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Large-scale movement patterns in a social vulture are influenced by seasonality, sex, and breeding region

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Abstract

Quantifying space use and segregation, as well as the extrinsic and intrinsic factors affecting them, is crucial to increase our knowledge of species-specific movement ecology and to design effective management and conservation measures. This is particularly relevant in the case of species that are highly mobile and dependent on sparse and unpredictable trophic resources, such as vultures. Here, we used the GPS-tagged data of 127 adult Griffon Vultures Gyps fulvus captured at five different breeding regions in Spain to describe the movement patterns (home-range size and fidelity, and monthly cumulative distance). We also examined how individual sex, season, and breeding region determined the cumulative distance traveled and the size and overlap between consecutive monthly home-ranges. Overall, Griffon Vultures exhibited very large annual home-range sizes of 5027±2123 km², mean monthly cumulative distances of 1776 ± 1497 km, and showed a monthly home-range fidelity of $67.8 \pm 25.5\%$. However, individuals from northern breeding regions showed smaller home-ranges and traveled shorter monthly distances than those from southern ones. In all cases, home-ranges were larger in spring and summer than in winter and autumn, which could be related to difference in flying conditions and food requirements associated with reproduction. Moreover, females showed larger home-ranges and less monthly fidelity than males, indicating that the latter tended to use the similar areas throughout the year. Overall, our results indicate that both extrinsic and intrinsic factors modulate the home-range of the Griffon Vulture and that spatial segregation depends on sex and season at the individual level, without relevant differences between breeding regions in individual site fidelity. These results have important implications for conservation, such as identifying key threat factors necessary to improve management actions and policy decisions.

Jon Morant and Juan Manuel Pérez-García contributed equally to this work.

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KEYWORDS

griffon vulture, home-range, scavenger, site fidelity, spatial segregation

TAXONOMY CLASSIFICATION

Behavioural ecology

Resumen

Cuantificar el uso y la segregación del espacio, así como los factores extrínsecos e intrínsecos que los afectan, es crucial para aumentar nuestro conocimiento de la ecología de movimientos de cada especie y diseñar medidas eficaces de gestión y conservación. Esto es especialmente relevante en el caso de especies con gran movilidad y dependientes de recursos tróficos escasos e impredecibles, como son los buitres. En este trabajo se utilizaron datos GPS de 127 buitres leonados Gyps fulvus adultos capturados en cinco regiones de cría diferentes en España para describir los patrones de movimiento (tamaño y fidelidad del área de campeo y distancia acumulada mensual). También examinamos cómo el sexo, la estación del año y la región de cría determinaban la distancia acumulada recorrida y el tamaño y solapamiento entre áreas de campeo mensuales consecutivas. En conjunto, los buitres leonados mostraron un área de campeo anual muy extensa de 5027±2123km², una distancia acumulada mensual media de 1776±1497km y una fidelidad mensual al área de campeo del 67.8 ± 25.5%. Sin embargo, los individuos de las regiones de cría más septentrionales mostraron áreas de campeo más pequeñas y recorrieron distancias mensuales más cortas que los de las más meridionales. En todos los casos, las áreas de campeo fueron mayores en primavera y verano que en otoño e invierno, lo que podría estar relacionado con las diferencias en las condiciones de vuelo y las necesidades tróficas asociadas a la reproducción. Además, las hembras mostraron mayores áreas de campeo y menor fidelidad mensual que los machos, lo que indica que estos últimos tienden a utilizar zonas similares durante todo el año. En conjunto, nuestros resultados indican que tanto los factores extrínsecos como los intrínsecos modulan el área de campeo del buitre leonado y que la segregación espacial depende del sexo y de la estación a nivel individual, sin que existan diferencias relevantes entre las regiones de cría en cuanto a la fidelidad individual al lugar. Estos resultados podrían tener importantes implicaciones para la conservación, como la identificación de los principales factores necesarios para mejorar las medidas de gestión y las decisiones políticas.

