix Σ_t is decomposed into an environmental (C) and demographic component (D_t) , $\Sigma_t = C + D_t$ (see, Mutshinda, O'Hara, and Woiwod 2011; Almaraz and 220 Oro 2011). The environmental matrix includes the variance of the stochastic environmental factors impacting on the dynamics of each life-stage in the main diagonal (σ^2) , as well 222 as the covariance terms for the pairwise (stage-by-stage) joint responses to these factors, $\zeta_{i,j}$ (for $i \neq j$), in the off-diagonal. The diagonal matrix $D_t = [\delta_f^2/n_f, \dots, \delta_a^2/n_a]^T$ reflects 224 the demographic variance affecting the transition of each demographic stage from time t-1 to t, which scales inversely with population size. See Appendix S1: Section S1 for details. 226

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Given the density-dependent Lefkovitch matrix in eqn. 3 it is possible to estimate 228 both the density-independent and density-dependent components of each vital rate. For 229 example, in eqn. 3 the parameter F is the maximum attainable fecundity at very low 230 population sizes (that is, when N_{t-1} is close to 0). Hence, it is a density-independent 231 quantity. In contrast, the density-dependent component of fecundity is $\frac{F}{1+\beta_1N^*}$, where 232 N^* is the total population size at equilibrium. This is the size at which the population 233 growth rate is 0. Indeed, it is straightforward to estimate a transient rate of increase, 234 encapsulating the realized rate at which the stage-structured population grows, and an 235 asymptotic rate of increase. This rate is, for a density-dependent model, the real part of 236 the dominant eigenvalue of the Lefkovitch matrix evaluated at the equilibrium N^* , and 237 should be 1 for a population to be stabilized through density-dependent mechanisms. See 238 Appendix S1: Section S4 for details. 239

$Observation \ model$

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The reliability of vital rates estimates depends on the correct classification of individuals according to a given demographic stage, either fledgling, sub-adult or adult. Individual 242 variations in phenotypic characteristics might introduce some error in the assignment of a 243 stage to an individual (see subsection Study area and fieldwork). To control for this we 244 introduced three observation equations, linking the real (latent, or unobserved) abundance of fledglings, sub-adults and adults, $n_{f,t}$, $n_{s,t}$ and $n_{a,t}$ in Eqn. 1, to the demographic stage 246 assignments made to every individual during the long-term monitoring of the colony. 247 We let $y_{f,t}$, $y_{s,t}$ and $y_{a,t}$ be the number of pairs assigned to the fledgling, sub-adult and 248 adult stage at time t, respectively, Then, we specified the real abundance for each stage as 249 following a Poisson distribution across time with the mean as the observed (field-assigned) 250 abundance: 251

$$y_{f,t} \sim \mathcal{P}(n_{f,t})$$

$$y_{s,t} \sim \mathcal{P}(n_{s,t})$$

$$y_{a,t} \sim \mathcal{P}(n_{a,t})$$

$$(4)$$

A Poisson distribution is suitable in this case given the discrete nature of abundance, and due to the linear scaling of the observation variance with the mean abundance. The set of equations in 4 are called observation equations, while the set of equations in 1 are called process equations. The linking of the observation and process equations is called a state-space model (King et al. 2010). This strategy efficiently separates the uncertainty arising from the observation process, in our case the uncertain assignment of a demographic stage to an individual, from the variability due to the ecological process under study. Thus, this approach is fully stochastic.

$Parameter\ specification,\ posterior\ estimation\ and\ model$ validation

The inverse state-space stage-structured model was fitted to the stage-structured time series of the Eurasian griffon vulture using Bayesian Markov Chain Monte Carlo (MCMC)

