

The use of social information in vulture flight decisions

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18 Abstract

19 Animals rely on a balance of personal and social information to decide when and where to
 20 move next in order to access a desired resource, such as food. The benefits from cueing on
 21 conspecifics to reduce uncertainty about resources availability can be rapidly overcome by the
 22 risks of within-group competition, often exacerbated toward low-ranked individuals. Being
 23 obligate soarers, relying on thermal updrafts to search for carcasses around which competition
 24 can be fierce, vultures represent ideal models to investigate the balance between personal and
 25 social information during foraging movements. Linking dominance hierarchy, social affinities
 26 and meteorological conditions to movement decisions of eight captive vultures, *Gyps spp.*,
 27 released for free flights in natural-like soaring conditions, we found that they relied on social
 28 information (i.e. other vultures using/having used the thermals) to find the next thermal updraft,
 29 especially in unfavourable flight conditions. Low-ranked individuals were more likely to
 30 disregard social cues when deciding where to go next, possibly to minimise the competitive
 31 risk of social aggregation. These results exemplify the architecture of decision-making during
 32 flight in social birds. It suggests that the environmental context, the context of risk and the
 33 social system as a whole calibrate the balance between personal and social information use.

34
 35 **Key-words:** griffon vulture, hierarchy, movement decision, landscape exploration, social information,
 36 unpredictable resource

1. Introduction

Animals must constantly decide where and when to move next in order to find resources such as food, water, shelter, or a mate, necessary for life. To make these decisions, they can rely on two sources of information: personal information and social information. Personal information includes knowledge of the spatiotemporal patterns of resource distribution that individuals may perceive or have memorised from previous encounters [1]. For example, food-storing birds are able to return to locations where they stored or saw food in the past, based on prior expectation of the resource availability [2]. Social information, on the other hand, is obtained by observing the behaviour of others [3–5]. Feeding, fleeing, or mating individuals provide discrete information about the availability and locations of food, predators, or potential mates.

For resources that are heterogeneously distributed in the environment, ephemeral and unpredictable, using only personal information for movement decisions may be prone to inaccuracies [6]. In such conditions, social animals may benefit from companions' knowledge and may follow the dominant or oldest individual(s) considered as knowledgeable (e.g. homing pigeons, *Columba livia*, or elephants, *Loxodonta africana*, [7,8]), follow the largest group through shared decision-making [9], or stay with preferred affiliates [10–12]. Because using social information can considerably reduce uncertainty in finding resources, individuals should favour this source of information to achieve cost-efficient movement [13–15]. However, relying heavily on social information can also lead individuals to aggregate on resources, potentially inducing competition by exploitation or interference if the resource is monopolizable and depletable [16]. Since both social and personal information are often available to social animals [1], they need to balance their relative importance, depending on the availability and predictability of the resource. When deciding on the next movement step, social animals must trade-off the decreased uncertainty of locating a resource through social information, with the potential increase in competition risk. Such a balance may be dictated by the immediate needs of the individual and its risk sensitivity [17] but also by the group social organisation. For example, low-ranked individuals are known to suffer more from within-group competition compared to high-ranked individuals [18] and should therefore be more reluctant to engage into social information use, which could eventually trigger proximity to despotic individuals [19].

Vultures rely on two unpredictable resources: carcasses to feed and thermal updrafts to move. During foraging flights, these large soaring birds gain altitude by circling into thermal

updrafts (i.e. masses of hot air rising from heated surfaces) and glide across the landscape to the next updraft while scanning the ground for carcasses [20]. Although some topographic features are clearly favourable to updrafts presence [21], at the individual level, challenging local meteorological conditions (e.g. high wind speed, low temperature, high cloudiness) can make thermal locations and availability hard to predict [22]. If they fail to detect an updraft, vultures may be forced to switch to flapping flight, or worse to land and take-off again, significantly increasing their energy expenditure [23,24]. While both thermals and carcasses are relatively unpredictable, thermals are not depletable contrary to carcasses. When a vulture discovers a carcass, its sharp drop in altitude while circling before landing is used as a signal by conspecifics, dragging tens of individuals to the food source in a few minutes [25,26]. As the number of vultures around the carcass increases (up to 100-120 individuals, [27,28]), individual feeding rates decrease due to reduced access to the resource, resource depletion by competitors and increased agonistic interactions [27]. Therefore, in these social birds, individuals should balance the advantage of conspecific presence to locate thermal updrafts [29] with the ultimate cost of competition around the carcasses that can be fierce [30–33]. As such, vultures are ideal models to investigate the role of conspecifics in shaping their foraging movement decisions.

Using a group of captive but freely-flying ‘griffon’ vultures, *Gyps fulvus* and *G. rueppellii*, tagged with high-resolution GPS loggers, we studied how conspecifics’ presence shapes individuals’ movement decisions during soaring flights. Despite being trained birds released for public shows, these individuals sometimes detected and fed on carcasses at surrounding farms (BN and YS, pers. obs.). We therefore consider these flights comparable to natural foraging flights. Focusing on the movement steps from thermal to thermal, we first assessed when do individuals preferentially discover new thermals (i.e. use of personal information) compared to using thermals already discovered by conspecifics (i.e. use of social information). We expected that vultures would favour the use of social information when unfavourable meteorological conditions increased thermal unpredictability and when flight conditions (e.g. low altitude) increased risks of landing [1]. Furthermore, given the hierarchy in vulture groups, we expected low-ranked individuals to be more prone to use personal information than high-ranked individuals to try to find the food source first, in order to avoid large aggregation [34,35]. Second, we investigated the drivers underlying thermal selection when individuals had to choose between simultaneously available thermals. We expected individuals to select thermals providing the maximal positive vertical speed (i.e. climb rate) as it may provide a reliable proxy of the thermal current strength helping them maximise their

height gain [29]. To decrease uncertainty about resource finding and risks mentioned above, we expect that individuals should favour thermals hosting the maximum number of individuals to maintain cohesion and secure the possibility to cue on as many conspecifics as possible [29]. Finally, social preferences may also influence decision, with individuals preferentially moving together with preferred affiliates [12,36,37], as it could reduce competition due to familiarity between individuals [38].

2. Materials and methods

2.1 Study site, vultures housing conditions and experimental settings

The study was carried out in 2021 and 2022 at the Rocher des Aigles falconry centre, Rocamadour, France, and divided between winter and summer periods each year. During winters, vultures were housed within an aviary (6.7 x 6 x 6 m) equipped with four perches: three of them measuring 3.10 m, placed at 1.7, 2.6 and 3.5 m from the ground, and one of the full width of the aviary at 4 m height. This setting was used to estimate vulture social bonds (see Social bond estimation). In addition, besides being fed daily on small pieces of meat to prevent conflicts, five feeding events (one each week during a five-week period) were organised in the aviary on a butchery carcass occurring after a one-day fasting (to motivate feeding). These feeding events were used to assess dominance hierarchy within the group (see Hierarchy estimation). In summer, these trained vultures were kept perching on individual logs, released several times per day to execute free flight shows for the public within a landscape composed of plateaus interspaced by canyons, similar to “Causses” landscape typically used by french wild vultures [39]. The falconry centre is located near a 120 m-deep canyon and offers natural soaring conditions for raptors, making this study site a great place to investigate natural group flight behaviour (see Group flights), [24].

We used GPS data and visual observations to characterise the social and flight behaviour of eight captive vultures (7 Eurasian griffon vultures, *Gyps fulvus*, and 1 closely-related Rüppell’s vulture, *Gyps rueppellii*), including five females and three males (Table S1). Each year, we conducted experiments on a group of six individuals (two griffon vultures were replaced in 2022, Table S1). Experiments followed the animal ethic guidelines of France and the Centre National de la Recherche Scientifique. Handling of birds to fit GPS loggers followed the protocol of telemetry study of vultures authorised in the Programme Personnel 961, coordinated by OD, under the supervision of the French ringing centre, CRBPO, Paris.

Furthermore, experiments, observations, handling and flight events were systematically performed under the guidance of the head of animal caretakers, BN.

2.1.1 Social bond estimation

During five weeks in both years (December/January 2020-2021 and November/December 2021), we recorded pictures of vultures in the aviary from 8:00 to 19:00 (local time) at 5 minutes interval, using three camera traps (Wosport Big Eye D3 and Reconyx HyperFire HC600).

We identified birds using repeated colours on plastic rings and marks on the ruff and backhead feathers, using harmless colour sticks (Raiderx GmbH, Figure 1A). We then processed recorded pictures to extract the individuals' ID and position (bill or head position), using a purpose-built image annotation program in Julia software, JuliaHub Inc., [39]. For subsequent analyses, we relied on R software (v 4.2.2, R software, 2022, [40]).