1 | INTRODUCTION

Animal movements are a consequence of an organism's internal state (e.g., sex, age, breeding stage) and environmental factors (e.g., food availability or weather), and can affect individual fitness and ecological processes at local and global scales (Hansson et al., 2014). Individual movements are also influenced by inter and intraspecific relationships (e.g., competition), which may lead to spatial compartmentalization and the maintenance of population-specific movement patterns over time (Nathan et al., 2008). Deciphering how these factors modulate individual movements and how the latter are compartmentalized in space and time is essential to understanding

population dynamics (Costa-Pereira et al., 2022) and identifying priority areas for conservation and management (Katzner & Arlettaz, 2020).

Home-range size and cumulative distance traveled are key elements in the study of animal movement ecology (Kie et al., 2010; Shaw, 2020; Thaker et al., 2019; Tucker et al., 2018), defining foraging patterns at the individual and population-level, and assessing their stability over time (e.g., Shaffer et al., 2017). For example, investigating variation in home-range size and cumulative distance traveled may reveal that certain individuals behave as centralplace foragers during only a specific period of their life cycle (e.g., the breeding season; Carrete & Donázar, 2005; Delgado-González et al., 2022). Similarly, the study of home-range overlap allows the analysis of attraction or repulsion relationships that may affect space use, for example, to avoid competition for resources (Bolnick et al., 2003; Cecere et al., 2018; Winner et al., 2018). This information could be used to explicitly map intra and interspecific meeting sites and prioritize high-quality habitats for communal roost or feeding hotspots (Cortés-Avizanda et al., 2014; Kane et al., 2015).

Vultures from the Gyps genus (which includes seven species) are among the largest flying birds, showing high sociality and covering large areas in search of ephemeral and unpredictable resources such as carrion (e.g., 162,824 km² annually on average in the case of Cape Vultures, Gyps coprotheres; Jobson et al., 2021). The Eurasian Griffon Vulture Gyps fulvus is a monomorphic social species that breeds colonially (Almaraz et al., 2022; Donázar, 1993; Harel et al., 2017; Zuberogoitia et al., 2018). The breeding period of the species expands from early December (when first copulates occurs) to late August, when fledglings fly from the nest (Donázar, 1993; Zuberogoitia et al., 2018). Individuals forage over vast areas to satisfy their energetic requirements (e.g., ranging annually from 1560 to 4233 km², Fluhr et al., 2021; Monsarrat et al., 2013; Nathan et al., 2012; Xirouchakis et al., 2021), frequently congregating around both wild and domestic ungulate carcasses (Cortés-Avizanda et al., 2010, 2012 but see also Delgado-González et al., 2022). Although information exists on Griffon vulture movement ranges (Arkumarev et al., 2021; Arrondo et al., 2018; Arrondo, Sanz-Aguilar, et al., 2020; Fluhr et al., 2021; García-Ripollés et al., 2011; Harel et al., 2017; Spiegel et al., 2013, 2015; Xirouchakis et al., 2021; Xirouchakis & Mylonas, 2007; Zuberogoitia et al., 2013), virtually nothing is known about the spatio-temporal variation in the movement patterns of adult individuals, or about the factors (e.g., sex, breeding region) governing the spatial ecology and home-range fidelity of this species from a mechanistic perspective.

In this paper, we gather movement data from 127 GPS-tagged adult Griffon Vultures captured in five breeding regions of peninsular Spain, the largest vulture population in Western Palearctic encompassing up to 37,000 breeding pairs (90% of all European populations) (Del Moral & Molina, 2018). Our main aim is to assess the effect of individual and environmental factors on movements and spatial use indicators. Specifically, our objectives are: (1) to estimate annual and monthly home-range sizes, monthly cumulative distances traveled, and monthly home-range site fidelity; and (2) to investigate the effect of season, sex and breeding regions on individual monthly home-range size, site fidelity, and cumulative distance. We hypothesize that adult Griffon Vultures, being a large monomorphic colonial species, will exhibit large home-ranges and will travel long distances to fulfill their requirements (mainly food), especially during the autumn and winter, when food availability is the lowest (Margalida et al., 2018; Spiegel et al., 2013). We also predict that the fidelity of monthly home-ranges should be similar between sexes due to the lack of dimorphism, but would differ between seasons, as foraging constraints are more likely during the breeding period (see Carrete & Donázar, 2005). Finally, we expect differences between

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breeding regions due to differences in resource availability (Morant et al., 2022).