integration through Gibbs sampling (Gelman et al. 2014). We performed a review of the 264 available literature searching for empirical natural history data on the vital rates of the 265 Eurasian griffon vulture (Appendix S1: Table S1). This data was used to construct weakly 266 informative priors for all the vital rates in our model (eqn. 1), which greatly improved 267 posterior convergence of parameters and latent states. A Stochastic Search Variable Selec-268 tion scheme (SSVS; George and McCulloch 1993) was implemented to automatically set 269 to 0 the density-dependent parameters with a negligible effect on demography during the MCMC simulation (see Mutshinda, O'Hara, and Woiwod 2011; Almaraz and Oro 2011 271 for further details). As a sparsity-inducing method (see Gelman et al. 2014.) with SSVS 272 it is possible to estimate the posterior probability of inclusion of a density-dependent 273 vital rate, and therefore evaluate the Bayes factor associated to each one. This allows evaluating the evidence in favor of including a given density-dependent vital rate. Finally, 275 given the use of time-series of abundance data it is also possible to estimate the relative 276 contribution of each vital rate and stochastic component in the stage-structured model 277 to the observed temporal variance in population dynamics (see Almaraz and Oro 2011; Mutshinda, O'Hara, and Woiwod 2011). This is analogous to Life Table Response Ex-279 periments (Caswell 2010) applied to inverse, time series models. The construction of the 280 model for the priors, the specification of SSVS and the description of variance component 281 estimation are described in detail in Appendix S1: Sections S1-S3. 282

The Bayesian model was written in the JAGS language (Plummer 2003), version 4.3.0, using the R environment (version 4.1.2, RCoreTeam 2021) through the runjags package (Denwood 2016). The JAGS code is available in DataS1. Three models were fitted to separate datasets: the first one models the demographic dynamics of the Eurasian griffon vulture prior to the BSE outbreak (from 1978 to 2001), the second one models the dynamics during the term of the Regulation EC 1774/2002 (2002-2011), and the third one models the dynamics after the ban was lifted (2012-2020). Three MCMC chains with diffuse random priors were run for 10⁶ iterations for each model. Posterior estimates for parameters, latent states, missing data and variance components were obtained after

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discarding the first 5×10^5 iterations as burn-in. Standard diagnostic tests (see Gelman et al. 2014) were conducted to assess the convergence of the chains to a stationary distribution, using the package ggmcmc (Fernández-Marín 2016).

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Posterior predicted checks

To check for potential issues with parameter identifiability of the proposed modelling ap-298 proach, we generated synthetic time series from scenarios with known demographic rates 299 and stochastic effects. The state-space density-dependent demographic model (Eqns. 1 300 and 4) was then fitted to each of these time-series, and the resulting distribution of poste-301 rior parameter values were compared with the ground-truth estimates. The results of this 302 exercise suggest that the model is highly successful in recovering the original demographic 303 rates (see Appendix S1: Section S5). We also conducted posterior predictive checks on the 304 fitted model (see Gelman et al. 2014) to assess model adequacy. We simulated 50 synthetic 305 stage-structured time series from the fitted models and fit the model to each one. If the 306 posterior estimates of vital rates are identifiable, we expect that the vital rates recovered 307 by the model with the synthetic time series will approach the values of the model fitted 308 to the observed data (Gelman et al. 2014). That is, a clustering of the posterior fits to 309 simulated data around the Y=X line is indicative of parameter identifiability. 310

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RESULTS

Temporal trends and shifts in breeding parameters

From 1978 to the BSE outbreak in 2001, population numbers expanded from 128 to 334 breeding pairs in the focal colony (Fig. 1A). The proportion of breeding pairs made up by either an adult plus a sub-adult (mixed-age pairs) or two sub-adults also increased consistently from 1985 to 2001, but these positive trends ended abruptly in 2001, indicating that most sub-adults withdrew from the reproducing population (Fig. 1B). These shifts are concurrent with the onset of the BSE outbreak, with a peak in 2003 (Fig. 1A). Moreover,

a phenological advancement in mean laying date of roughly two weeks from 1985 ended 319 with an abrupt shift in 2001-2003, amounting to an average delay in one month up to 320 2011 (Fig. 1C). Another clear abrupt shift to very low breeding success took place in 2004 321 for both (Fig. 1D). These shifts in breeding structure and phenology were accompanied 322 by similar trends in the stage-structured time series (Fig. 1A). From the BSE outbreak 323 in 2001 to the end of the restrictive regulation (2011), the breeding population dropped 324 with a time lag similar to the age of first breeding in the Eurasian griffon vulture (4-5 years), while the non-breeding population stabilized. However, a particularly dramatic 326 crash was observed in the number of fledglings throughout this period (Fig. 1A). After the ban was lifted (2011-2020), mean-laying date abruptly dropped to pre-outbreak levels 328 and breeding success increased for both the adults only and adults plus sub-adults pairs.