We considered the social bond between a dyad of individuals i and j based on spatial proximity following the Simple Ratio association Index (SRI, equation 1, [41,42])

$$\text{Equation 1: } SRI_{i,j} = SRI_{j,i} = \frac{n_{tog}}{n_{tot}},$$

where n_{tog} is the number of pictures in which individuals i and j were on the same perch at a Euclidean distance of less than 1.55 m and n_{tot} is the total number of pictures in which individuals i and j were both detected on the same perch. SRI values varied between 0 and 1, where 0 represented dyads that were never seen associated and 1 represented dyads that were always observed sitting at less than 1.55 m from each other. The distance of 1.55 m was chosen as matching to the mode of the inter-individual distances distribution (Figure S1). This was also consistent with the aviary setting, as it corresponded to half the length of most available perches. Our analyses were robust to other choices for this distance threshold (see Supplementary Material, ESM01).

2.1.2 Hierarchy estimation

Each winter, we estimated hierarchy within the vulture group by monitoring feeding interactions during the five carcass-based feeding events in the aviary (10 in total, Figure 1B) using a remotely-controlled video camera (GoPro Hero 4, GoPro Inc.) fixed at 2 m height on the aviary wall. These feeding events lasted on average 34 min (SD $\pm$ 4 min).

We computed individuals' rank relying on the randomised Elo-rating approach [43,44], which accounts for potential temporal instability of the rank using permutations in the agonistic interaction series ('elo_scores' function, *aniDom* package, [44,45]; using 1000 randomisations and fixing the rank adjustment speed along the series, K-factor, to 200). The interaction series consisted in identifying the "wins" and "losses" for a given individual in agonistic interactions [46] with other individuals in each video (annotated with BORIS video analysis software, [47]). We used the ethograms from Bose & Sarrazin (2007, [30]) and Valverde (1959, [48]) to characterise griffon vulture feeding behaviour and between-individual interactions. An individual won the interaction when it interrupted another individual's feeding bout (by pecking it, displacing it or engaging in a fight), and finally accessed the carcass before its opponent. In other cases, the interaction was considered as a "loss" for the initiator. We assessed the reliability of the dominance hierarchy through individual Elo-rating repeatability ('estimate_uncertainty_by_repeatability' function, *aniDom* package, [44]).

2.1.3 Group flights

We recorded vulture flights decisions during 42 flight sessions (21 sessions each year) in the vicinity of the Rocher des Aigles. In general, birds were released for a flight session three times per day (in rare occasions from 2 to 4 times), at around 11:00, 14:30 and 16:00 (local time) for a mean duration of 26.03 min (SD $\pm$ 14.15 min) of flight. These captive vultures are trained to fly freely, searching for thermals, gaining altitude and coming back to their trainers (Supplementary Video 1). Vultures were equipped with a high-resolution GPS logger (4 Hz, TechnoSmart, models Gipsy 1, Gipsy 5 or Axytreck) positioned at their lower back using a Teflon leg-loop harness (Figure 1C, [49]). They were released in two groups, built according to social preferences with the three most socially-bonded birds together, at 2-min intervals. Release order alternated between consecutive days. For each flight session, we recorded and considered as stable the cloudiness (i.e. the proportion of clouds covering the sky, on a scale from 0 - no clouds - to 8 - sky fully covered by clouds), horizontal wind speed and temperature (all extracted from meteofrance.com based on local meteorological models).

To further investigate how vulture thermal choices were shaped by personal and social information, we pre-processed flight tracks in three consecutive steps. We subsampled individuals' tracks from 4 to 1 GPS fix per second, and segmented their flight behaviour into gliding, linear soaring and circular soaring. We then created spatio-temporally dynamic maps of thermal availability based on the spatial clustering of individual's circular soaring phases. Leaning on these maps, we retraced the history of thermal use/choice by individuals.

2.1.3.1 Thermal use identification

To segment vulture flight between circular soaring, linear soaring and gliding flight we first calculated turning angle and vertical speed between consecutive locations using the *move* R package [50]. We applied a moving window of 30 s to calculate the absolute cumulative sum of the turning angles (hereafter cumulative turning angle) and a moving window of 5 s to calculate the average vertical speed. We then applied a k-means approach ($k = 2$, ‘kmeans’ function, *stats* R package) on the smoothed vertical speed (positive speed when flying upwards) to distinguish between soaring (ascending flight) and gliding (descending flight, [51,52]). We further classified soaring locations into circular soaring (indicating use of thermal updrafts) and linear soaring (also called slope soaring, expected to occur outside of thermals), with circular soaring being associated with a cumulative turning angle ≥ 300 degrees. A result of segmentation is illustrated in Figure 2B. Finally, we inferred the use of a thermal when the individual engaged in circular soaring for more than 30 s, with no interruption of more than 5 s of gliding.

2.1.3.2 Dynamic mapping of available thermals

Within each flight session, we created a dynamic map of thermals (Figure 3). First, we spatially clustered vulture circular soaring locations (reflecting the use of the same thermal updraft) independently of time by using a 3D density-based spatial clustering approach (‘dbscan’ function, *dbscan* R package, [53]). This algorithm relies on a spherical neighbourhood to perform density-based neighbour joining, i.e. clustering (Figure 2C). We assumed this neighbourhood to be of a 40-m radius, and a minimum number of five locations within this range for the algorithm to consider the neighbourhood further. This 40 m threshold corresponded to the largest 4-nearest-neighbour distance observed when considering locations attributed to thermal use only (‘kNNdistplot’ function, *dbscan* R package) and matched with empirical expectations of radius during circular soaring phases [54].

We then made those maps dynamic in time by considering the lifetime of each thermal. We considered a thermal as “available” from the moment when the first individual entered it until the last individual left it (Figure 3).

2.2 Statistical analyses

We defined collective flight events as any time of a flight session when at least two individuals were flying. For each of these events, we first analysed the use of social information (the tendency to join thermals already discovered by conspecifics) as a function of external

(meteorological) and internal drivers (individual traits). We then used step selection functions to define, at each movement step, which drivers determined the selection of the chosen next location (thermal updraft) relative to other potential locations.

2.2.1 Drivers of social information use

We investigated the effect of local meteorological context, individual traits and flight mechanics on the use of social information, defined here as the tendency to join thermals already discovered by conspecifics. We considered that an individual discovered a thermal when it was the first, among all individuals, to adopt circular soaring flight into it. For the analysis, we discarded the discovery of the first thermal in each flight session (as this thermal was necessarily discovered).

To investigate the drivers underlying the use of social information we modelled the probability to join a thermal already discovered by others using generalised linear mixed models (GLMMs) with binomial error structure and a logit link function [55]. Our full model contained the following ten fixed effects: **meteorological variables** with the (i) wind speed (categorical predictor), (ii) cloudiness and (iii) temperature (both continuous predictors); **social variables** with (iv) the age (continuous predictor) and (v) rank in the dominance hierarchy of the individual (ordinal categorical predictor), and variables related to the **mechanic of flight** with (vi) the glide-ratio (horizontal distance travelled during a 1-m altitude loss, only measured on glides with straightness > 0.95 in each flight), (vii) the altitude of and (viii) the 3D distance to the exit location from the previous thermal used (all continuous predictors). We also added (ix) the group in which individuals have been released (first or second group released for the flight) and (x) the time elapsed since the first individual take-off (continuous predictor) as control variables. Individual ID was considered as a random factor.

To compare the relative importance of the fixed effects we scaled all non-categorical variables to use their estimate as dimensionless effect size [56]. We examined the significance of each variable by comparing the goodness of fit of models with and without the variable of interest using a likelihood ratio test ('drop1' function, *stats* R package). Assumptions required for these statistical approaches (homoscedasticity, Gaussian distribution of residuals) were checked with plot diagnosis (histogram of residuals, residual Q-Q plot, distribution of residuals vs fitted values, *DHARMA* R package, [57]). We also tested for the presence of outliers, and calculated the variance inflation factor (VIF) to test for collinearity (VIF values ≥ 3 suggesting a strong collinearity [58]). We did not detect collinearity in our predictors (VIF_{max} = 1.74) (Figure S3). Furthermore, we extracted the marginal coefficient of determination (R_m^2) and the

conditional coefficient of determination (R_c^2) which describe, respectively, the proportion of variance explained by fixed effects and by the fixed and random effects combined [59]. Finally, as the flight time period, and the tested individuals differed, we cross-compared models fitting on the two years separately (see Supplementary Material ESM01).

2.2.2 Drivers of thermal updraft selection

To study the drivers underlying thermal selection, we embedded our work in the Step Selection framework [60]. We considered the series of thermals used by each individual. In that series, we focused on movement steps involving a flight to a thermal previously (or currently) used by a conspecific when other thermals were available. Using a conditional logistic regression, we compared the “chosen” thermal characteristics to those “available” but not chosen. The conditional logistic regression included seven predictors, respectively characterising the **thermal profitability** with (i) the distance to it and (ii) maximum vertical speed reached in the thermals by any individual since the focal individual has been released in the flight session (continuous predictors), **individual personal experience** considering whether (iii) the thermal was previously used by the focal individual (binary predictor), and **social information** with (iv) the presence of the focal individual’s preferred affiliates in the thermal or not (binary predictor), (v) the number of individuals present in the thermal, (vi) the weighted mean (by the number of previous visits to the thermal) of the social bond with individuals that used the thermals, and (vii) the negative cubed difference of ranks between the focal individual and those in the thermals (all continuous predictors, set to 0 for the two latter if no individuals used it/were present). We used the negative cubed difference to consider an attraction-repulsion effect (high rank toward low rank and the opposite respectively) by translating a linear rank difference to a relative hierarchy scale which enhances large rank differences. For example, following the curve of the negative cube function, if the difference of rank was five (e.g. the focal individual is ranked 6th - a low rank, a conspecific in another thermal is ranked 1st - a high rank) the probability that the focal individual joined the conspecific should be drastically decreased, mimicking a repulsion effect.