2 | METHODS

2.1 | Capture and tagging of vultures

From 2014 to 2022, we captured 127 adult Griffon Vultures (43 males and 84 females) in five breeding regions distributed across northern (Alto Ebro: 49 individuals, Pyrenees: 21 individuals), central (Segovia: 15 individuals), and southern Spain (Cádiz: 12 individuals and Cazorla: 30 individuals) (see Figure 1). Breeding regions were delimited according to the proximity between nesting sites and the biogeographic characteristics of each area where nests are located. Birds were trapped using remotely activated cannon nets and cage traps baited with livestock carcasses. Individuals were tagged with yellow or blue plastic alphanumeric and metal rings and equipped with solar-powered GPS/GSM transmitters (Ecotone https://ecoto ne-telemetry.com/en, Ornitela https://www.ornitela.com/, and e-Obs https://e-obs.de/). Devices were attached using a Teflon tape backpack harness. The total weight of the transmitters and rings did not exceed 64 g, which represented less than 3% of the body weight of the individuals (Bodey et al., 2018). The age of individuals was estimated from plumage molt and other external features such as the color of the culmen and the eyes (Donázar, 1993; Zuberogoitia et al., 2013), while sex was determined using molecular sexing techniques from body feather samples (Fridolfsson & Ellegren, 1999).

Tracking devices were programmed to record fixes (i.e., GPS positions) at 5–10-min intervals from 1 h before sunrise to 1 h after sunset (see Table 1 for details of the tracking devices and sampling frequency). GPS data were incorporated into the Movebank online data repository (www.movebank.org). Data were standardized by resampling the GPS fixes to 15 min for each individual to homogenize our dataset. Vultures were tracked on average 1040±809 days with a mean number of fixes per individual of 41,335±40,493.

2.2 | Estimation of home-range, cumulative distance traveled, and site fidelity

We estimated annual and monthly home-ranges using the 95% kernel density estimator contours (KDE) and the cumulative distance traveled for each tagged individual using the "amt" package (Signer et al., 2019). We selected KDE function instead of AKDE since this had no qualitative improvement to our results and notably increased computational time compared to KDE function (see for instance Silva et al., 2022). Cumulative distance traveled per month was measured as the total length of each track (in km) (Edelhoff et al., 2016). Individuals with less than ten fixes per day were discarded (n = 4 cases).

We measured the individual home-range fidelity by calculating the percentage of overlap between consecutive monthly homerange (Fieberg & Kochanny, 2005). We selected the 95% KDE as an

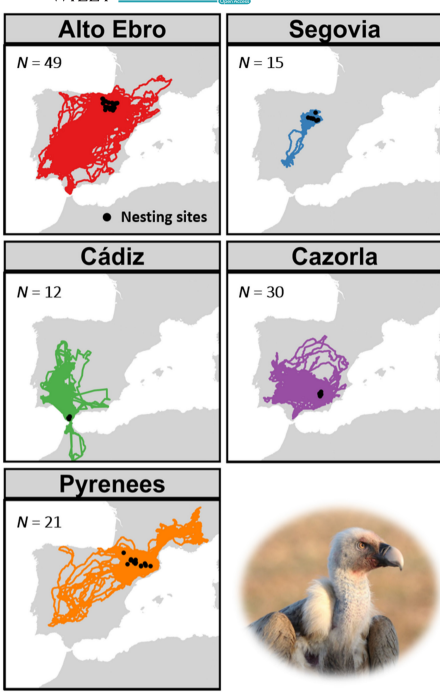


FIGURE 1 Movements and nest
locations of 127 GPS-tagged adult Griffon
Vultures in northern (Alto Ebro and
Pyrenees), central (Segovia), and southern
(Cádiz and Cazorla) Spain. N represents
the number of tagged vultures for each
breeding region. Photo credit: Manuel de
la Riva.

	Home-range (km ²)	Cumulative distance traveled (km)		
Annual	5027±2123 (1981-9863)	15,090±11,256 (219-39,298)		
Monthly	4889±1753 (1908-6822)	1776±1497 (108-7172)		

TABLE 1 Annual and monthly homerange size (in km²; estimated at 95% KDE), and cumulative distance traveled (in km) of 127 adult Griffon vultures tagged with GPS in Spain.

Note: All values area shown with mean (\pm SD) and their range (minimum and maximum values) in brackets.

estimator of home-range to evaluate differences in foraging areas. The percentage of overlap was quantified using the Bhattacharyya coefficient, which ranges between 0 (complete segregation) and 1 (perfect overlapping; Fieberg & Kochanny, 2005; Winner et al., 2018).

2.3 | Data analysis

We analyzed the effect of sex, season, and breeding regions on the monthly home-range sizes, cumulative monthly distances traveled (normal error distribution, identity link function), and percentage

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of monthly overlap (beta error distribution, identity link function) using generalized linear mixed models (GLMMs; glmmTMB package, Brooks et al., 2017). Sex, season, and breeding region were included as fixed factors, while individual identity and year were included as random terms in the models to avoid pseudo replication.

We did not divide the period into breeding stages since we could not rule that all individuals were breeding each monitoring year and, therefore, followed the same breeding cycle periods. Instead, we here referred to the season as the division of the year based on changes in weather, ecology, and the number of daylight hours in a given region. In the case of the breeding region, we regard this variable, not only as an indicator of differences in the periods suitable for soaring (mediated by photoperiod; see Scacco et al., 2019) but also as a potential proxy of differences in the food available in the field (e.g., some regions encompass much more carrion than others in absolute numbers; Morant et al., 2022) and population differences in the levels of exposure to known threats (i.e., risk of collision, electrocution or poisoning; Arrondo, Sanz-Aguilar, et al., 2020).

Finally, we did not consider the interaction between the aforementioned variables due to (1) the complexity of the understanding that is implicit to them (e.g., sex differences in movement associated with particular regional conditions) and (2) to avoid speculations without any empirical foundations in the interpretation of the results.

Models were compared using the Akaike Information Criterion corrected for small sample sizes (AICc; Burnham & Anderson, 2002). The best model was the one with the lowest AICc value. All models with a difference of Δ AICc < 2 were considered alternatives (Burnham & Anderson, 2002). For the best model, homogeneity of variance and normality of residuals was inspected by using the "ggresid" package to check the goodness-of-fit of our best models (Goode & Rey, 2019). We estimated the variance explained by

TABLE 2 Models obtained to assess the effects of individual sex, season, and population on the movement and spatial segregation patterns of 127 GPS-tagged adult Griffon Vultures in Spain.

	Model	k	AIC _c	∆AIC _c	AIC _{cw}	R ² fixed	R ² randor
Movement patterns	;						
Home-range size (km²)	Season + sex + breeding region	6	59582.1	0.00	0.993	28.18	20.61
	Breeding region + season	5	59612.7	30.56	0.007		
	Breeding region + sex	5	59849.0	266.88	0.000		
	Season + sex	5	59862.2	280.05	0.000		
	Season	4	59877.6	295.45	0.000		
	Breeding region	4	59877.7	295.58	0.000		
	Sex	4	60132.9	550.79	0.000		
	Null	3	60147.5	565.39	0.000		
Cumulative	Season + sex + breeding region	6	67835.8	0.00	0.988	33.29	19.25
distance (km)	Season+breeding region	5	67844.6	8.74	0.012		
	Season + sex	5	67941.3	105.48	0.000		
	Season	4	67952.4	116.59	0.000		
	Breeding region + sex	5	69656.5	1820.6	0.000		
	Breeding region	4	69665.9	1830.	0.000		
	Sex	4	69771.4	1935.6	0.000		
	Null	3	69784.4	1948.5	0.000		
patial segregation							
Monthly	Season + sex + breeding region	6	-1415.9	0.00	0.998	0.842	1.974
home-range	Season + breeding region	5	-1392.5	23.45	0.002		
fidelity (%)	Season + sex	5	-1371.2	44.73	0.000		
	Season	4	-1353.0	62.90	0.000		
	Breeding region + sex	5	-1230.5	185.46	0.000		
	Breeding region	4	-1209.2	206.70	0.000		
	Sex	4	-1189.4	226.54	0.000		
	Null	3	-1171.3	244.57	0.000		

Note: The null model was included in our set of models. For each best model, variability (as a percentage) explained by the fixed and random predictors (R^2) are shown. The best models ($\Delta AICc < 2$) are highlighted in bold.

Abbreviations: AICc: Akaike Information Criterion corrected for small sample sizes; k: number of parameters; w: Akaike weight; Δ AICc: difference between the AICc of model i and that of the best model (i.e., the model with the lowest AICc).

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the fixed and random factors by using the "performance" package (Lüdecke et al., 2021), while differences between their levels were assessed through Tukey's post hoc tests using the "emmeans" package (Lenth, 2022). All tests were two-tailed, statistical significance was set at $\alpha = 0.05$, and all results were shown as mean \pm standard deviation. Results of the Tukey's post hoc tests included the marginal means and its standard error. Spatial and statistical analyses were done in R version 4.0.3 (R Core Team, 2021).

3 | RESULTS

Griffon Vultures exploited very large annual home-range sizes (ca. 5000 km^2), showing average monthly cumulative distances traveled of 1776 km (Table 1), and a monthly home-range fidelity of $67.77 \pm 25.05\%$. Monthly home-ranges differed between sexes, seasons, and breeding regions (Tables 2 and 3; Figure 2). Males showed a smaller monthly home-range size than females (Table 3). Birds also

showed larger home-ranges during spring and summer compared to autumn and winter (Table S2; Figure 2). Individuals from southern breeding regions (Cazorla, Cádiz), and central Pyrenees showed, on average, larger home-ranges than those of central (Segovia) and northern breeding regions (Alto Ebro) (Table S2 and Figure 2).

Monthly cumulative distances traveled were similar for both females and males (Tables 2 and 3; Figure 3). However, birds traveled longer distances during spring and summer than during autumn and winter (Table S2). Individuals from Alto Ebro, Segovia, and Pyrenees traveled larger monthly distances than those of Cazorla and Cádiz (Table S2; Figure 3).

The monthly home-range fidelity was slightly higher for males than females, and during spring and summer, being the lowest in autumn and winter. Likewise, there were differences in fidelity between breeding regions. Individuals from Central Spain (Segovia) showed higher fidelity over time than those from the north (Alto Ebro, Pyrenees) and southern Spain (Cádiz, Cazorla) (Tables 2 and 3 and Table S2; Figure 4).

TABLE 3 Estimates for fixed terms of the best models of monthly home-range size, cumulative distance traveled per month and monthly home-range fidelity for 127 GPS-tagged adult Griffon Vultures in Spain.

Response variable	Predictors	$Estimate \pm SE$	z value	p-value
Monthly home-range size	Season (spring)	2250.6 ± 193	11.660	<.001
	Season (summer)	2195.7 ± 195.6	11.225	<.001
	Season (autumn)	204 ± 193.9	1.053	.292
	Sex (male)	-1294.5 ± 291	-4.449	<.001
	Breeding region (Segovia)	-899.8 ± 586.8	-1.533	.125
	Breeding region (Cádiz)	1670.7 ± 507.2	3.294	.001
	Breeding region (Cazorla)	5750 ± 319.8	17.980	<.001
	Breeding region (Pyrenees)	1920.3 ± 397.9	4.827	<.001
Cumulative distance	Season (spring)	1393.69 ± 47.05	29.622	<.001
	Season (summer)	1925.70 ± 48.43	39.763	<.001
	Season (autumn)	162.46 ± 49.79	3.263	.001
	Sex (male)	41.15 ± 73.88	0.557	.577
	Breeding region (Segovia)	169.34 ± 174.01	0.973	.331
	Breeding region (Cádiz)	950.40 ± 139.34	6.821	<.001
	Breeding region (Cazorla)	419.20 ± 84.79	4.944	<.001
	Breeding region (Pyrenees)	649.23 ± 108.33	5.993	<.001
Monthly home-range fidelity	Season (spring)	0.439 ± 0.046	9.502	<.001
	Season (summer)	0.629 ± 0.046	13.549	<.001
	Season (autumn)	0.315 ± 0.047	6.653	<.001
	Sex (male)	0.267 ± 0.058	4.543	<.001
	Breeding region (Segovia)	0.691 ± 0.147	4.688	<.001
	Breeding region (Cádiz)	-0.099 ± 0.135	-0.729	.465
	Breeding region (Cazorla)	0.253 ± 0.096	2.635	.008
	Breeding region (Pyrenees)	0.255 ± 0.112	2.279	.022

Note: Season, Sex, and Breeding region variables were coded as a factor, being "Winter," "Female," and "Alto Ebro" the reference values for statistical comparisons. Significant values are highlighted in bold. Abbreviation: SE, Standard error. 20457758, 2023, 2, Downloaded from https //onlinelibrary.wiley.com/doi/10.1002/ece3.9817 by Juan Pére Readcube (Labtiva Inc.) , Wiley Online Library on [30/01 1/2024] See the Wiley Online Library for ;OA article are governed by the applicable Creative Comi

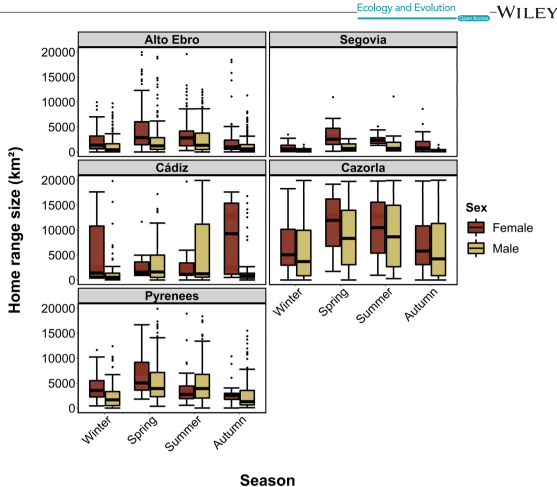


FIGURE 2 Mean monthly home-range size (in km², based on 95% KDE_± SD) of 127 adult Griffon Vultures (43 females and 84 males) tagged in northern (Alto Ebro and Pyrenees), central (Segovia), and southern (Cádiz and Cazorla) Spain. The standard deviation is shown as error bars.

4 | DISCUSSION

Griffon Vulture movements varied between seasons, breeding regions and sexes. We found that movements were larger during spring and summer, which is similar to other soaring raptors such as Bearded Vulture or Bonelli's Eagle (see Margalida et al., 2016; Pérez-García et al., 2013). This could be related to the food requirements associated with reproduction, which may force vultures to prospect larger areas, seeking for dispersed and unpredictable carcasses to satisfy the food requirements of the offspring and themselves (Carrete & Donázar, 2005). However, seasonal differences could be also explained by longer days (and therefore more time to forage) and better atmospheric conditions particularly during summer, minimizing energy expenditure during long-range movements (see Martin-Díaz et al., 2020). Similarly, differences in monthly home-range size and cumulative distance traveled between regions could be explained by differences in both the importance of predictable and nonpredictable food availability and the ability to move due to better flight conditions due to the presence of thermal uplifts (Scacco et al., 2021). For instance, the southern populations may experience a higher thermal uplift availability (mainly due to warmer climatic conditions in summer), thus minimizing the energy

expenditure while increasing the movement capacity of birds (e.g., see Scacco et al., 2019).

Differences in foraging performance between sexes are common in mammals and birds and are often due to differences in body size and parental duties (Lewis et al., 2002), In monomorphic species such as the Griffon Vulture, these differences might be associated with energetic and nutritional requirements for reproduction (Bennison et al., 2022; Pinet et al., 2012). Our results showed that females have larger home-ranges and travel farther than males. These findings align with other studies in which the same dataset was analyzed and where females exhibited larger traveled distances than males during reproduction (see Delgado-González et al., 2022; Gangoso et al., 2021), including other vulture species (see Bamford et al., 2007; García-Jiménez et al., 2018; Kane et al., 2015; Krüger et al., 2014; Margalida et al., 2016).

Contrary to our expectations, we also found sex differences in the monthly home-ranges fidelity. Males showed greater fidelity than females, indicating that the latter use different areas throughout the year. This gender variation in seasonal fidelity may respond to differences in resource selectivity (Delgado-González et al., 2022; Hertel et al., 2020). In fact, according to Fernández-Gómez et al. (2022) males may be more prone to feed

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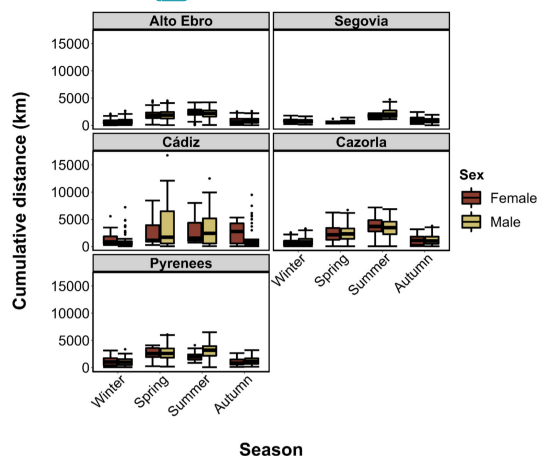


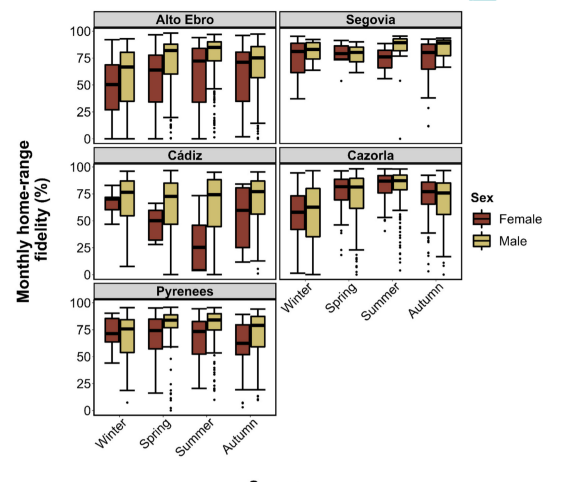
FIGURE 3 Mean monthly cumulative distance traveled (in km) of 127 adult Griffon Vultures (43 females and 84 males) tagged in northern (Alto Ebro and Pyrenees), central (Segovia), and southern (Cádiz and Cazorla) Spain. The standard deviation is shown as error bars.

on predictable resources such as supplementary feeding stations or vulture restaurants, while females may rely on more ephemeral and less clumped food resources. Thus, there may be parallel strategies in the large-scale exploitation of space and, therefore, sexual spatial segregation (see also Perrig et al., 2021). Alternatively, the fact that males show a greater fidelity of their monthly homeranges throughout the year and that it is higher during the springsummer period could be related to their greater territoriality. Males may be more involved in nest building and guarding (Xirouchakis & Mylonas, 2007), which might be a response to competence for nesting sites (see Zuberogoitia et al., 2018). Interestingly, females not only showed higher extension of home-ranges than males but also exhibited lower site fidelity. All this reinforces the argument that females might forage more in different locations throughout the year, therefore, showing lesser home-range fidelity over time than males. These sex and seasonal differences in individual home-range fidelity were similar between breeding regions except in the case of Cádiz and Segovia, where females exhibited lower fidelity during spring/summer.

Differences (or lack of them) between breeding regions in individual movement patterns indicate that local effects not explored in this work may be affecting them. This is evidenced by the low variance explained by the fixed factors of the best models and the high variance explained by random factors (e.g., home-range size fidelity models). The latter indicates that high interindividual differences in home-range area and fidelity exist. Moreover, there are variables, such as distance to trophic resources, colony size, or habitat type, that perhaps could improve the results from our models and better explain breeding region level differences in the evaluated parameters (see Cecere et al., 2018; Delgado-González et al., 2022; Harel et al., 2017). Finally, it should be taken into account that the effect of differences in the sampling duration of individuals of some breeding regions also affects the observed home-range sizes which may have led to an underestimation of this and other parameters such as cumulative distance or home-range fidelity.