Inverse stage-structured demographic modelling

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Model-based estimates of vital rates obtained with the state-space inverse demographic 331 model suggest a severe drop in fecundity after the BSE outbreak, from 0.629 ± 0.027 (1SD) 332 chicks per breeding adult prior to the outbreak, to 0.465 ± 0.047 during the BSE epidemic 333 (Fig. 2A). After the BSE outbreak (when the food-shortage period ended) fecundity in-334 creased again (0.487 \pm 0.034). These figures agree very well with the data on breeding 335 success obtained through the individual-based long-term monitoring of the colony (Fig. 336 1D). Sub-adult recruitment to the breeding adult population dropped during the epidemic, 337 with a correlated rise in sub-adult survival (a larger fraction of sub-adults remained as 338 such during the epidemic; Fig. 2A). Adult survival, in contrast, remained constantly high 339 during the 42-year period (0.992 ± 0.008) during the pre-BSE period, 0.991 ± 0.007 during 340 the food-shortage, and 0.985 ± 0.017 after the food-shortage; Fig. 2A).

The relative impact of vital rates on the temporal variance of the stage-structured population suffered a significant shift after the BSE outbreak: prior to 2001 adult survival explained a large proportion of the variance observed at the population level (83.1%), followed by fecundity (10.6%, Fig. 2B). During the BSE outbreak the impact of adult

survival decreased to 71.3%, and a further decrease to a 53.8% was modeled but after this event (Fig. 2B). Moreover, after the BSE outbreak the variance component of environmental stochasticity increased to 42.5%, while it was negligible during the previous periods (Fig. 2B). Overall, the shifts in demographic rates translated to shifts in the transient rate of increase, which was larger before (1.036 ± 0.004) and after (1.059 ± 0.010) than during the BSE outbreak $(0.990 \pm 0.006, \text{ Fig. 3A})$. Note that the transient rate of increase of the stage-structured population overlapped 1 during the BSE outbreak (95% Credible Interval, CI: 0.971-1.002), but neither before (95% CI: 1.028-1.044) nor after this period (95% CI: 1.040-1.079).

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$_{ extsf{57}}$ Impact of density-dependent vital rates on population stabilization

The density at equilibrium predicted by the stage-structured density-dependent model 358 differed significantly among the three time periods (Appendix S1: Section S4): $793.397 \pm$ 359 59.925 (1SD) individuals before the BSE outbreak, 153.854 ± 30.887 individuals during 360 the BSE outbreak and 398.113 ± 185.558 individuals after the BSE outbreak. Compared 361 to the observed number of individuals across time (Fig. 1A), this suggests that prior to the 362 BSE outbreak the Eurasian griffon vulture population was approaching the equilibrium density, while both during the BSE outbreak and after this period it was fluctuating above 364 its carrying capacity. The posterior asymptotic rates of increase of the density-dependent model evaluated at the population size at equilibrium were centered around 1 in all three 366 periods, consistent with a long-term stabilized population (Fig. 1B; see Appendix S1: Section S4). 368

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The probability of detecting density-dependence across the life cycle was relatively low for all time periods (Appendix S1: Table S2), ranging from 0.280 before the BSE outbreak to 0.264 during the BSE outbreak and 0.255 after the BSE outbreak. The posterior probability of density dependence was indeed very low for most vital rates during all time periods, which overall suggest weak density-dependent regulation across the life

cycle. However, the posterior probability of density-dependence in sub-adult survival be-375 fore the BSE outbreak was of 0.466, with an associated Bayes factor of 2.248. This is 376 regarded as barely worth mentioning evidence according to the Kass-Raftery scale (Ap-377 pendix S1: Table S2). Both during and after the BSE outbreak, very weak evidence for 378 density-dependence was found for sub-adult recruitment, but not survival. Due to the low 379 sample sizes of these latter periods, Type II error in the detection of density dependence 380 cannot be ruled out. Finally, breeding success of adult and mixed-aged pairs showed a slight decrease from 1985 as a likely consequence of density-dependent processes, only 382 significant for the time series of mixed-aged pairs (r = 0.66, p-value = 0.003; see Fig. 1D). 383

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The impact of density-dependence on the different vital rates can be assessed in Figure 2A. While most vital rates are only weakly depressed at equilibrium in all time periods, sub-adult survival is severely reduced when the population approaches the equilibrium population size. As expected, this is particularly the case before the BSE outbreak, when the evidence for density-dependence is significant.