Also for this model, we scaled all non-categorical variables to better compare their relative importance. We fitted the conditional regression considering all individuals together, yet considering data stratified at the individual-step level. We finally reported the relative selection strength (RSS) of significant variables which provides the magnitude of estimated selection coefficients, holding all other covariates fixed [61,62].

3. Results

Vulture dominance hierarchy was steep (Figure S2) and reliably inferred (individual Elo-rating repeatability = 0.82 and 0.83 in 2021 and 2022 respectively). The rank orders among individuals present in both years were relatively consistent and unrelated to sex or age (Table S1). During the 21 flight sessions performed each year, we identified a total of 520 and 578 thermalling events in 2021 and 2022. On average, 63% (SD $\pm$ 7%, Table S1) of these circular soaring behaviours took place in thermals discovered by a conspecific.

3.1 Flight risks and hierarchy shapes the use of social information

Our model was significantly better than the null model (considering only control effects; AIC = 1237.4 and 1412.7 respectively) and explained 30% of the variance (Table S2). The probability for an individual to use a thermal previously discovered by a conspecific decreased with temperature (from 0.74 at 17°C to 0.43 at 31°C, Figure 4A, Figure 5A, Table S2), but tended to increase with cloudiness and wind speed (Figure 4A, Table S2). This probability dropped also with the distance from the previous thermal and the altitude at which the bird left it (from 0.63 when being at a distance of 12 m from the last thermal used to 0.16 at a distance of 6776 m and from 0.76 when exiting the last thermal at an altitude of 195 m to 0.039 at 1574 m of altitude, Figure 4A, Figure 5B, C, Table S2). Individuals lower in the dominance hierarchy were approximately twice as likely to discover new thermals than high-ranked individuals (Figure 4A, Figure 5D). We did not detect significant effects of age and glide-ratio on the probability to use thermal previously discovered by conspecifics (Figure 4A, Table S2). Fitting the same model structure on 2021 and 2022 data separately yielded the same overall results, suggesting that the observed pattern was robust to changes in hierarchy and between-year conditions (Figure S4, Table S2).

3.2 Vultures select thermal updrafts hosting the most conspecifics

We identified 178 movement steps where an individual entered a thermal while at least one other thermal was available simultaneously. Individuals were significantly more likely to select thermals hosting the largest number of conspecifics at time of decision (Relative Selection Strength [95% confidence interval] = 27.94 [5.99, 131.63]; Figure 4B, Table S3). On the contrary, the probability to choose a thermal tended to decrease when the preferred affiliate was using it. The distance to the previous thermal and the maximal vertical speed reached in the thermal and whether individuals used this thermal in the past did not significantly affect

thermal selection (Figure 4B, Table S3). At time of decision, the difference in dominance ranks as well as the presence of its preferred affiliate did not drive the individual's probability of selecting the thermal. This pattern was consistent when considering only movement steps where individuals had to choose between thermals used at time of decision (N = 61, Figure S5, Table S3). Furthermore, considering all decision events, the sensitivity analysis on the inter-individual distance threshold for the social bond strength estimation yielded the same results (i.e. 1.55 m, 1.30 m and 1 m; see Supplementary Material ESM01, Figure S6, Table S4).

4. Discussion

Using a combination of high-resolution tracking and social structure monitoring, we identified contextual drivers for the differential weighting of personal and social information in movement decisions. We showed that vultures' movement decisions predominantly relied on social information, especially in unfavourable flight conditions that increased thermal unpredictability or put individuals at risk of undesired landing. Overall, individuals preferentially joined thermals with the largest number of conspecifics. However, the use of social information depended on the individual social status: low-ranking individuals were more inclined to use personal information and discovered more thermals on their own than high-ranking individuals.

We found that low-ranked individuals, likely the ones suffering the most from interference competition, had higher probabilities of discovering new thermals, thus likely exploring their environment more intensively than the high-ranked individuals. Such flight strategy would enable subdominant individuals to reach carcasses first, or at least to arrive at the beginning of the feeding event when the rate of interference is lower [27] hence avoiding to loose opportunities due to conformity with conspecific behaviour [64]. From this may emerge a producer-scrouter dynamic [65,66] wherein the use of personal information from low-ranked individuals to arrive at food sources with lower competition levels would be exploited by dominant individuals to reduce their own searching effort [16,66,67]. This is coherent with previous observations of low-ranked vultures being “pioneers”: the very first individuals to land and feed on the carcasses before being displaced by high-ranked individuals arriving afterwards [27]. This influence of dominance on foraging tactics where low-ranked individuals explore and find food while dominant profit has also been observed in other social bird species such as common cranes *grus grus*, oystercatcher *Haematopus ostralegus*, house sparrows, *Passer domesticus*, and barnacle goose, *Branta leucopsis*, [18,67–69]. Eviction of

subordinates from food patches has even recently been identified as a trigger for collective movements [70]. In contrast, in activities where individuals do not experience competition, such as tool-use learning in chimpanzees, naïve individuals will generally copy dominant (and knowledgeable) individuals [71]. Our study hence stands as a clear cut illustration of the “copy when asocial learning is costly” rule [72]: the vulture position in the dominance hierarchy, through the costs it imposes on access to food, seems to calibrate the balance between the use of personal and social information in foraging movements. In some cases, however, trading personal information in favour of social information is inevitable.

When the environment is largely unpredictable or whenever using error-prone personal knowledge can be energetically costly, individuals should tend to eavesdrop, and rely more on information provided by conspecifics to reduce uncertainty about resources availability [15,73]. Here, we evidenced both cases. First, vultures prioritised the use of social information when the temperature was low and tended to when cloudiness and wind speed increased. These weather conditions may translate into fewer and weaker thermals, drifting into the wind, making them less predictable [74–78]. Second, they also favoured social information when the altitude at which they left their previous thermal was low. When exiting a thermal at low altitude, individuals have limited time to glide to the next thermal before having to shift to flapping flight to stay aloft, or else landing in an undesired place, which both would add high energetic cost associated with flapping and take-off [23,24,79,80]. Reaching high altitudes quickly to avoid this risk may also explain why vultures used more thermals previously discovered by conspecifics if those were close to the last thermal they used. While vultures are able to cope with difficult flight conditions (e.g. turbulence and strong wind) by adjusting their banking angles [55], anticipating such risky events may remain the most efficient way to maximise the trade-off between time, energy and risk which largely dictates their flight strategy [81]. Adult individuals, through experience, are generally better at coping with difficult flight conditions [82], yet we did not evidence an effect of age relative to the use of social information, as observed in other group living species (e.g. [83]). More than age *per se*, the familiarity of individuals with a given situation might shape their tendency to rely or not on social knowledge (e.g. in spider monkeys, *Ateles geoffroyi*, during collective foraging [84]). The captive individuals tested in this experiment are all adults and fly in the same landscape every day since their birth, thus they are probably very familiar with the areas favourable to thermal emergence. This could explain why we did not detect any effect of age on the use of social information, but also indicates that the relative importance of this source of information is probably underestimated due to the birds' familiarity with the surroundings. An interesting

complementary experiment (though technically difficult) to disentangle the effect of familiarity with the landscape would be to move the whole group and repeat the experiment in an unfamiliar area to better estimate the strength of personal versus social information use.

When faced with a choice between simultaneously available thermals, the previous experience of individuals (i.e. whether the thermal was used previously or not by the focal) or current expertise of the group (i.e., relative age/hierarchy difference) impacted very little vulture movement decisions compared to other social cues, contrasting with previous findings from insects to mammals, including birds [85–89]. In the current system, ascending currents can be very ephemeral phenomena, sometimes only lasting a few minutes [90,91]. Certainly, a “live report” is therefore better provided by the accumulation of convergent information sources (i.e. numerous conspecifics, [92]) rather than relying on a unique individual source (i.e. the individual itself or one reference individual). In that line, and surprisingly, the presence of one preferred affiliate in a thermal tended to reduce the probability to join it. There is evidence that social bonds assessed “on the ground” are often unrelated to co-flight preferences [93]. It therefore questions whether collective flights might be used by vultures to strengthen initially weak social bonds. Maintaining in-flight bounds can indeed be important, as evidenced in the migratory behaviour of other soaring bird species to enable accurate collective mapping of the distribution of uplifts [94,95]. Furthermore, for soaring birds, the presence of conspecifics should provide not only information on the location and strength of updrafts [20,95] but could also indicate flight speed and circling radius needed to optimise climb rate, by remaining close to the centre of the thermal where uplift is highest [55]. Yet, the maximum speed reached by individuals using the thermal little affected vulture decision choices. Possibly, climb rate or individual speed are not as easy to assess at a distance, compared to the number of conspecifics. In other words, vultures tended to favour quantity signals (with the number of conspecifics) over quality signals (maximal vertical speed) [96]. The “power of the group” may indeed in turn drive cohesion, which could itself make social information even more profitable [96,97].