5 | CONCLUDING REMARKS

Our work adds evidence to the spatial requirements of large soaring birds such as vultures, and the importance of individual and regional differences in explaining their movement patterns. Contrary to our expectations, we found sex-dependent spatial segregation in this monomorphic species, maybe related to behavioral differences between males and females, particularly during the breeding period. Moreover, we observed that vultures showed larger home-ranges



Season

FIGURE 4 Differences in the monthly home-range fidelity (in %) at an individual level between females and males in different seasons of 127 adult Griffon Vultures (43 females and 84 males) tagged in northern (Alto Ebro and Pyrenees), central (Segovia), and southern (Cádiz and Cazorla) Spain. The standard deviation is shown as error bars.

and cumulative distance traveled during the breeding period, when site-fidelity is higher, with females traveling further. Finally, despite high levels of variation in home-range sizes between breeding regions, there were no clear differences in site fidelity between breeding regions over time, which may indicate stability in the homerange. Our findings open new promising avenues for research on interindividual differences in optimal foraging, and on the intrinsic and extrinsic factors operating at multiple levels (Williams & Safi, 2021). Beyond this, increasing our knowledge of how these patterns translate into regional differences (or lack thereof) is crucial in predicting population dynamics through movement ecology (Shaw, 2020).

Importantly, our results add information that is crucial may be decisive for the effective management and conservation of highly mobile species which require protective measures to be implemented at large spatial scales. For example, differences in the home-range sizes between females and males may indicate different levels of exposure to well-known threats to species (e.g., poisoning). This in fact could have direct effects on population dynamics in the long term, in particular, in those areas where overall individual survival is low (see Arrondo, Sanz-Aguilar, et al., 2020). Additionally, the observed patterns of interpopulation and seasonal differences in movement ranges highlight the need to disentangle the risk factors for affecting different populations. In particular, in those places where installation of wind energy facilities is expected in the near future that may pose serious threat for the species (Pérez-García et al., 2022; Serrano et al., 2020). Finally, the large movement patterns of this species highlight the need to implement transboundary conservation plans (see Lambertucci et al., 2014) and closely evaluate the sensitivity to different risk factors operating at each region to design a coordinated response of all countries involved in species conservation.

AUTHOR CONTRIBUTIONS

Jon Morant: Conceptualization (equal); data curation (lead); formal analysis (lead); investigation (equal); methodology (lead). Eneko Arrondo: Supervision (supporting); validation (supporting); writing – original draft (supporting); writing – review and editing (supporting). José Antonio Sánchez-Zapata: Supervision (supporting); validation (supporting); writing – original draft (supporting); writing – review and editing (supporting). José Antonio Donázar:

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Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Ainara Cortés-Avizanda: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Manuel De La Riva: Data curation (supporting); supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Guillermo Blanco: Supervision (supporting); validation (supporting); writing original draft (supporting); writing - review and editing (supporting). Félix Martínez: Supervision (supporting); validation (supporting). Juan Oltra: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Martina Carrete: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Antoni Margalida: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Pilar Oliva-Vidal: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing review and editing (supporting). Jose María Martínez: Supervision (supporting); validation (supporting). David Serrano: Supervision (supporting); validation (supporting); writing - original draft (supporting); writing - review and editing (supporting). Juan Manuel Pérez-García: Conceptualization (equal); data curation (equal); formal analysis (equal); investigation (equal); methodology (equal); project administration (equal); supervision (equal); validation (equal); writing - original draft (equal); writing - review and editing (equal).

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CONFLICT OF INTEREST STATEMENT

The authors declare no competing interests.

DATA AVAILABILITY STATEMENT

The data and scripts used in the analyses are publicly available at Zenodo data repository (https://doi.org/10.5281/zenodo.7500857).

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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