Posterior predictive checks and model validation

With only 50 synthetic time series, the inverse demographic model was able to successfully recover the original vital rates estimates with simulated stage-structured data (Fig.
4A-C): most of the posterior predicted vital rates cluster around the Y=X regression line.
Importantly, fecundity and adult survival, which are the vital rates most strongly impacting on temporal dynamics (Fig. 2B) are also the vital rates more accurately recovered by
the posterior predictive checking (Fig. 4). The posterior predicted abundance time-series
agree very well with the true abundance for all demographic stages and temporal periods
(Fig. 4D-F).

DISCUSSION

Based on 42 years of monitoring of one of the largest colonies of Eurasian griffon vul-401 tures in Europe, we found that temporal, abrupt variability in food availability derived 402 from human activities and induced by an epidemic outbreak had profound effects on the 403 population dynamics of this long-lived species. The influence of food availability on the 404 dynamics of wildlife populations has been widely discussed (see review in Ostfeld and 405 Keesing 2000). Human activities have played an important role in ecosystem function-406 ing by generating anthropogenic food subsidies (Oro et al. 2013). Vertebrate scavengers 407 are one of the most susceptible group to changes in the availability of these subsidies 408 (Cortés-Avizanda et al. 2016). Several studies suggest a negative impact of food scarcity 409 arising from the application of the European sanitary policy on the populations of some 410 scavengers of conservation concern. For example, dietary changes in vultures (Donázar, 411 Cortés-Avizanda, and Carrete 2010) and other avian scavengers (Blanco 2014), as well as 412 large carnivores (Lagos and Bárcena 2015; Llaneza and López-Bao 2015; Northrup and 413 Boyce 2012), impacts on the movement patterns (Arrondo et al. 2018) or changes in de-414 mographic parameters of vultures (e.g. decrease in survival and breeding success or delay 415 in egg-laying dates; Donázar, Margalida, and Campión 2009; Martínez-Abraín et al. 2012; 416 Margalida et al. 2014; Donázar et al. 2020). However, while some studies have suggested 417 that food shortage derived from sanitary regulations may affect the Eurasian griffon vul-418 ture (Margalida and Colomer 2012), few studies have attempted to study the effects of 419 food shortages on the basic demographic parameters of this species on the long-term. 420

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Our inverse Bayesian demographic modelling approach uses long time-series of stage-structured abundance data (Wood 1994; Gross, Craig, and Hutchinson 2002) to decompose the temporal variability of population abundance into the relative impact of the constituent vital rates. Our method thus allows for the estimation of the effects of transient perturbations on the long term demographic variability and population dynamics by using only stage-structured abundance data. The results suggests that some vital rates of this keystone avian scavenger, in particular fecundity, might be very sensitive to severe

food shortages derived from the shifts in sanitary regulations (i.e., reduction of carcasses 429 availability in the field). In the Iberian Peninsula, the Eurasian griffon vulture primarily 430 depended on free-ranging livestock in the past, especially sheep in lowland areas and cattle 431 in mountain ranges (Donazar 1993). The declining trend in the abundance of extensive 432 herds over the last decades, especially of sheep and goats, along with the sanitary regula-433 tions forbidding the abandonment of cattle carcasses in the countryside, were concurrent 434 with an increase in the number of factory farms of fattening pigs and poultry and with increasing stabled conditions of ungulate livestock (Blanco 2014). As a consequence, the 436 populations of this and other vultures now largely depend on livestock carcasses from intensive exploitations in the study area and other regions across its distribution range in 438 Spain (Camiña and Montelío 2006; Donázar, Cortés-Avizanda, and Carrete 2010; Blanco 439 et al. 2019). It should be noted, though, that the application of the policy that banned 440 the abandonment of carcasses in the countryside due to the BSE crisis was not applied homogeneously across time and space, but it supposed a general crash in the availability 442 of carrion in most of the range of distribution of vultures in Spain (Donázar, Margalida, and Campión 2009; Margalida et al. 2010). In general, the application of the restrictive 444 policy was not conducted immediately after the new regulation, but it delayed a variable time period depending on regions with different government administrations. In addition, 446 the discard of carcasses in the countryside continued occurring illegally in some regions in the first years after the implementation of the sanitary policy, and still occurs, but this 448 practice was increasingly persecuted later. Overall, these factors led to a time lag between 449 the new regulation and its effective application, which was reflected in the lowest carcass 450 availability around 1-2 years after the emergence of the BSE crisis (Donázar, Margalida, 451 and Campión 2009; Blanco 2014). 452