Altogether, our results provide insights into the architecture of decision-making during movement in a social bird. It highlighted the trade-offs between personal and social information these birds have to consider in order to optimise both their flying efficiency and their foraging success. As a first approximation, we considered social cues as coming from “conspecifics”. Strictly speaking however, our study included two species, Griffon vulture and Rüppell’s vulture, albeit phylogenetically close and with similar biology. The one Rüppell’s vulture in fact, used social information provided by surrounding vultures and did not stand out as an outlier in its behaviour. It is known that even phylogenetically distant individuals could

426 be an important source of social information, not only about the presence of carcasses [98], but
 427 also about the availability of thermals when sharing the same airspace (e.g. from black kites,
 428 *Milvus migrans*, or common swifts, *Apus apus*, [99,100]). Interactions with heterospecifics can
 429 indeed drastically affect animals' daily life [101], up to shaping the cognitive machinery
 430 underpinning their foraging decisions [102]. How heterospecific cues are used when foraging
 431 remains clearly overlooked. Future studies in this direction could provide valuable insights into
 432 understanding the fundamental rules dictating how animals decide where to go.

433 CRediT authors' contributions

434 **Yohan Sassi:** Conceptualization, Methodology, Software, Investigation, Formal Analysis,
 435 Visualization, Writing - original draft, Writing - Review & Editing.
 436 **Basile Nousièrès:** Investigation, Resources.
 437 **Martina Scacco:** Software, Writing - Review & Editing.
 438 **Yann Tremblay:** Conceptualization, Writing - Review & Editing.
 439 **Olivier Duriez:** Conceptualization, Investigation, Supervision, Writing - Review & Editing.
 440 **Benjamin Robira:** Conceptualization, Methodology, Software, Visualization, Validation,
 441 Supervision, Writing - original draft, Writing - Review & Editing

442 Declaration of competing interest

443 The authors declare to have no conflict of interest

444 Data availability

445 Scripts for review and supplementary video are available here:
 446 <https://github.com/YohanSassi/UpdraftsDecisions>

447 A perennial storage of data and scripts will be provided after revision (e.g. Zenodo, Dryad)

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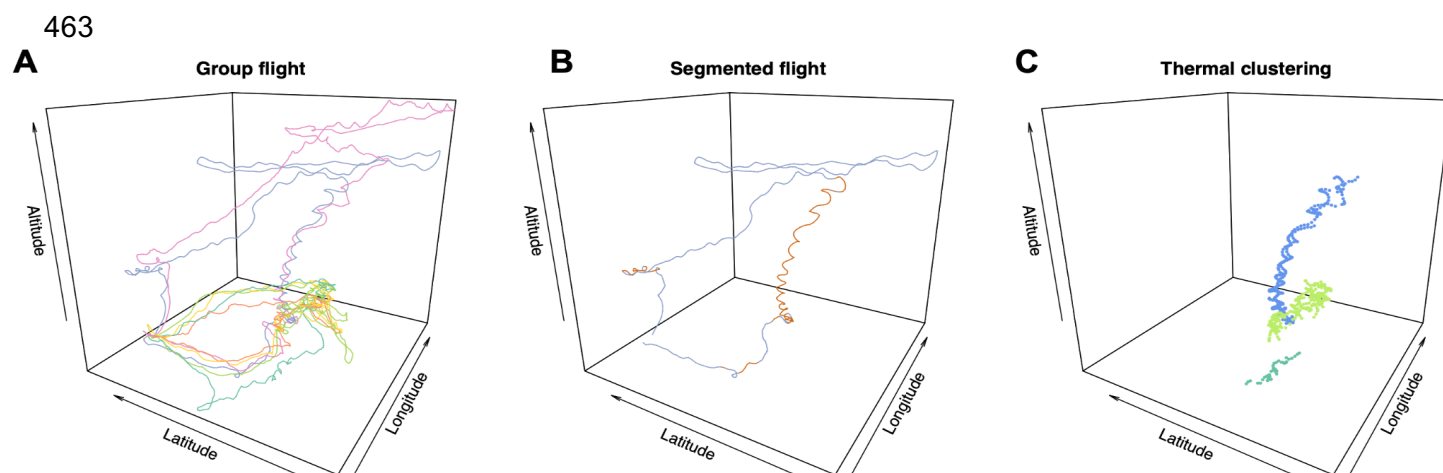
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457 Main text figures



458 **Figure 1. Data collection.** (A) Perched vultures. Distance between vultures during perching
 459 events were used to estimate social-bond strength. (B) Feeding event around a butchery carcass.
 460 Agonistic interactions during those feeding events were used to estimate dominance hierarchy.
 461 (C) Flying vulture. Vultures were released for free flight into a 120-m canyon, equipped with
 462 high-resolution GPS loggers.



464 **Figure 2. Flight data pre-processing.** Pre-processing steps of group flight GPS data, example
 465 of one flight session. The altitude ranges from 200 m to 600 m. (A) shows a group flight (see
 466 Supplementary Video 1), with colours corresponding to each individual. (B) illustrates the
 467 segmentation of an individual's flight (blue individual in (A)), with the orange segments
 468 corresponding to circular soaring phases. (C) illustrates the 3D density-based spatial clustering
 469 of individuals circular soaring phases, with colours indicating the three thermals identified in
 470 this flight session.

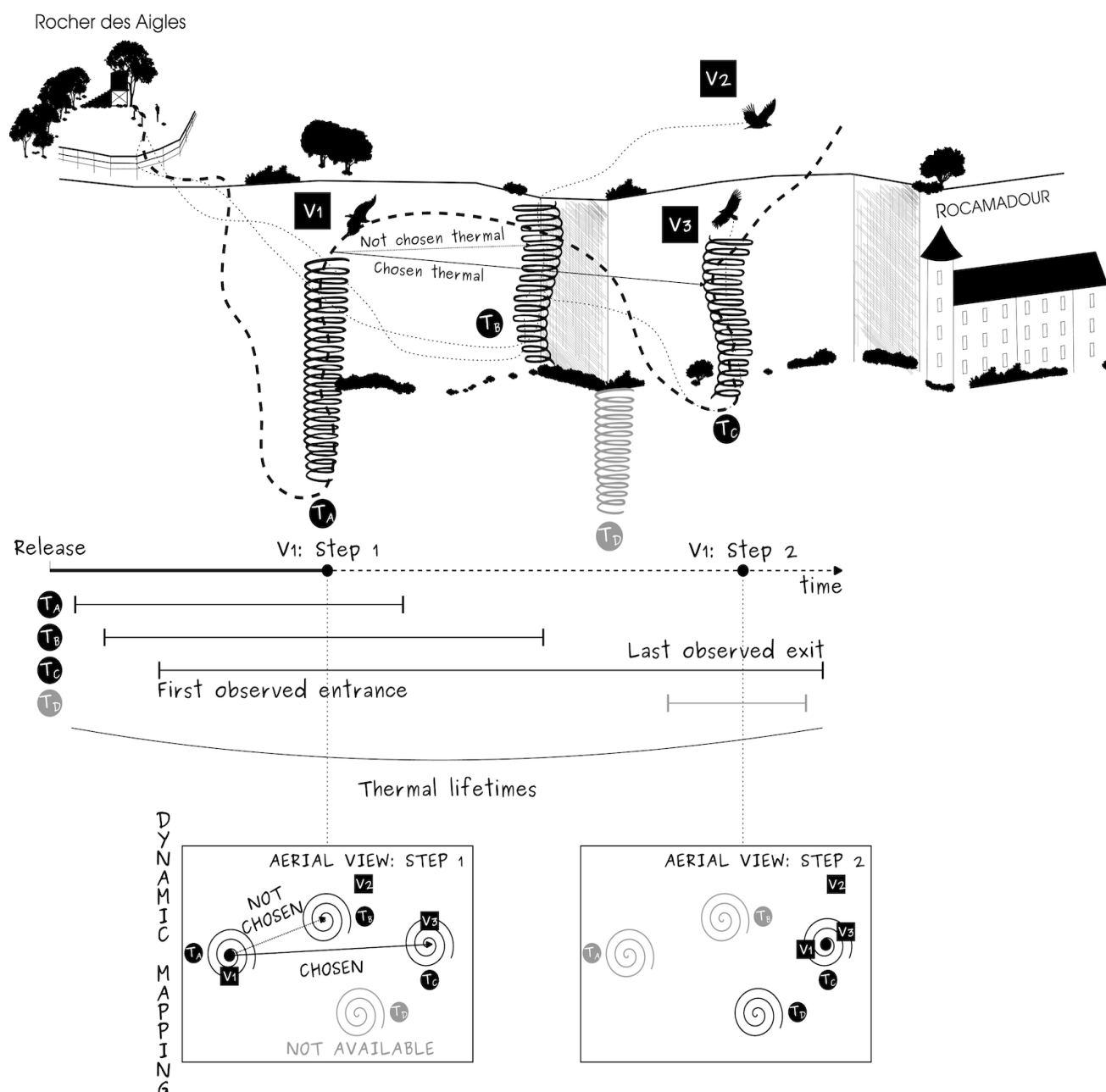
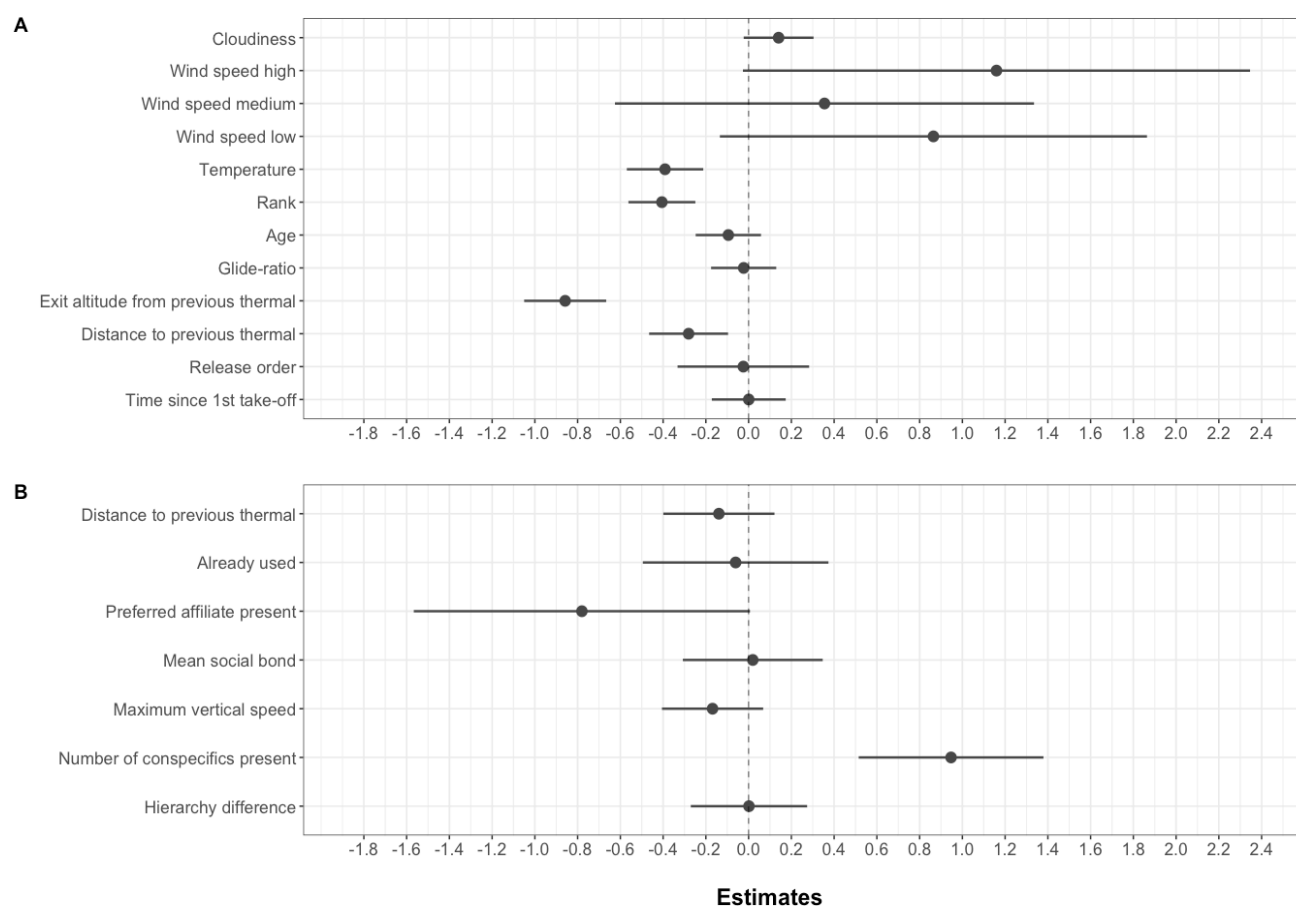


Figure 3. Illustration of the step selection framework used to investigate thermal selection. We focused on the movement of vultures released from the Rocher des Aigles when flying from thermal to thermal (i.e. a step). To do this, we mapped each thermal used during a flight session based on movement segmentation and clustering (see method section) to create dynamic maps of thermal availability over the flight session (as represented by the aerial views). The illustrated example focuses on the decision of a vulture (V₁; step 1) when leaving the thermal (T_A) and having to choose between two available thermals (T_B, close but not currently used by another vulture, and T_C, further away but currently used by another individual). T_D was not available until step 2, when it was discovered and used by another individual, and is therefore shown in grey at step 1. At step 2, V₁ joined V₃ in T_C and both

481 thermals T_A and T_B were no longer available. A thermal was available from the moment when
482 the first individual entered it until the last individual left it. Therefore, the number of available
483 thermals could change during the flight session (see differences between maps in step 1 and 2).



484 **Figure 4. Estimates of models investigating the drivers of social information use (A) and**
 485 **thermal selection (B).** Rows correspond to each predictor. Each point represents the
 486 standardised estimate value. Segments give the associated 95% confidence intervals.

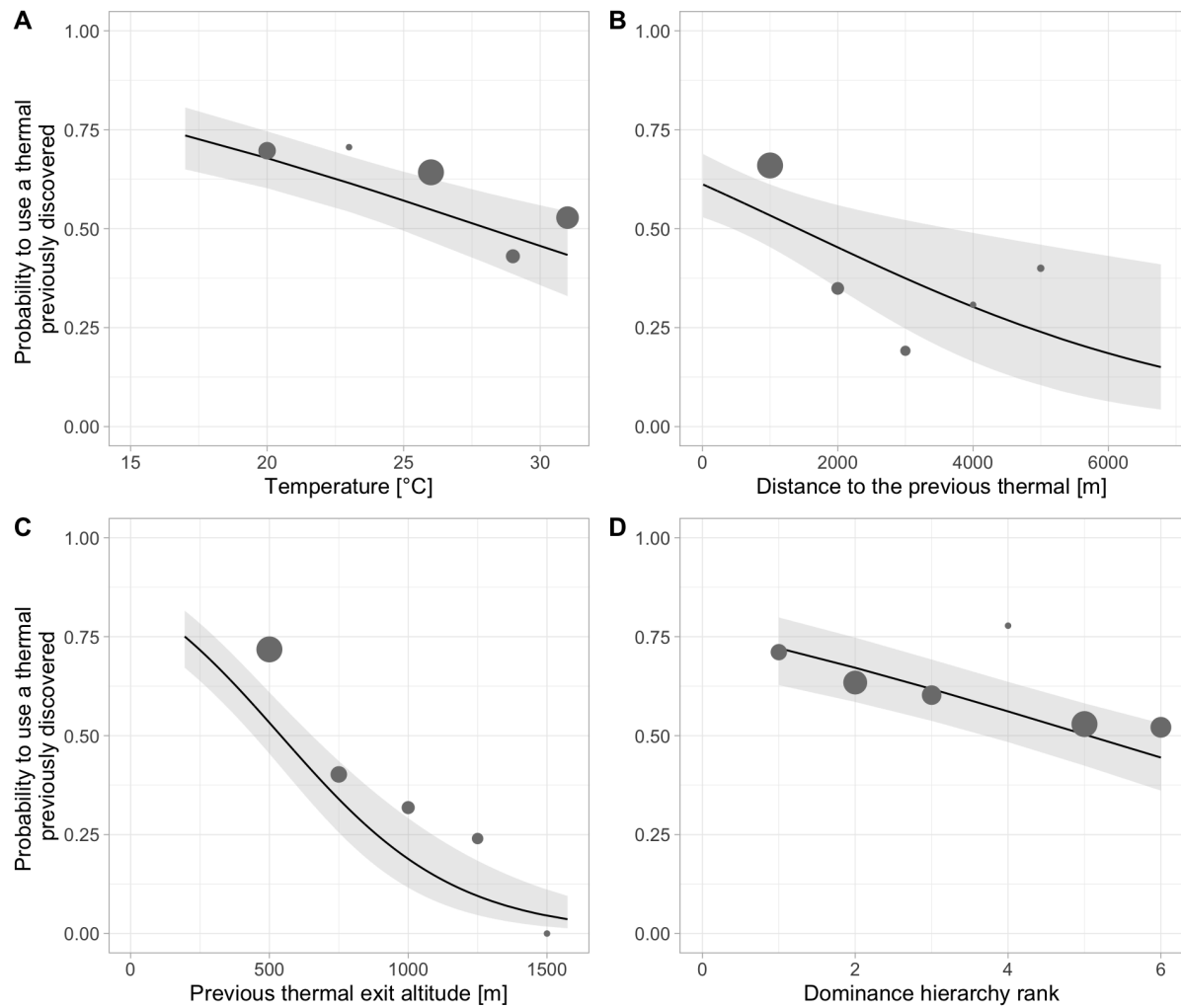


Figure 5. The probability to use a thermal already discovered by conspecifics decreases with temperature, distance to the thermal, flight altitude and hierarchy rank. Points represent the probability of using a thermal already discovered by a conspecific, estimated on the raw data. Their size is relative to the sample size. To do so, for (A), (B) and (C), predictors were binned. Black lines with grey shades show the GLMM estimated probability with its 95% confidence interval (N = 1098 thermals).

493 Reference

- 494 1. Dall SRX, Giraldeau L-A, Olsson O, McNamara JM, Stephens DW. 2005 Information and
495 its use by animals in evolutionary ecology. *Trends Ecol. Evol.* **20**, 187–193.
496 (doi:10.1016/j.tree.2005.01.010)
- 497 2. Shettleworth SJ, Krebs JR, Horn G. 1997 Spatial memory in food-storing birds. *Philos.*
498 *Trans. R. Soc. Lond. B. Biol. Sci.* **329**, 143–151. (doi:10.1098/rstb.1990.0159)
- 499 3. Valone TJ. 2007 From eavesdropping on performance to copying the behavior of others:
500 a review of public information use. *Behav. Ecol. Sociobiol.* **62**, 1–14.
501 (doi:10.1007/s00265-007-0439-6)
- 502 4. Danchin É, Giraldeau L-A, Valone TJ, Wagner RH. 2004 Public Information: From Nosy
503 Neighbors to Cultural Evolution. *Science* **305**, 487–491. (doi:10.1126/science.1098254)
- 504 5. Thiebault A, Mullers R, Pistorius P, Meza-Torres MA, Dubroca L, Green D, Tremblay Y.
505 2014 From colony to first patch: Processes of prey searching and social information in
506 Cape Gannets. *The Auk* **131**, 595–609. (doi:10.1642/AUK-13-209.1)
- 507 6. Fawcett TW, Fallenstein B, Higginson AD, Houston AI, Mallpress DEW, Trimmer PC,
508 McNamara JM. 2014 The evolution of decision rules in complex environments. *Trends*
509 *Cogn. Sci.* **18**, 153–161. (doi:10.1016/j.tics.2013.12.012)
- 510 7. Flack A, Pettit B, Freeman R, Guilford T, Biro D. 2012 What are leaders made of? The
511 role of individual experience in determining leader–follower relations in homing pigeons.
512 *Anim. Behav.* **83**, 703–709. (doi:10.1016/j.anbehav.2011.12.018)
- 513 8. McComb K, Shannon G, Durant SM, Sayialel K, Slotow R, Poole J, Moss C. 2011
514 Leadership in elephants: the adaptive value of age. *Proc. R. Soc. B Biol. Sci.* **278**, 3270–
515 3276. (doi:10.1098/rspb.2011.0168)
- 516 9. Strandburg-Peshkin A, Farine DR, Couzin ID, Crofoot MC. 2015 Shared decision-
517 making drives collective movement in wild baboons. *Science* **348**, 1358–1361.
518 (doi:10.1126/science.aaa5099)
- 519 10. Bode NWF, Wood AJ, Franks DW. 2011 The impact of social networks on animal
520 collective motion. *Anim. Behav.* **82**, 29–38. (doi:10.1016/j.anbehav.2011.04.011)
- 521 11. King AJ, Sueur C, Huchard E, Cowlshaw G. 2011 A rule-of-thumb based on social
522 affiliation explains collective movements in desert baboons. *Anim. Behav.* **82**, 1337–
523 1345. (doi:10.1016/j.anbehav.2011.09.017)
- 524 12. Farine DR, Strandburg-Peshkin A, Berger-Wolf T, Ziebart B, Brugere I, Li J, Crofoot MC.
525 2016 Both Nearest Neighbours and Long-term Affiliates Predict Individual Locations
526 During Collective Movement in Wild Baboons. *Sci. Rep.* **6**, 27704.
527 (doi:10.1038/srep27704)
- 528 13. Dunlap AS, Nielsen ME, Dornhaus A, Papaj DR. 2016 Foraging Bumble Bees Weigh the
529 Reliability of Personal and Social Information. *Curr. Biol.* **26**, 1195–1199.
530 (doi:10.1016/j.cub.2016.03.009)
- 531 14. Rafacz M, Templeton JJ. 2003 Environmental Unpredictability and the Value of Social
532 Information for Foraging Starlings. *Ethology* **109**, 951–960. (doi:10.1046/j.0179-
533 1613.2003.00935.x)
- 534 15. Williams HJ, Safi K. 2021 Certainty and integration of options in animal movement.
535 *Trends Ecol. Evol.* **36**, 990–999. (doi:10.1016/j.tree.2021.06.013)
- 536 16. Lee AEG, Ounsley JP, Coulson T, Rowcliffe JM, Cowlshaw G. 2016 Information use
537 and resource competition: an integrative framework. *Proc. R. Soc. B Biol. Sci.* **283**,
538 20152550. (doi:10.1098/rspb.2015.2550)
- 539 17. Rubenstein D I. 1982 Risk, uncertainty and evolutionary strategies. *Curr. Probl.*
540 *Sociobiol.* , 91–111.
- 541 18. Stillman RA, Caldow RWG, Goss-Custard JD, Alexander MJ. 2000 Individual variation in
542 intake rate: the relative importance of foraging efficiency and dominance. *J. Anim. Ecol.*
543 **69**, 484–493. (doi:10.1046/j.1365-2656.2000.00410.x)
- 544 19. Goss-custard JD, Cayford JT, Lea SEG. 1998 The changing trade-off between food

- finding and food stealing in juvenile oystercatchers. *Anim. Behav.* **55**, 745–760.
(doi:10.1006/anbe.1997.0680)
20. Pennycuik CJ. 1972 Soaring behaviour and performance of some East African birds observed from a motor glider. *Ibis* **114**, 178–218.
21. Scacco M, Flack A, Duriez O, Wikelski M, Safi K. 2019 Static landscape features predict uplift locations for soaring birds across Europe. *R. Soc. Open Sci.* **6**, 181440.
(doi:10.1098/rsos.181440)
22. Alarcón PAE, Morales JM, Donázar JA, Sánchez-Zapata JA, Hiraldo F, Lambertucci SA. 2017 Sexual-size dimorphism modulates the trade-off between exploiting food and wind resources in a large avian scavenger. *Sci. Rep.* **7**, 11461. (doi:10.1038/s41598-017-11855-0)
23. Horvitz N, Sapir N, Liechti F, Avissar R, Mahrer I, Nathan R. 2014 The gliding speed of migrating birds: slow and safe or fast and risky? *Ecol. Lett.* **17**, 670–679.
(doi:10.1111/ele.12268)
24. Duriez O, Kato A, Tromp C, Dell'Omo G, Vyssotski AL, Sarrazin F, Ropert-Coudert Y. 2014 How Cheap Is Soaring Flight in Raptors? A Preliminary Investigation in Freely-Flying Vultures. *PLoS ONE* **9**, e84887. (doi:10.1371/journal.pone.0084887)
25. Jackson AL, Ruxton GD, Houston DC. 2008 The effect of social facilitation on foraging success in vultures: a modelling study. *Biol. Lett.* **4**, 311–313.
(doi:10.1098/rsbl.2008.0038)
26. Cortés-Avizanda A, Jovani R, Donázar JA, Grimm V. 2014 Bird sky networks: How do avian scavengers use social information to find carrion? *Ecology* **95**, 1799–1808.
(doi:10.1890/13-0574.1)
27. Bosè M, Duriez O, Sarrazin F. 2012 Intra-specific competition in foraging Griffon Vultures *Gyps fulvus*: 1. Dynamics of group feeding. *Bird Study* **59**, 182–192.
(doi:10.1080/00063657.2012.658639)
28. Moreno-Opo R, Trujillano A, Margalida A. 2015 Optimization of supplementary feeding programs for European vultures depends on environmental and management factors. *Ecosphere* **6**, art127. (doi:10.1890/es15-00009.1)
29. Williams HJ, King AJ, Duriez O, Börger L, Shepard ELC. 2018 Social eavesdropping allows for a more risky gliding strategy by thermal-soaring birds. *J. R. Soc. Interface* **15**, 20180578. (doi:10.1098/rsif.2018.0578)
30. Bosè M, Sarrazin F. 2007 Competitive behaviour and feeding rate in a reintroduced population of Griffon Vultures *Gyps fulvus*: Competition for food in Griffon Vultures. *Ibis* **149**, 490–501. (doi:10.1111/j.1474-919X.2007.00674.x)
31. Kirk DA, Houston DC. 1995 Social dominance in migrant and resident turkey vultures at carcasses: evidence for a despotic distribution? *Behav. Ecol. Sociobiol.* **36**, 323–332.
(doi:10.1007/BF00167793)
32. Donázar JA, Travaini A, Ceballos O, Rodríguez A, Delibes M, Hiraldo F. 1999 Effects of sex-associated competitive asymmetries on foraging group structure and despotic distribution in Andean condors. *Behav. Ecol. Sociobiol.* **45**, 55–65.
(doi:10.1007/s002650050539)
33. van Overveld T, García-Alfonso M, Dingemanse NJ, Bouten W, Gangoso L, de la Riva M, Serrano D, Donázar JA. 2018 Food predictability and social status drive individual resource specializations in a territorial vulture. *Sci. Rep.* **8**, 15155. (doi:10.1038/s41598-018-33564-y)
34. Duriez O, Herman S, Sarrazin F. 2012 Intra-specific competition in foraging Griffon Vultures *Gyps fulvus*: 2. The influence of supplementary feeding management. *Bird Study* **59**, 193–206. (doi:10.1080/00063657.2012.658640)
35. Moreno-Opo R, Trujillano A, Margalida A. 2016 Behavioral coexistence and feeding efficiency drive niche partitioning in European avian scavengers. *Behav. Ecol.* **27**, 1041–1052. (doi:10.1093/beheco/arw010)
36. Kohn GM, Meredith GR, Magdaleno FR, King AP, West MJ. 2015 Sex differences in familiarity preferences within fission–fusion brown-headed cowbird, *Molothrus ater*, flocks. *Anim. Behav.* **106**, 137–143. (doi:10.1016/j.anbehav.2015.05.023)

37. Voelkl B, Fritz J. 2017 Relation between travel strategy and social organization of migrating birds with special consideration of formation flight in the northern bald ibis. *Philos. Trans. R. Soc. B Biol. Sci.* **372**, 20160235. (doi:10.1098/rstb.2016.0235)
38. Utne-Palm AC, Hart PJB. 2000 The effects of familiarity on competitive interactions between threespined sticklebacks. *Oikos* **91**, 225–232. (doi:https://doi.org/10.1034/j.1600-0706.2000.910203.x)
39. Fluhr J, Benhamou S, Peyrusque D, Duriez O. 2021 Space Use and Time Budget in Two Populations of Griffon Vultures in Contrasting Landscapes. *J. Raptor Res.* **55**, 425–437. (doi:10.3356/JRR-20-14)
40. Bezanson J, Karpinski S, Shah VB, Edelman A. 2012 Julia: A Fast Dynamic Language for Technical Computing. (doi:10.48550/arXiv.1209.5145)
41. R Core Team. 2022 R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
42. Whitehead H. 2008 *Analyzing Animal Societies: Quantitative Methods for Vertebrate Social Analysis*. University of Chicago Press.
43. Hoppitt WJE, Farine DR. 2018 Association indices for quantifying social relationships: how to deal with missing observations of individuals or groups. *Anim. Behav.* **136**, 227–238. (doi:10.1016/j.anbehav.2017.08.029)
44. Elo AE. 1978 *The rating of chessplayers, past and present*. Arco Publication. New York.
45. Sánchez-Tójar A, Schroeder J, Farine DR. 2018 A practical guide for inferring reliable dominance hierarchies and estimating their uncertainty. *J. Anim. Ecol.* **87**, 594–608. (doi:10.1111/1365-2656.12776)
46. Farine D, Sánchez-Tójar. 2018 aniDom: inferring dominance hierarchies and estimating uncertainty.
47. Drews C. 1993 The Concept and Definition of Dominance in Animal Behaviour. *Behaviour* **125**, 283–313. (doi:10.1163/156853993X00290)
48. Friard O, Gamba M. 2016 BORIS: a free, versatile open-source event-logging software for video/audio coding and live observations. *Methods Ecol. Evol.* **7**, 1325–1330. (doi:10.1111/2041-210X.12584)
49. Valverde JA. 1959 Moyens d'expression et hiérarchie sociale chez le vautour fauve *Gyps fulvus*. *Alauda* **1**, 1–15.
50. Anderson D *et al.* 2020 A practical guide to methods for attaching research devices to vultures and condors. (doi:10.17863/CAM.58032)
51. Kranstauber B, Smolla M, Scharf AK. 2018 move: Visualizing and analyzing animal track data. R package version 3.1.0.
52. Williams HJ, Shepard ELC, Duriez O, Lambertucci SA. 2015 Can accelerometry be used to distinguish between flight types in soaring birds? *Anim. Biotelemetry* **3**, 45. (doi:10.1186/s40317-015-0077-0)
53. Khosravifard S, Venus V, Skidmore AK, Bouten W, Muñoz AR, Toxopeus AG. 2018 Identification of Griffon Vulture's Flight Types Using High-Resolution Tracking Data. *Int. J. Environ. Res.* **12**, 313–325. (doi:10.1007/s41742-018-0093-z)
54. Hahsler M, Piekenbrock M, Arya S, Mount D. 2017 DbSCAN: density based clustering of applications with noise (DBSCAN) and related algorithms.
55. Williams HJ, Duriez O, Holton MD, Dell'Omo G, Wilson RP, Shepard ELC. 2018 Vultures respond to challenges of near-ground thermal soaring by varying bank angle. *J. Exp. Biol.* **221**, jeb174995. (doi:10.1242/jeb.174995)
56. Bolker BM, Brooks ME, Clark CJ, Geange SW, Poulsen JR, Stevens MHH, White J-SS. 2009 Generalized linear mixed models: a practical guide for ecology and evolution. *Trends Ecol. Evol.* **24**, 127–135. (doi:10.1016/j.tree.2008.10.008)
57. Schielzeth H. 2010 Simple means to improve the interpretability of regression coefficients. *Methods Ecol. Evol.* **1**, 103–113. (doi:https://doi.org/10.1111/j.2041-210X.2010.00012.x)
58. Hartig F, Lohse L. 2020 DHARMA: Residual Diagnostics for Hierarchical (Multi-Level / Mixed) Regression Models.
59. Zuur AF, Ieno EN, Elphick CS. 2010 A protocol for data exploration to avoid common

- statistical problems. *Methods Ecol. Evol.* **1**, 3–14. (doi:<https://doi.org/10.1111/j.2041-210X.2009.00001.x>)
60. Nakagawa S, Schielzeth H. 2013 A general and simple method for obtaining R² from generalized linear mixed-effects models. *Methods Ecol. Evol.* **4**, 133–142. (doi:<https://doi.org/10.1111/j.2041-210x.2012.00261.x>)
61. Thurfjell H, Ciuti S, Boyce MS. 2014 Applications of step-selection functions in ecology and conservation. *Mov. Ecol.* **2**, 4. (doi:[10.1186/2051-3933-2-4](https://doi.org/10.1186/2051-3933-2-4))
62. Avgar T, Lele SR, Keim JL, Boyce MS. 2017 Relative Selection Strength: Quantifying effect size in habitat- and step-selection inference. *Ecol. Evol.* **7**, 5322–5330. (doi:[10.1002/ece3.3122](https://doi.org/10.1002/ece3.3122))
63. Fieberg J, Signer J, Smith B, Avgar T. 2021 A ‘How to’ guide for interpreting parameters in habitat-selection analyses. *J. Anim. Ecol.* **90**, 1027–1043. (doi:[10.1111/1365-2656.13441](https://doi.org/10.1111/1365-2656.13441))
64. Webster MM, Laland KN. 2012 Social information, conformity and the opportunity costs paid by foraging fish. *Behav. Ecol. Sociobiol.* **66**, 797–809. (doi:[10.1007/s00265-012-1328-1](https://doi.org/10.1007/s00265-012-1328-1))
65. Barnard CJ, Sibly RM. 1981 Producers and scroungers: A general model and its application to captive flocks of house sparrows. *Anim. Behav.* **29**, 543–550. (doi:[10.1016/S0003-3472\(81\)80117-0](https://doi.org/10.1016/S0003-3472(81)80117-0))
66. Barta Z, Giraldeau L-A. 1998 The effect of dominance hierarchy on the use of alternative foraging tactics: a phenotype-limited producing-scrounging game. *Behav. Ecol. Sociobiol.* **42**, 217–223. (doi:[10.1007/s002650050433](https://doi.org/10.1007/s002650050433))
67. Stahl J, Tolsma PH, Loonen MJJE, Drent RH. 2001 Subordinates explore but dominants profit: resource competition in high Arctic barnacle goose flocks. *Anim. Behav.* **61**, 257–264. (doi:[10.1006/anbe.2000.1564](https://doi.org/10.1006/anbe.2000.1564))
68. Liker A, Barta Z. 2002 THE EFFECTS OF DOMINANCE ON SOCIAL FORAGING TACTIC USE IN HOUSE SPARROWS. *Behaviour* **139**, 1061–1076. (doi:[10.1163/15685390260337903](https://doi.org/10.1163/15685390260337903))
69. Alonso JC, Bautista LM, Alonso JA. 1997 Dominance and the dynamics of phenotype-limited distribution in common cranes. *Behav. Ecol. Sociobiol.* **40**, 401–408. (doi:[10.1007/s002650050356](https://doi.org/10.1007/s002650050356))
70. Papageorgiou D, Farine DR. 2020 Shared decision-making allows subordinates to lead when dominants monopolize resources. *Sci. Adv.* **6**, eaba5881. (doi:[10.1126/sciadv.aba5881](https://doi.org/10.1126/sciadv.aba5881))
71. Kendal R, Hopper LM, Whiten A, Brosnan SF, Lambeth SP, Schapiro SJ, Hoppitt W. 2015 Chimpanzees copy dominant and knowledgeable individuals: implications for cultural diversity. *Evol. Hum. Behav.* **36**, 65–72. (doi:[10.1016/j.evolhumbehav.2014.09.002](https://doi.org/10.1016/j.evolhumbehav.2014.09.002))
72. Laland KN. 2004 Social learning strategies. *Anim. Learn. Behav.* **32**, 4–14. (doi:[10.3758/BF03196002](https://doi.org/10.3758/BF03196002))
73. Riotte-Lambert L, Matthiopoulos J. 2020 Environmental Predictability as a Cause and Consequence of Animal Movement. *Trends Ecol. Evol.* **35**, 163–174. (doi:[10.1016/j.tree.2019.09.009](https://doi.org/10.1016/j.tree.2019.09.009))
74. Bradbury T. 1989 *Meteorology and flight: a pilot's guide to weather*. A&C Black.
75. Shannon HD, Young GS, Yates MA, Fuller MR, Seegar WS. 2002 American White Pelican Soaring Flight Times and Altitudes Relative to Changes in Thermal Depth and Intensity. *The Condor* **104**, 679–683. (doi:[10.1093/condor/104.3.679](https://doi.org/10.1093/condor/104.3.679))
76. Shamoun-Baranes J, Baharad A, Alpert P, Berthold P, Yom-Tov Y, Dvir Y, Leshem Y. 2003 The effect of wind, season and latitude on the migration speed of white storks *Ciconia ciconia*, along the eastern migration route. *J. Avian Biol.* **34**, 97–104. (doi:[10.1034/j.1600-048X.2003.03079.x](https://doi.org/10.1034/j.1600-048X.2003.03079.x))
77. Shamoun-Baranes J, Bouten W, van Loon EE, Meijer C, Camphuysen CJ. 2016 Flap or soar? How a flight generalist responds to its aerial environment. *Philos. Trans. R. Soc. B Biol. Sci.* **371**, 20150395. (doi:[10.1098/rstb.2015.0395](https://doi.org/10.1098/rstb.2015.0395))
78. Kerlinger P, Gauthreaux SA. 1984 Flight behaviour of sharp-shinned hawks during

- migration. I : Over land. *Anim. Behav.* **32**, 1021–1028. (doi:10.1016/S0003-3472(84)80216-X)
79. Shepard ELC. 2022 Energy economy in flight. *Curr. Biol.* **32**, R672–R675. (doi:10.1016/j.cub.2022.02.004)
80. Williams HJ, Shepard ELC, Holton MD, Alarcón PAE, Wilson RP, Lambertucci SA. 2020 Physical limits of flight performance in the heaviest soaring bird. *Proc. Natl. Acad. Sci.* **117**, 17884–17890. (doi:10.1073/pnas.1907360117)
81. Harel R et al. 2016 Decision-making by a soaring bird: time, energy and risk considerations at different spatio-temporal scales. *Philos. Trans. R. Soc. B Biol. Sci.* **371**, 20150397. (doi:10.1098/rstb.2015.0397)
82. Harel R, Horvitz N, Nathan R. 2016 Adult vultures outperform juveniles in challenging thermal soaring conditions. *Sci. Rep.* **6**, 27865. (doi:10.1038/srep27865)
83. Dugatkin LA, Godin J-GJ. 1993 Female mate copying in the guppy (*Poecilia reticulata*): age-dependent effects. *Behav. Ecol.* **4**, 289–292. (doi:10.1093/beheco/4.4.289)
84. Palacios-Romo TM, Castellanos F, Ramos-Fernandez G. 2019 Uncovering the decision rules behind collective foraging in spider monkeys. *Anim. Behav.* **149**, 121–133. (doi:10.1016/j.anbehav.2019.01.011)
85. Baciadonna L, McElligott AG, Briefer EF. 2013 Goats favour personal over social information in an experimental foraging task. *PeerJ* **1**, e172. (doi:10.7717/peerj.172)
86. Heinen VK, Stephens DW. 2016 Blue jays, *Cyanocitta cristata*, devalue social information in uncertain environments. *Anim. Behav.* **112**, 53–62. (doi:10.1016/j.anbehav.2015.11.015)
87. Leadbeater E, Florent C. 2014 Foraging bumblebees do not rate social information above personal experience. *Behav. Ecol. Sociobiol.* **68**, 1145–1150. (doi:10.1007/s00265-014-1725-8)
88. Grüter C, Czaczkes TJ, Ratnieks FLW. 2011 Decision making in ant foragers (*Lasius niger*) facing conflicting private and social information. *Behav. Ecol. Sociobiol.* **65**, 141–148. (doi:10.1007/s00265-010-1020-2)
89. Biro D, Inoue-Nakamura N, Tonooka R, Yamakoshi G, Sousa C, Matsuzawa T. 2003 Cultural innovation and transmission of tool use in wild chimpanzees: evidence from field experiments. *Anim. Cogn.* **6**, 213–223. (doi:10.1007/s10071-003-0183-x)
90. Reddy G, Celani A, Sejnowski TJ, Vergassola M. 2016 Learning to soar in turbulent environments. *Proc. Natl. Acad. Sci.* **113**, E4877–E4884. (doi:10.1073/pnas.1606075113)
91. Westbrook JK, Eyster RS. 2017 Atmospheric Environment Associated with Animal Flight. In *Aeroecology* (eds PB Chilson, WF Frick, JF Kelly, F Liechti), pp. 13–45. Cham: Springer International Publishing. (doi:10.1007/978-3-319-68576-2_2)
92. King AJ, Cowlshaw G. 2007 When to use social information: the advantage of large group size in individual decision making. *Biol. Lett.* **3**, 137–139. (doi:10.1098/rsbl.2007.0017)
93. Sharma N, Anglister N, Spiegel O, Pinter-Wollman N. 2022 Social situations differ in their contribution to population-level social structure in griffon vultures. (doi:10.22541/au.165107196.66953827/v1)
94. Nagy M, Couzin ID, Fiedler W, Wikelski M, Flack A. 2018 Synchronization, coordination and collective sensing during thermalling flight of freely migrating white storks. *Philos. Trans. R. Soc. B Biol. Sci.* **373**, 20170011. (doi:10.1098/rstb.2017.0011)
95. van Loon EE, Shamoun-Baranes J, Bouten W, Davis SL. 2011 Understanding soaring bird migration through interactions and decisions at the individual level. *J. Theor. Biol.* **270**, 112–126. (doi:10.1016/j.jtbi.2010.10.038)
96. Fernández-Juricic E, Kacelnik A. 2004 Information transfer and gain in flocks: the effects of quality and quantity of social information at different neighbour distances. *Behav. Ecol. Sociobiol.* **55**, 502–511. (doi:10.1007/s00265-003-0698-9)
97. Bose T, Reina A, Marshall JA. 2017 Collective decision-making. *Curr. Opin. Behav. Sci.* **16**, 30–34. (doi:10.1016/j.cobeha.2017.03.004)
98. Kane A, Jackson AL, Ogada DL, Monadjem A, McNally L. 2014 Vultures acquire

765 information on carcass location from scavenging eagles. *Proc. R. Soc. B Biol. Sci.* **281**,
766 20141072. (doi:10.1098/rspb.2014.1072)
767 99. Hedrick TL, Pichot C, de Margerie E. 2018 Gliding for a free lunch: biomechanics of
768 foraging flight in common swifts (*Apus apus*). *J. Exp. Biol.* **221**, jeb186270.
769 (doi:10.1242/jeb.186270)
770 100. Santos CD, Hanssen F, Muñoz A-R, Onrubia A, Wikelski M, May R, Silva JP. 2017
771 Match between soaring modes of black kites and the fine-scale distribution of updrafts.
772 *Sci. Rep.* **7**, 6421. (doi:10.1038/s41598-017-05319-8)
773 101. Tremblay Y, Thiebault A, Mullers R, Pistorius P. 2014 Bird-Borne Video-Cameras Show
774 That Seabird Movement Patterns Relate to Previously Unrevealed Proximate
775 Environment, Not Prey. *PLOS ONE* **9**, e88424. (doi:10.1371/journal.pone.0088424)
776 102. Robira B, Perez-Lamarque B. 2023 Primate sympatry shapes the evolution of their brain
777 architecture. *Peer Community J.* **3**. (doi:10.24072/pcjournal.259)