Prior to the BSE outbreak, an advancement in the mean laying date, along with an in-454 crease in the proportion of mixed-aged and sub-adults pairs suggest that food availability was relatively high in the study colony and other areas (Blanco, Martinez, and Traverso 1997; Blanco 2014; Parra and Tellería 2004). During the implementation of the Euro-

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pean sanitary regulation a dramatic reduction of available livestock carcasses induced a 458 severe environmental deterioration that triggered the modeled structural change in stage 459 structure, productivity and phenology. Among these effects the delay in laying date and 460 the dramatic drop in fecundity were particularly large. This is likely due to the high 461 cost of reproduction and other life-history traits of vultures, evolved as a result of unpre-462 dictable food conditions (see Donazar 1993; Bennett and Owens 2002; Carrete, Donázar, 463 and Margalida 2006). In contrast, as a long-lived species, adult survival is expected to be very high and its temporal variability relatively low, owing to the canalization of this 465 vital rate (Stearns and Kawecki 1994; Saether and Bakke 2000; Pfister 1998; Sæther et al. 2013). Eurasian griffon vultures would exhibit relatively constant adult survival 467 but large plasticity in other traits like reproduction (Fargallo et al. 2018), as has been 468 described for different organisms with a slow pace of life (e.g. Bennett and Owens 2002; 469 Benton, Plaistow, and Coulson 2006; Sæther et al. 2013). Thus, the combination of fixed and variable traits might have dampened the effects of sanitary policies and allow for a 471 quick recovery of population structure and dynamics once the EU sanitary policies allowed for the disposal of carcasses (Margalida et al. 2010; Blanco 2014). Interestingly however, 473 while most of the observed and modeled demographic parameters promptly reversed to pre-outbreak values after the ban was lifted, their relative impact on population dynam-475 ics shifted across the 42-year period: the variance component associated to adult survival 476 consistently decreased across time even though this rate remained constantly high, while 477 the environmental stochastic component of the dynamics increased dramatically. This 478 suggest that the relative dynamical impacts of constant and variable vital rates across 479 time can vary in rather non-intuitive ways in stochastic environments. 480

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Several studies have shown the importance of food competition driven by density-482 dependent processes on population dynamics of long-lived birds such as gulls (see e.g. 483 Payo-Payo et al. 2016). Likewise, breeding productivity of the semi-colonial cinereous vulture (Aegypius monachus) and the territorial bearded vulture (Gypaetus barbatus) was affected by density-dependent mechanisms (Carrete, Donázar, and Margalida 2006; 486

Fernández-Bellon et al. 2016). For the Eurasian griffon vulture in our study area, ob-487 served breeding success consistently declined across time, particularly for the mixed-aged 488 pairs formed by an adult and a sub-adult. Moreover, our modeling results indicated that 489 before the BSE outbreak, density-dependent sub-adult survival likely played a role on 490 population regulation: at the onset of the sanitary regulation the population was fluctu-491 ating close to the equilibrium population size, the size at which the population growth 492 rate is 0. From this moment to the end of the time-series, the fluctuations were above the estimated equilibrium population size, particularly during the BSE outbreak. In other 494 raptor species, density-dependence also operates through sub-adult demographic stages (e.g., Carrete et al. 2006), since juveniles disperse to other areas or may skip breeding 496 at high population densities. In our case, the reduced sub-adult survival suggests that a 497 fraction of sub-adult individuals abandoned the population when the colony approached 498 the transient carrying capacity. This is reflected in the abrupt drop in the proportion of sub-adult and mixed-aged pairs during the BSE outbreak. Finally, a consistent delay in 500 laying date for adults, sub-adults and mixed-aged pairs during the term of the sanitary 501 regulation point to a severe environmental deterioration. Life history theory indeed pre-502 dicts that pulse perturbations, such as severe food shortages, may induce long-lived species 503 to reduce breeding effort and hence maintain large survival rates across time (e.g. Saether 504 and Bakke 2000). While age and sex might influence differential patterns of survival in 505 some vulture species (Sanz-Aguilar et al. 2017), data from 66 GPS-tagged adult Eurasian 506 griffon vultures throughout Spain suggest that mortality was mainly related to landscape 507 anthropization and only secondarily to sex and sub-population (Arrondo et al. 2020), but 508 no information is available for fledglings and sub-adults. 509

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Overall, our findings suggest that the changes in food availability related to shifts in European sanitary regulations have had negative consequences on key demographic parameters of the colonial Eurasian griffon vultures with significant detrimental effects on the population dynamics of this species. Vultures have coevolved with a rich but ephemeral food resource (DeVault, Rhodes, and Shivik 2003). Carcass availability and predictability

can be largely variable depending among others on the source of food (i.e. wild vs. do-516 mestic ungulates) and the processes involved (Moleón et al. 2014b). Vultures have been 517 able to cope with this variability for eons (Moleón et al. 2014a). In contrast, other en-518 vironmental changes increasing mortality such as illegal poisoning, the use of veterinary 519 drugs like diclofenac, or even collision with wind turbines (Martínez-Abraín et al. 2012; 520 Margalida et al. 2021) are known to negatively impact on vultures' populations (Green 521 et al. 2006; Serrano et al. 2020). Our results confirm the impact of domestic ungulate 522 carcasses availability as the major source of food resources for Eurasian griffon vultures in 523 Spain, in spite of the importance of wild ungulate populations (Mateo-Tomás et al. 2015; 524 Blanco et al. 2019). Nearly twenty years after the BSE outbreak, the consensus among 525 scientists, conservationists and managers led to the implementation of a new European 526 regulation (EC 142/2011), which allows farmers to leave the carcasses of livestock in the 527 field. Our results show that, although most of the changes in demographic parameters of 528 Eurasian griffon vultures promptly reversed to pre-BSE outbreak scenario after the ban 529 was lifted and the new legislation EC 142/2011 was implemented, the dynamical impacts 530 of the abrupt demographic changes can still be seen in the population more than a decade 531 after the BSE outbreak. Nevertheless, although the new and encouraging legislation rep-532 resents an important improvement in the conservation of European scavengers and the 533 environment, some aspects should still be improved to ensure the long-term conserva-534 tion of vultures (Mateo-Tomás et al. 2019; Morales-Reyes et al. 2017; Blanco et al. 2016; 535 Blanco et al. 2019). Given the tight intertwining of sanitary and conservation policies, 536 further research is needed to evaluate the demographic changes derived from the applica-537 tion of new sanitary regulations that may impact vulture conservation and the ecosystem 538 services they provide. 539

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SUPPORTING INFORMATION

- Additional supporting information may be found online at: http://onlinelibrary.
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Figure captions

Figure 1. Long-term dynamics of the abundance, breeding parameters and phenology 812 of the Eurasian griffon vulture in Central Spain from 1980 to 2020. A. Time series of 813 adult breeding pairs (red dots), non-breeding pairs made up by sub-adults (blue dots), 814 and fledglings (green dots). The impact of observation uncertainty arising from the stage 815 classification errors is shown as shaded bands. These areas encompass the 90% credi-816 ble intervals of the posterior estimates for the latent abundances of adults (red shade), 817 fledglings (green shade) and sub-adults (blue shades). The orange time-series stand for 818 the yearly reported cases of Bovine Spongiform Encephalopathy in Spain provided by the 819 Spanish Ministry of Agriculture (MAPAMA 2018). B. Proportion of breeding pairs (%) 820 formed by adults only (red dots), by an adult plus a sub-adult (green dots) and pairs 821 formed by sub-adults only (blue dots). C. Mean laying date in Julian days for the three 822 types of breeding pairs. D. Breeding success, estimated as the number of fledglings per 823 breeding pair are shown structured by stages. In all figures, the gray shaded rectangle 824 covers the time window during which the European legislation banned carrion disposal in 825 the wild. 826

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Figure 2. Demographic modelling of the Eurasian griffon vulture in Central Spain. An 828 inverse stage-structured model was fitted to the population dynamics prior to the BSE out-829 break (PreBSE, 1978-2000), for the dynamics during the term of the sanitary regulation 830 EC 1774/2002 (BSE, 2001-2011) and after the sanitary regulation was lifted (PostBSE, 831 2012-2020). A. The posterior distribution of the vital rates estimates considered in the 832 life cycle during the three time periods, shown for both the density-independent (yel-833 low) and density-dependent rates (blue, see main text); F = Fecundity; Gf = Fledglingrecruitment; Gs = Sub-adult recruitment; Ss = Sub-adult survival; and Sa = Adult sur-835 vival. B. The relative impact of the set of vital rates, and environmental and demographic 836 stochasticity, on the temporal variance of the stage-structured population (measured as 837 the % of explained variance in abundance). In the box-plot, horizontal black line stand 838 for the posterior median of each rate and stochastic component; the box represents the 839

inter-quantile range and whiskers show the 95\% percentiles. 840

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Figure 3. Rates of increase of the stage-structured Eurasian griffon vulture population during a 42-year period. A. Posterior distributions of the transient rates of increase be-843 fore the BSE outbreak (PreBSE, 1978-2001), during the BSE outbreak (BSE: 2002-2011), 844 and after the BSE outbreak (PostBSE: 2012-2020). B. Posterior distributions of the 845 asymptotic rates of increase, λ_s , predicted from the model evaluated at the equilibrium population size N^* for the three time periods (Appendix S1: Section S4). In all figures, 847 vertical dotted lines indicate the mean of each posterior distribution, and the solid black line denotes the rate of increase 1, at which the population achieves long-term stability. 849

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Figure 4. Results of the numerical experiment testing the ability of the inverse demographic model to recover the demographic dynamics of the Eurasian griffon vulture before, 852 during and after the BSE outbreak. A-C. The posterior estimates for the vital rates of 853 the inverse state-space demographic model fitted to each of the 50 posterior predicted 854 time series are plotted against the sampling posterior estimate of the real dataset used to 855 generate them during the three temporal periods. The box-plots show the median (hori-856 zontal line), inter-quantile range (box) and 95% percentiles (whiskers). The thickness of 857 the box is proportional to the posterior density of the estimates within the inter-quantile 858 range. The Y=X regression line (in black) is plotted as a reference. **D-F**. Time series of 859 the observed abundance for the three demographic stages (adults: blue dots; sub-adults: 860 green dots; fledglings: red dots) during the three demographic periods (Before, during and 861 after de BSE outbreak). The time series of the abundance predicted by the 50 posterior 862 simulated models are shown for each stage as lines (average of predicted abundance) and 863 shaded regions (95% credible interval). 864

Figure 1:

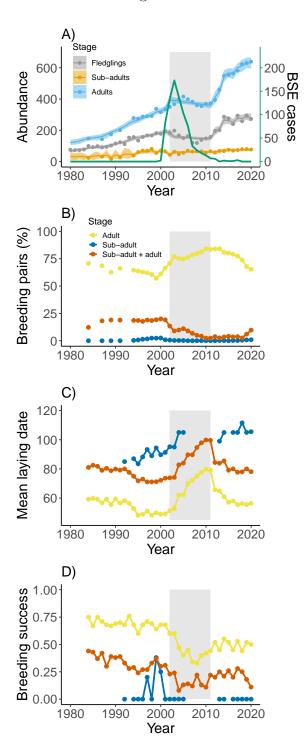


Figure 2:

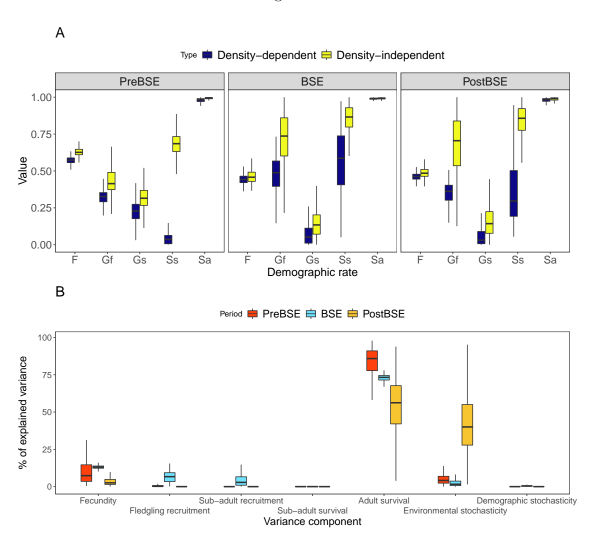


Figure 3:

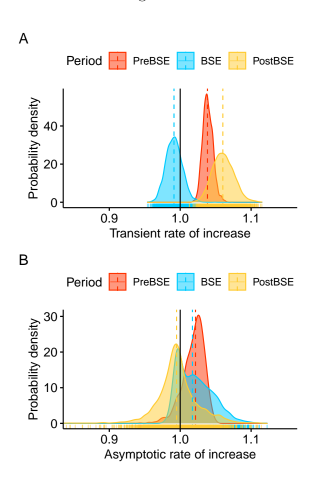


Figure 4:

