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Optic flow helps explain gulls' altitude control over seas

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For studies of how flying animals control their flight, seabirds are of particular interest to track with a biologger because they forage offshore where the visual environment can be simply modeled by a flat world textured by waves. This study suggests that optic flow can explain gull's altitude control over seas. In particular, a new flight model that includes both energy and optical invariants (called the *ventral optic* flow regulation) explain the dynamics of gulls' altitude control during offshore takeoff and cruising flight. A linear statistical model applied to 352 flights from 16 individual lesser black backed gulls (Larus fuscus) gave a strong correlation between wind assistance and gulls' altitude. Thereafter, an optic flow-based flight model was applied to 18 offshore takeoff flights from 9 individual gulls. By introducing an upper limit in climb rate in a non-linear first order parametric model on the gull's elevation dynamics, coupled with an optic-flow set-point, the predicted altitude gives an optimized fit factor value of 63% on average (min value: 30%, max value: 83%) with respect to GPS data. We conclude that the optic-flow regulation principle (here running close to $25^{\circ}/s$) allows gulls to adjust their altitude over sea without having to directly measure their current altitude.

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2 1. Introduction

Understanding how a bird decides to fly at a given altitude during a specific manoeuver is a difficult task because it is strongly dependent on the atmospheric conditions and flight capacity of the bird (see review [59]). Seabirds such as albatrosses and petrels flying close to the sea surface take advantage of the logarithmic increase in wind speeds to support dynamic soaring [50, 52, 53, 66], which works only at very low altitudes from ca. 0-10 m (see e.g. Fig. 5 in [56]). Birds flying by flapping flight at low altitudes over the sea could also use this windspeed gradient to reduce their transport costs. Under tailwinds, birds should fly higher where wind speed is high, while under headwinds birds should fly lower where wind speed is low. In terms of energy, a bird minimizing 10 its transport cost should adjust its airspeed with respect to wind by increasing it in headwinds 11 and decreasing it in tailwinds [26, 48]. This prediction comes from a U-shaped function between 12 power required to fly and airspeed, which defines characteristic speeds for achieving minimum 13 power V_{mp} and maximum range V_{mr} . During migratory [38] and homing flights [33] birds utilize 14 wind assistance to minimize the transport cost and adjust airspeed accordingly to fly at the wind 15 dependent V_{mr} . 16

Groundspeed is the combined effect of airspeed and wind speed (actually the airspeed and wind vectors). Wind assistance alone cannot be used by the bird to select a given groundspeed

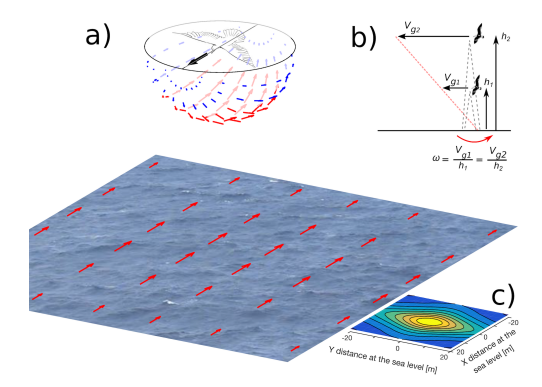


Figure 1. (a) A gull flying over the sea generates a vector field of optic flow. Such a vector field is perceived by a gull based on the contrasts created by waves and white-crested waves (also called white-horses). Inspired by [19]. (b) The magnitude of the vector of the optic flow, ω , is determined by the gull's groundspeed, V_g , and its altitude, h. If ω is held constant by adjusting the altitude, h will always tend (through the bird dynamics) to be proportional to V_g (only a linear combination -red dashed line- between h and V_g is asymptotically possible). (c) Optic flow magnitude in the ventral field of view at 10m-height where the magnitude of the ventral optic flow $\omega(\phi, \theta) = \frac{V_g}{h} sin^2\theta \times cos\phi$ is projected at the sea level with ϕ the azimuthal angle and θ the elevation angle. (The magnitude of vertical optic flow is the maximum and is $\omega(\phi = 0^\circ, \theta = -90^\circ) = \frac{V_g}{h}$)

and a flight altitude. The altitude could be set by surrounding visual information seen by 19 the bird. A bird can access information about its own motion with respect to its surrounding 20 environment via the optic flow field through its early visual processing [4], as flying insects do 21 in similar situations [4, 58]. The optic flow field perceived by an agent (a flying insect, a bird, 22 or a human) is particularly dependent on the structure of the environment [19, 35, 45, 67]. Optic 23 flow can be defined by a vector field of the apparent angular velocities of objects, surfaces, and 24 edges in a visual scene caused by the relative motion between the agent and the scene (Fig. 25 1). The translational optic flow component is particularly interesting for birds positioning in 26 space because it depends on (i) the ratio between the relative linear groundspeed of an object 27 in the scene with respect to the bird, and (ii) the distance from obstacles in the surrounding 28 environment. Consequently, optic flow requires neither groundspeed nor distance measurement, 29 which is particularly useful to explain how birds perceive the world because birds are likely 30 unable to sense directly their own groundspeed nor the 3D structure of the environment in which 31 the binocular vision plays a minor role [39]. 32

During flight manoeuvers, various optic flow parameters (such as the magnitude, the direction, the focus of expansion, the time-to-contact of optic flow) can be collected by birds to control their lateral position in straight tunnels (in budgerigars [7]), to decrease their speed in a converging tunnel (in budgerigars [57]), to plunge into water (in gannets [36]), to hover (in hummingbirds [20, 54]), and finally to land (in hawks [12] and in hummingbirds [37]).

In this study, we address the question of how seabirds control their altitude during offshore takeoffs and cruise flights with respect to wind. Here, two working hypotheses were compared about altitude control:

- a first hypothesis based on a direct measurement and regulation of optic flow that adjusts the altitude, and,
 - a second hypothesis based on a direct measurement of the barometric pressure that directly regulates the altitude itself.

To test these alternative hypotheses, a statistical analysis of 352 flights comprising 16 individual lesser black-backed gulls (*Larus fuscus*) in various wind conditions were conducted. Then, 18 offshore takeoffs followed by a cruise flight were analyzed by taking into account morphological parameters from 9 individual gulls.

2. Operating point in flight in terms of speed and altitude: a theoretical approach

(a) How is bird speed deducted from aeraulic effects?

⁵² The relationship between:

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- the bird's ground speed V_g
- the bird's airspeed V_{air}
- $_{55}$ the wind speed V_w
- ⁵⁶ is given by equation (2.1):

$$V_g = V_{air} + V_w \tag{2.1}$$

The basis for deriving predictions about bird flight is the so-called flight mechanical theory, which combines the relationship between power output P and airspeed V_{air} in flapping flight as follows:

$$P(V_{air}) = a + b \cdot V_{air}^{-1} + c \cdot V_{air}^{3}$$
(2.2)

where *a*, *b*, and *c* represent various physical, morphological and physiological properties of the bird and air [47, 49, 51]. If the objective is to minimize the energy cost per unit distance (i.e., cost of transport), the optimal flight speed is the maximum range speed V_{mr} [26, 47]. The maximum range speed V_{mr} is obtained from the U-shaped power curve [24, 27, 51] by the condition: bioRxiv preprint doi: https://doi.org/10.1101/569194; this version posted March 8, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

$$\left(\frac{\partial P}{\partial V_{air}}\right)_{V_{air}=V_{mr}} = \frac{P(V_{mr})}{V_{mr}} \tag{2.3}$$

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64 Indeed, a gull's homing flight is similar to a migratory flight, in that it is assumed that 65 the flight's objective is principally for transportation, as opposed to outbound foraging flights when the bird is likely also searching for food. Seabirds' homing flight over the sea is therefore 66 a relatively straight path between two locations. During transport flight gulls are expected to 67 minimise overall energy expenditure or time, thus cost of travel per unit distance should be 68 minimised rather than instantaneous energy expenditure. If minimising the cost of travel per 69 unit distance birds will travel at maximum range speed (V_{mr}) not minimum power airspeed 70 (V_{mp}) . V_{mr} refers to V_{air} rather than V_g . If a bird experiences a tailwind, its cost of travel per 71 unit distance decreases, thus V_{mr} also declines. Conversely under headwinds V_{mr} increases. In 72 a recent work, it was analyzed how lesser black-backed gulls (and guillemots) modulate their 73 airspeeds in relation to winds [17]. It was found that gulls increased airspeeds under headwinds 74 75 and decreased airspeeds under tailwinds [17], and similar behaviour has been observed during longer distance homing flights [42]. These results suggest that gulls are flying at V_{mr} rather than 76 V_{mp} , since V_{mp} should not be affected by winds like V_{mr} [17]. 77

78 (b) Optic flow vector field

⁷⁹ Consider a bird flying over the sea, assumed as flat in the optic flow calculation, then based on ⁸⁰ groundspeed V_g only (neglecting vertical speed V_z) the magnitude of the ventral optic flow field ⁸¹ ω can be expressed as follows:

$$\omega(\phi,\theta) = \frac{V_g}{h} \sin^2\theta \times \cos\phi \tag{2.4}$$

with *h* the altitude, θ the elevation angle and ϕ the azimuth angle.

The magnitude of the ventral optic flow field is plotted in Fig. 1a with the projection of its elevation and azimuth angles over the sea. The larger projection of vector magnitude of optic flow over the sea is shown using a contour plot in Fig. 1c in the case of a bird flying at a height of 10 m. The bird may be able to perceive the optic flow maximum from a non-negligible area of its field of view (Fig. 1c). The maximum magnitude of the ventral optic flow is always vertically downwards from the bird in the direction of the sea :

$$\omega(\phi = 0^{\circ}, \theta = -90^{\circ}) = \frac{V_g}{h}$$
(2.5)

(c) How the model predicts the bird's flight height from the ventral optic flow regulation principle

The ventral optic flow regulation principle tends to keep constant the vertically downward optic flow whatever the speed or height of flight by adjusting the altitude [18, 55]. Here, it introduces this asymptotic proportionality relationship for birds: the bird's height of flight *h* will always tend (through the bird dynamics) to be proportional to the bird's ground speed V_g (Fig. 1b) as :

$$\omega_{sp} = \omega(\phi = 0^{\circ}, \theta = -90^{\circ}) = \frac{V_g}{h} = \text{constant}$$
(2.6)

where ω_{sp} is the ventral optic flow set-point. Besides, the wind profile power-law is often used to estimate the horizontal wind speed [31] as follows:

$$V_w = V_{ref} \cdot \left(\frac{h}{h_{ref}}\right)^{\alpha} \tag{2.7}$$

with the parameter α is the power-law exponent (that is usually specified as a function of stability as well as the roughness of the surface 0< α <1 (here over seas $\alpha = 0.11$ see [30]), the speed

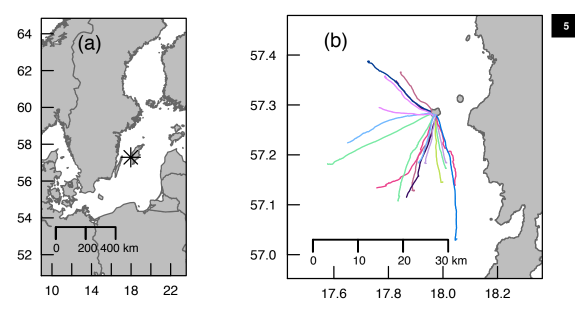


Figure 2. (a) Study location of the island of Stora Karlsö (indicated by *asterisk), Baltic Sea, Sweden. (b) From this site Lesser Black-backed Gull (*Larus fuscus*) inbound flights were tracked with GPS (18 flights from 9 individual gulls, coloured lines).

 V_{ref} as being the wind speed at a reference height h_{ref} (10m). By combining (2.6) and (2.7) into (2.1), we obtain:

$$\omega_{sp} \cdot h = V_{air} + V_{ref} \cdot \left(\frac{h}{h_{ref}}\right)^{\alpha}$$
(2.8)

To find the bird's steady-state flight height *h* reached during a takeoff as function of the wind profile, it requires to solve the equation f(h) = 0 with the function *f* defined as follows:

$$f(h) = V_{air} + V_{ref} \cdot \left(\frac{h}{h_{ref}}\right)^{\alpha} - \omega_{sp} \cdot h$$
(2.9)

In the variation table of the function f (Tab. S1), we observe that only one unique altitude h103 exists, enabling f(h) = 0 during an offshore takeoff manoeuvre. We can therefore conclude that 104 both the minimisation of the rate of energy consumption and regulating the ventral optic flow 105 enable a bird to fix both its groundspeed and its altitude above the sea. The bird's steady-state 106 flight height h cannot be considered as a "target flight height" or a "desired flight height", but as an 107 "optimal flight height" because the bird's altitude is adjusted as a function of the wind conditions 108 (higher under tailwinds but lower under headwinds) and thereby maximizing positive effects as 109 well as minimizing adverse effects of the wind gradient. 110

3. Gulls' trajectory recording

¹¹² 16 lesser black-backed gulls (*Larus fuscus*) were GPS tracked from their breeding colony on Stora
¹¹³ Karlsö island, Sweden (17.972° E, 57.285° N) during May to September of 2013-2015. The island
¹¹⁴ is a small offshore island (2.5 km²) located in the western central Baltic Sea, sited 7 km west of the
¹¹⁵ much larger island of Gotland (Fig. 2a). During breeding the gulls perform central-place foraging
¹¹⁶ trips [46], flying out from their island to forage, either at sea or on land [32].

Gulls were caught during late incubation (late May) using walk-in traps set over their nests. They were weighed and sexed from morphological measurements [11] or genetically [23] from a

few breast feathers taken at capture. An 18 g solar-powered UvA-BiTS GPS tracker with remote 119 download capacity [9] was mounted using either a full body or wing harnesses [64] constructed 120 of tubular TeflonTM ribbon (Bally Ribbon Mills 8476-.25") (full tagging procedure given in [32], 121 see Fig. 3). Data were downloaded and programs uploaded to the GPS devices remotely using 122 a network of four antennas providing good coverage of the colony area. GPS tracking was 123 continuous though the location intervals varied depending on the requirements of parallel studies 124 (e.g. [32]). At a 6 seconds interval on a white stork (Ciconia ciconia) on its nest, it was quantified a 125 mean altitude error of 2.77 m and a mean speed error of 0.02 m/s of the UvA-BiTS GPS tracker 126 **[9**]. 127



Figure 3. Lesser black-backed gull (*Larus fuscus*) equipped with an 18 g solar-powered UvA-BiTS GPS tracker (see [9] for GPS tracker details). Photographic credit: the authors.

The continuous GPS tracks were segmented into foraging trips and within these, sections of 128 continuous flight, with the final flight of a foraging trip considered a homing flight, as the gulls 129 returned from presumed foraging at sea (only marine trips were used in this study, c.f. [32]) to 130 the island colony. 18 takeoffs from 9 individual gulls with high resolution data were selected (i.e. 131 10 or 15-second intervals), and we selected only takeoffs reaching a steady-state altitude - i.e. 132 not those with a constantly fluctuating altitude. In addition, the final altitude had to be greater 133 than 10 m with variation in altitude during the ascent until reaching a steady-state altitude. 134 Flight GPS points were annotated with wind data extracted from a global weather model, ERA-135 interim data [13] provided by the European Centre for Midrange Weather Forecasts (ECMWF, 136 http://www.ecmwf.int/en/research/climate-reanalysis/era-interim), which gives variables at 137 3-hour intervals and is gridded with a spatial resolution of approximately 79 km. These were 138 extracted using the environmental-data automated track annotation (Env-DATA) system [14] 139 hosted by MoveBank (http://www.movebank.org/). 140

4. Full flights' dataset analysis: statistical model

¹⁴² The dataset here includes all inbound (returning to the island colony) over sea flights by the lesser ¹⁴³ black-backed gulls (383 flights, 16 gulls). The dataset is composed of median altitudes *h* calculated ¹⁴⁴ per flight, median wind speed measured at 10m-height (from ECMWF data), V_{ref} , and the gull ¹⁴⁵ identifier. After excluding the flights endowed with a median altitude below zero meters, the data ¹⁴⁶ comprise 352 observations of 16 individual gulls.

¹⁴⁷ A nonlinearity of wind profile power law (2.7) was introduced to estimate the wind speed ¹⁴⁸ $V_w(h)$ experienced by gulls at their median altitude *h* calculated per flight. A linear mixed effect ¹⁴⁹ model was designed using *lmer* in R software for the ordinates (β_i is the constant random effect)

150 as follows:

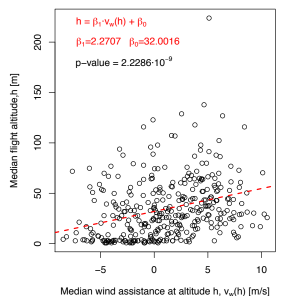


Figure 4. Gull's median altitude h versus median wind assistance V_w (head or tail wind) at median altitude h for 352 flights (16 distinct gulls). The regression line using β_1 and β_0 is plotted in red.

$$h = \beta_1 \cdot V_w(h) + \beta_0 + \beta_i + \varepsilon_{i, V_w} \tag{4.1}$$

with the regression parameters: $\beta_1 = 2.2707$ and $\beta_0 = 32.0016$. The Kenward-Roger corrected Ftest was used to calculate the significance level of the linear mixed model (ndf:1, ddf: 347.89, Fstat: 37.722, p.value: 2.2286·10⁻⁹, F.scaling: 1). The parameter β_1 was highly significant (Fig. 4). Using the coefficient $\beta_1 = 2.2707$, an identification of the ventral optic flow set-point $\omega_{sp-lmer}$ can be performed using the equation (2.8) that includes the wind profile power law as follows:

$$\omega_{sp-lmer} = \frac{1}{\beta_1} = 0.4403 \text{ rad}/s = 25.23^{\circ}/s \tag{4.2}$$

This statistical analysis tells us that gulls tend to maintain a ventral optic flow close to 25.23°/s whatever the wind conditions are while flying above the sea.

5. Takeoff time series analysis: individually tuned parametric model

¹⁶⁰ In this section, 18 takeoffs are treated as independent observations despite these being recorded ¹⁶¹ on 9 individual birds. Indeed, the weather, the wind, the state of the sea, the moment, and the ¹⁶² fishing area were uncontrolled and different from one flight to another (Fig. 2b).

(a) Parametric model estimation

The linear parametric models about each gull's elevation dynamics were estimated with the System Identification Toolbox from the Matlab software (parameters : time constant τ_h and static gain $\frac{1}{\omega_{en}}$ in (5.2). The maximum climbing speed V_{zmax} (5.1) was computed from [24, 48]:

$$V_{zmax} = \frac{2.16 \cdot m_m \cdot f}{m} - \frac{1.92 \cdot m^{\frac{2}{3}}}{\rho^{\frac{1}{2}} \cdot b^{\frac{3}{2}}}$$
(5.1)

where m_m is the mass of the flight muscles, f is the observed flapping frequency (3.26 Hz on average, see page 162 in [17]), m is the total mass including any added load, ρ is the air density (1.205 kg/m³ at 20°C) and b is the wing span. The vertical wind is low over the sea, consequently ¹⁷⁰ in flight, we neglected the vertical wind. For each of the 18 offshore takeoffs followed by a cruise

¹⁷¹ flight, we took into account the morphological parameters of each gull.

(b) Computation of the predicted altitude

The model output, i.e. the predicted altitude, hest, was computed with the Simulink environment 173 from the Matlab software. The best fit factor of the optic flow-based control model is obtained 174 by adjusting the flight muscle fraction $(\frac{m_m}{m})$ instead of the bird mass *m*, because the bird mass 175 was known without any prey load. The fit factor considered was the goodness of fit between 176 optimized simulated data (h_{est}) and actual GPS data (h_{GPS}) using a Matlab function with a 177 normalized mean square error cost function (called NRMSE cost function). NRMSE fit factor 178 varies between minus infinity (worse fit) to 1 (perfect fit). According to the table 15 in [21], the 179 flight muscle ratio ($\frac{m_m}{m}$) is relatively constant across birds species at 0.18 ± 0.05 (MEAN ± SD, 180 with n = 221). Our simulated model has been adjusted with the flight muscle ratio in order to get 181 the best fit factor, then adjusting the maximum climbing speed in the elevation dynamics model. 182 For our group of 9 individual lesser black-backed gulls, we obtained the best fit factor with a 183 corresponding distribution of flight muscle ratio $(\frac{m_m}{m})$ of 0.18 \pm 0.03, which is quite similar 184 to prediction 9 from [24]. The optic flow-based control model takes into account the observed 185 correlation between the groundspeed V_q and the altitude h coming from gulls' GPS data. The 186 proportionality factor is called here a ventral optic-flow set-point ω_{sp} (2.6). Once the best fit factor 187 has been reached by adjusting the flight muscle fraction $\frac{m_m}{m}$, each gull's altitude is re-computed 188 by considering an altitude control model that directly feeds the elevation dynamics with a "target 189 flight altitude", noted an altitude set-point h_{sp} , which is computed when the gull reached its 190 steady-state altitude. 191

(c) Optic flow-based altitude control model

¹⁹³ We consider two scales of time. The gull's forward dynamics responds faster than the gull's ¹⁹⁴ upward dynamics (constrained by V_{zmax} see (5.1)) because the height of flight arises from the ¹⁹⁵ response of a first order differential equation by considering the forward speed as a step input ¹⁹⁶ (5.2). The bird's elevation dynamics is represented in Fig. 5a, this includes both the first order ¹⁹⁷ upward dynamics (5.2) and the maximum climbing speed V_{zmax} (5.1).

$$r_h \cdot \frac{dh}{dt} + h(t) = \frac{1}{\omega_{sp}} \cdot V_g(t)$$
 (5.2)

¹⁹⁸ An explicit solution of equation (5.2) can be written, if we consider a step response at a given ¹⁹⁹ positive amplitude V_{g0} value, as follows:

$$h(t) = \frac{V_{g0}}{\omega_{sp}} \cdot \left(1 - e^{-\frac{t}{\tau_h}}\right)$$
(5.3)

For each gull trajectory, we consider only one takeoff followed by a cruise flight, and then we perform a first order system identification described by the differential equation (5.2). In this model, a proportionality factor $\frac{1}{\omega_{sp}}$ is introduced, which is the inverse of the ventral optic-flow set-point ω_{sp} (2.6), and the input of the upward dynamics (5.2) is the groundspeed V_g , which correlates the altitude h and the groundspeed V_g . If the gull's groundspeed is constant during takeoff as well as during cruising flight, then the predicted altitude profile will be the same with both models.

The inter-flight variability of the climb time constant ($\tau_h = 97.3s \pm 68.0s$, with n = 18 takeoffs) was derived on the basis of morphological properties of the birds (*inter alia* age, wingspan, body mass including the load of prey and sex).

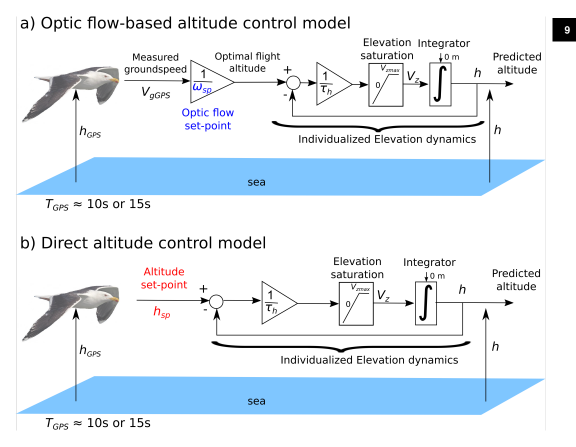


Figure 5. (a) Optic flow-based altitude control model including an individualized gull's elevation dynamics. Once the gull has reached the minimum groundspeed to takeoff, groundspeed is then relatively constant during its flight, an optic-flow-based control system can be switched on and lead the gull to a given altitude depending on both its groundspeed V_g and its ventral optic flow set-point ω_{sp} . The ventral optic-flow set-point ω_{sp} is an internal parameter used by the gull to tend asymptotically to its optimal flight altitude proportionally to its current groundspeed to its current altitude. (b) Direct altitude control model. Here, the model only includes an individualized elevation dynamics and an altitude set-point h_{sp} . This model does not impose asymptotically any proportionality between groundspeed and altitude. The altitude set-point h_{sp} is an internal parameter used by the gull to select its "desired" or "target" flight altitude.

210 (d) Direct altitude control model

Here, the bird's elevation dynamics is represented in Fig. 5b, which includes both the first order upward dynamics (5.4) and the maximum climbing speed V_{zmax} (5.1).

$$\overline{h} \cdot \frac{dh}{dt} + h(t) = h_{sp}$$
(5.4)

An explicit solution of equation (5.4) can be written, if we consider a step response at a given altitude h_{sp} value, as follows:

τ

$$h(t) = h_{sp} \cdot \left(1 - e^{-\frac{t}{\tau_h}}\right) \tag{5.5}$$

The "target flight altitude", also called the altitude set-point is denoted h_{sp} , which is computed from when the gull reached its steady-state altitude, i.e. the gull's mean altitude when $t > 3\tau_h$ or $t > 5\tau_h$, depending on data availability. In this model, there is no correlation between altitude and groundspeed.

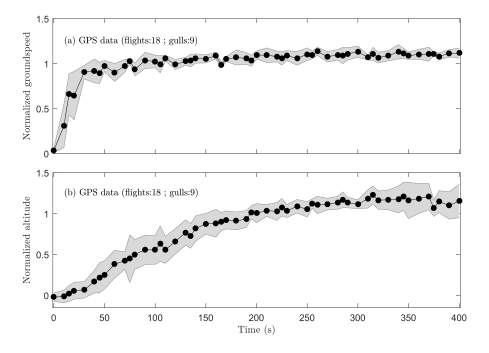


Figure 6. (a) Normalized groundspeed coming from GPS speed measurements $\frac{V_{gGPS}}{mean(V_{gGPS})}$, which is computed by the current groundspeed to average groundspeed ratio. (b) Normalized altitude coming from GPS data, which is computed by the current altitude to average altitude (by removing the first 100 seconds) ratio $\frac{h_{GPS}}{mean(h_{GPS}(100s:end))}$. Black dots represent GPS data recorded at a sampling time 10 *s* (12 trajectories) or 15 *s* (6 trajectories). Each dot represents the median value and shaded areas represent the median absolute deviation (MAD) of the GPS data collected from 18 flights.

(e) Comparison between optic flow-based and direct altitude control models

A set of 18 trajectories representing 9 different gulls are individually shown in the horizontal plane in Fig. 2b. The set of GPS data are clustered and shown in Fig. 6 for the initial 400 seconds of each flight. It allows us not only to show the increase in speed during the gulls takeoff (Fig. 6a), but also their level flight along the vertical plane (Fig. 6b). Both groundspeed and altitude have been individually normalized by the steady state value reached by the gull's groundspeed and altitude, respectively (Fig. 6). Consequently, both curves reach a steady state close to a value of one (Fig. 6).

A linear 1st order parametric model on the data (18 trajectories) gives a fit factor value 228 (i.e., a normalized mean square error cost function, called NRMSE cost function) of 40.4% 229 on average (range: 10-80%). Then, by introducing a constraint on the climb rate according to 230 prediction 10 in [24, 48], a direct altitude control model based on a non-linear 1st order parametric 231 model combined with an altitude set-point h_{sp} (see Fig. 5b for details) gives a fit factor of on 232 average 57.1% (range: 11-77%). However, by adding to the previous model a correlation between 233 groundspeed and altitude, which is linked to what we call an optic flow set-point ω_{sp} (see Fig. 234 5a for details), an optic flow-based control model gives a fit factor of 63.4% on average (range: 235 30-83%). 236

Examples comparing an optic flow-based control model to a direct altitude control model for one takeoff is given in Fig. 7b (the 17 other takeoffs are shown in Supplemental Information, Figs. S3-S19). We observe that in each case the fit factor was higher with an optic flow-based control model (blue dots in Fig. 7b rather than a direct altitude control model (red dots in Fig. 7b).

The set of normalized predicted altitudes (n = 18) computed with an altitude control model 241 (Fig. 5b) is shown in Fig. 8a, and with an optic flow-based control model (Fig. 5b) is shown 242 in Fig. 8b. Residuals, which are the errors between altitudes coming from GPS data and 243 predicted altitudes coming from models, are represented in Figs. 8c-d. We compared the residuals 244 distribution between the two models in transient response (white shaded boxes in Fig. S2) and 245 in steady-state response (gray shaded boxes in Fig. S2). The median value of the residuals (Figs. 246 8c-d) coming from the optic flow-based model was significantly higher in transient response (one-247 sided Wilcoxon rank sum test, n = 27, $p \ll 0.001$) and was also significantly higher in steady-state 248 response (one-sided Wilcoxon rank sum test, n = 27, $p \ll 0.001$). Consequently for both parts, the 249 response predicted by the optic flow-based control model was better than the response predicted 250 by the altitude control model. Finally, the average value of the residuals coming from each 251 control model in transient response, then in steady-state response, were compared to a normal 252 distribution centred around zero. The distributions of residuals with the optic flow-based control 253 model (white shaded boxes in Fig. S2) were not significantly different from a normal distribution 254 centred around zero (t-test, n = 27, p = 0.95 in transient response, and p = 0.07 in steady-state 255 response). Residuals with the direct altitude control model (gray shaded boxes in Fig. S2) were 256 significantly different from a normal distribution centred around zero (t-test, n = 27, p < 0.01 in 257 transient response and $p \ll 0.001$ in steady-state response). This statistical analysis shows that the 258 optic flow-based control model is the most established model. Besides, for 13 out of 18 flights, 259 we observe a significant correlation (Spearman's test on GPS data) between groundspeed and 260 altitude (ρ from 0.22 to 0.83, 13 flights). We therefore conclude that our optic flow-based control 261 model (Fig. 5a) better explains the gulls' GPS tracking data than the direct altitude control model 262 (Fig. 5b). 263

264 6. Discussion

(a) Comparison of optic flow set-points identified by both analyses

We compared the distribution of ventral optic flow set-points coming from the tuned parametric 266 model obtained from the takeoff time series ($\omega_{sp} = 22^{\circ}/s \pm 9^{\circ}/s$ with n = 18, Shapiro normality 267 test: p = 0.16) and the parameter $\omega_{sp-lmer} = 25.23^{\circ}/s$ obtained from the linear mixed effect 268 model (4.2), respectively. No significant difference was observed between the ω_{sp} distribution 269 and the value $\omega_{sp-lmer}$ (t-test, t:1.5296, df: 16, p-value:0.1457). This suggests shows that both 270 analyses identify optic-flow set-points that are in the same range and not significantly different. 271 As a consequence, both the takeoff time-series and the full dataset support the ventral optic flow 272 regulation hypothesis in a consistent manner. 273

(b) Effect of wind on the birds' altitude

An additional outcome of the ventral optic flow regulation hypothesis [18, 55] is that any increase in headwind will lead to a decrease in gull flight altitude in order to maintain the ventral optic flow constant (Fig. 9a). Conversely, any increase in tailwind will lead to an increase in bird altitude (Fig. 9c). A bird can adjust its ground speed by adjusting its airspeed or its heading relative to ground (and wind), thus allowing it to minimize its cost of transport in flight. The altitude control system based on optic flow is therefore consistent with previous observations on speed adjustment with respect to winds in migrating birds [2].

The small Hellman exponent α over relatively smooth surfaces, such as the sea, means that wind speed increases more rapidly than over a rough surface (e.g. a forest). Thus at higher altitudes (i.e., from 10 m to 100 m) wind speed will not vary much, but below 10 m wind speed can double going from 1 m to 10 m. Around the sea's waves wind is deflected leading to a pattern of updrafts and downdrafts [50, 53, 68]. Together these effects are used by soaring seabirds in dynamic soaring, gust soaring or "sweeping flight" [50, 53, 68], and the characteristic meandering flight style that results has been termed "wave-meandering wing-sailing" [61]. Flapping seabirds

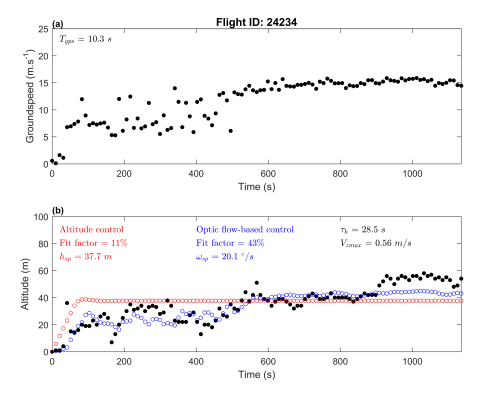


Figure 7. (a) Groundspeed of the gull ID24234 tracked with the GPS. (b) Altitude of the gull: black dots represent the GPS data, red dots represent the gull altitude on the basis of an altitude-based control model (fit factor: 11%), and the blue dots represent the gull altitude on the basis of an optic flow-based control model (fit factor: 43%). A significant correlation was observed between groundspeed and altitude of the GPS data ($\rho = 0.83$, $p \ll 0.001$ by Spearman's test).

can also use these features to gain a higher climb rate at the start of a take-off maneuver, taking off facing into the wind in the updraft formed by the deflection of the wind over a wave (see page 268 in [51], and [33]), which therefore reduces the effort required to take-off and accelerate to reach the maximum range speed V_{mr} . Seabirds may also use the "ground effect" while flying very close to the sea to reduce their energetic expenditure [8], which is helpful for takeoff at sea.

(c) Effect of altitude on optic flow

According to prediction 3 in [24, 48], the optimal altitude for a migratory bird is that where it can 295 get just sufficient oxygen to maintain its cruising airspeed. This arises from the power required to 296 fly at maximum range speed decreasing with altitude due to decreasing air density. Consequently, 297 at an altitude of 6000 m, where the air density is half that at sea level, a bird should theoretically 298 fly $\sqrt{2}$ times faster. On the other hand, at a given optic-flow set-point working in a 100 m altitude 299 range, the optic flow would be divided by a factor $\frac{60}{\sqrt{2}}$ at an altitude of 6000 m. Therefore the optic flow would be too small to be maintained at the amplitude of the one generated in a 100 m altitude 300 301 range. Recently, McLaren and colleagues (2016) analysing flights of lesser black-backed gulls 302 flying between south-east England and The Netherlands recorded much greater flight altitudes 303 than those observed here during homing flights to the breeding colony, with maximal values 1,240 304 m [42], even though typical values were lower at 100-150 m. On migratory flights, the gulls have 305 been recorded flying higher still, though that is overland, with maximal altitudes around 5,000 306 m (unpublished data). Consequently, an optic flow based altitude control system can only work 307

below a 100-meter altitude range where the optic flow is significant and detectable by the visual
 system of the birds.

(d) Are groundspeed and altitude still proportional at higher altitudes?

Birds making lower altitude flights (<100-150 m) will generate a detectable optic flow. However, 311 when on long distance or migratory flights birds may fly higher at hundreds to thousands of 312 meters (see above), optic flow values will then be extremely low, thus unlikely to be suitable for 313 regulating a given optic flow set-point. This relates to the finding for common swifts (Apus apus) 314 by Hedenström & Åkesson (2017), that the swifts did not compensate for head- and tail winds as 315 expected from flight mechanical theory when flying at high altitudes (>1000 m), but they did so at 316 low altitudes (<100 m) [25]. This was interpreted as a failure to detect small changes in optic flow 317 due to winds by the swifts' visual system at high altitudes. In addition, for altitudes higher than 318 400 m, lesser black-backed gulls were observed to compensate less for cross-wind disturbance 319 than they did at lower altitudes: fractional compensations were observed to decrease from about 320 1.3 (on average) to less than 0.5 at 900 m height [42]. At altitudes above 400 m, gulls' groundspeed 321 may be highly dependent on the wind speed: no altitude increase or decrease can be predicted 322 with respect to the optic flow-based control model as optic flow is low thus its changes with 323 altitude would be difficult to detect by the gulls' visual system. 324

(e) Bird navigation in the vertical dimension: can birds use barometric pressure to determine altitude?

The birds' mechanoreceptive paratympanic organ (PTO) is located in the middle ear, and it is probably used by birds to detect barometric pressure [65]. Birds appear to use the PTO not only as a barometer to predict the onset of inclement weather [10, 60, 65], but also as a genuine altimeter to adjust their flight altitude during migration. Birds can fly level within ± 20 m for distances of 2–3km at altitudes of 700 – 1,100m, even at night [22], i.e. without visual cues. However, it is still an open question whether birds can use changing barometric pressure directly to measure their current altitude in real time.

A mechanoreceptive scale sensory organ found in fish [5] may play the same sensory function 334 as the PTO in birds. It is known that fish can determine their depth using hydrostatic pressure 335 [29, 63]. On this point, it was demonstrated that the dynamic depth sensing in fish is less than 1 336 m at a depth of 100 m [63]. However, water density is approximately 1,000 times higher than air 337 density, and the pressure gradient in flight is therefore particularly low generating extremely low 338 339 frequencies in the feedback signal to the bird's elevation dynamics. Therefore, it would be difficult to adjust the flight altitude for a short period of time, only being practical for long periods of time 340 such as for example during longer distance migratory flights. 341

(f) Effect of wave motion on the optic flow pattern

The flight model assumes that the sea-surface, over which the gulls fly, provides a stationary 343 reference frame: no data are currently available on the wave speed. Therefore, the optic flow 344 experienced by the gulls is solely modeled as a function of their own movement (groundspeed 345 and altitude). Previous studies on bird navigation over water suggest that the seascape (or more 346 specifically the wavescape) is not a fixed reference frame [1], as the wave patterns move, usually 347 in roughly the same direction as the wind but at a slower speed. Therefore the perceived optic 348 flow will be different than the physical optic flow. Alerstam & Petterson (1976) suggested that the 349 motion of the wave scape allows birds to only partially compensate for wind-drift over the sea 350 351 [3], thus presumably a similar constraint may apply to using the ventral optic flow for control of flight altitude. 352

Overall, the wave pattern will reduce the adjustment of altitude if a fixed optic flow set-point was used, as under headwinds perceived optic flow will be higher than otherwise, i.e. even

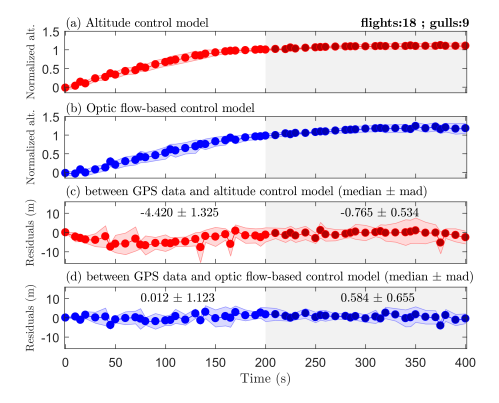


Figure 8. Red dots (altitude control model) or blue dots (optic flow-based control model) represent predicted altitude ((a) and (b)) or residuals ((c) and (d)) at a same sampling time 10 *s* (12 trajectories) or 15 *s* (6 trajectories) like GPS data (see Fig. 6). Each dot represents the median value and shaded areas represent the median absolute deviation (MAD) of data (n = 18). The white shaded areas represent the transient response (time <200 *s*) during takeoff and ascent, and the gray shaded areas represent the steady state response (time >200 *s*) once in cruising flight. The duration 200 $s \approx 2 \cdot \tau_h$ represents about 86% of the step response of a 1st order dynamic system (see (5.3)). (a) Normalized predicted altitude using an **altitude control model** (Fig. 5b), which is computed by current predicted altitude average predicted altitude (by removing the first 100 seconds) ratio $\frac{h}{mean(h(100s:end))}$. (b) Normalized predicted altitude (by removing the first 100 seconds) ratio $\frac{h}{mean(h(100s:end))}$. (c) Residuals between GPS data (Fig. 6b) and altitude computed with the optic-flow based control model (data in (a)). (d) Residuals between GPS data (Fig. 6b) and altitude computed with the optic-flow based control model (data in (b)).

³⁵⁵ as groundspeed approaches zero there will still be a perceived optic flow if the wavescape is ³⁵⁶ moving, which would lead to higher flight altitudes than expected. While under tailwinds optic ³⁵⁷ flow is somewhat reduced, as the sea surface pattern will be moving in the same direction as the ³⁵⁸ bird, and hence lower than expected flight altitudes would result. The wave pattern distorts the ³⁵⁹ ventral optic flow perceived: such disturbances could be added to the flight model once data or a ³⁶⁰ methodology of how to obtain wave pattern becomes available.

However, for optic flow to be useful ripples above the sea are essential to form a textured surface. In fact, it was observed by Heran & Lindauer (1963) that a great number of honeybees plunged into the water when the water surface was mirror smooth [28]. An altitude control system based solely on a ventral optic flow regulation irrevocably pulls any flying animal down whenever its eye fails to measure an optic flow [18]. This did not happen in honeybees when the water surface was rippled [28, 62] or when a floating bridge provided a visual contrast [28].

At this level of reasoning, we may wonder if the visual pattern produced by waves was textured enough during the gulls' flights for an optic flow field to be perceived. To investigate

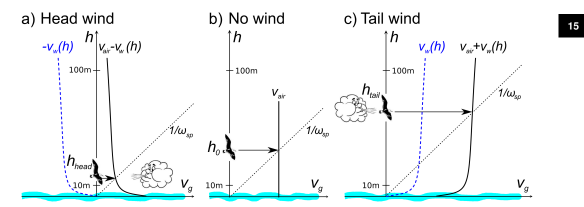


Figure 9. Gull's speed and altitude for three different wind scenarios under the hypothesis that the gull adjusts its vertical lift to maintain constant its ventral optic flow. The red straight line $\frac{1}{\omega_{sp}}$ indicates the set of possible pairs of altitudes and groundspeeds allowed by the ventral optic flow regulation hypothesis. (a) In the presence of a head wind, given that the wind speed increases with the altitude, the groundspeed profile $V_{air} - V_w(h)$ intersects the straight line $\frac{1}{\omega_{sp}}$ at a lower altitude h_{head} than in absence of wind. (b) In the absence of wind, the ground speed and hence the altitude depend only on the airspeed produced by the agent: the vertical line $V_g = V_{air}$ intersects the line $\frac{1}{\omega_{sp}}$ at the altitude h_{0} . (c) In the presence of a tail wind, the ground speed profile $V_{air} + V_w(h)$ intersects the straight line $\frac{1}{\omega_{sp}}$ at a greater altitude h_{tail} than in absence of wind. Modified from [55].

this, knowing that the average significant wave height of the Baltic Sea in 1991-2015 was in the 369 range 0.44–1.94 m [34], which corresponds to a Beaufort number of 3 (gentle breeze, mean wind 370 speed equivalent from 3.4 m/s to 5.4 m/s) to 4 (moderate breeze, mean wind speed equivalent 371 from 5.5 m/s to 7.9 m/s) [6]. We deduce that gulls could see scattered or fairly frequent white-372 crested waves at an effective height of 10 m above the sea level. However for Beaufort numbers 373 from 0 to 2, the sea has a smooth appearance, which makes for poor visual conditions to perceive 374 an optic flow field. Interestingly, the wind conditions corresponding to a Beaufort number from 3 375 to 4 fit not only with the wind conditions of gulls in flight (Fig. S20), but also with their altitude 376 (see page 166 in [17]). We can conclude that wind is an important parameter to generate an optic 377 flow field cue, and to help gulls to control their flight above the sea. 378

Little is known about the visual system of gulls. The spatial acuity of seabirds can be more 379 than four times lower than that in humans [43], with a maximum spatial acuity of about 60 380 cycles/degree in humans. Moreover, in seabirds rods are evenly distributed across the entire 381 retina [15], which allow them to conveniently detect the optic flow coming from the sea. Most 382 of the seabirds have a maximum binocular field width in the $15^{\circ} - 30^{\circ}$ range (about 120° in 383 humans), which is limited, suggesting that binocular vision plays only a minor role in seabirds' 384 flight control system [39]. We conclude that the optic flow field is the major visual cue used by 385 seabirds to control their flight above the sea. 386

(g) Optic-flow set-point: differences between honeybees and gulls

There are a number of differences in flight behaviours expressed by birds and flying insects 388 [4]. Typically, the average maximum airspeed of honeybees is approximately 7.5 m/s with a 389 minimum power speed of their power U-curve at 3.3 m/s [44]. In free-flight natural conditions, 390 honeybees have been observed to fly from 3.3 m/s to 5.1 m/s [44]. However, lesser black-backed 391 392 gulls typically fly at an airspeed in natural offshore conditions at an average 12.3 m/s \pm 2 m/s (see [17], page 166) with a minimum power speed of their power U-curve at 9.3 m/s (computed 393 for lesser black-backed gull, see [27]). Hence, lesser black-backed gulls can fly 3 times faster than 394 honeybees by comparing their minimum power speed. 395

In honeybees, average maximal flight height is about 2.5 m over natural terrain [16, 28]. In general, lesser black-backed gulls fly at an altitude over sea of up to 130 m with a distribution of 31 m \pm 29 m on average (see [17], pages 166-167) during foraging flights. We conclude that lesser black-backed gulls fly much higher than honeybees during foraging flights, which reduces optic flow emanating from the sea.

Consequently, we can conclude from these two last points that the ventral optic-flow set-point 401 of lesser black-backed gulls is much lower than that typically experienced by honeybees, knowing 402 that the ventral optic-flow set-point of honeybees is close to $200^{\circ}/s$. Our statistical analysis 403 estimates that the ventral optic-flow set-point of lesser black-backed gulls is close to $25^{\circ}/s$ on 404 average (see section 4), which is a detectable value by the gulls' visual system [39, 40, 41]. A recent 405 review indicates that pigeons' fast LM neurons (pretectal nucleus lentiformis mesencephali) 406 respond to optic flow stimuli of their preferred backward direction (front to back visual stimuli: 407 temporal to nasal on the retina) in this same angular velocities range [69]. 408

409 7. Conclusion

A mathematical model of optic flow-based offshore takeoff control system in lesser black-backed 410 gulls was developed in this study to understand what visual cue can be used by seabirds to 411 control their takeoff and to cruise over a sea surface. This mathematical model introduced an 412 optic flow set-point parameter, which aims to be maintained constant by seabirds during take-413 off manoeuvers and cruising foraging flights. Besides, the model takes into account the bird's 414 individual morphology through its elevation dynamics. Finally, both analyses on the takeoff time-415 series and the full dataset support the ventral optic flow regulation hypothesis in a consistent 416 manner. 417

We conclude that the optic-flow regulation principle allows seabirds to control their altitude 418 over sea at low flight altitudes without having to measure their current altitude directly by 419 another method. To do this, they just have to measure the optic flow perceived from the sea 420 to adjust their vertical thrust in order to maintain the ventral optic flow at a given value, called 421 the optic-flow set-point, as previously suggested for flying insects [18, 55]. According to both the 422 airspeed and altitude ranges of lesser black-backed gulls measured during flight in their natural 423 environment, we demonstrate that gulls could control their altitude by regulating the ventral 424 optic-flow at a value of $25^{\circ}/s$ on average, allowing them to fly jointly up to 130 m in altitude at 425 a groundspeed up to 20 m/s, while maintaining visual contact with the sea. The introduction of 426 this asymptotic proportionality relationship for birds also accounts very nicely for the transient 427 altitude response during takeoff. Overall, gulls need such accurate altitude control based on optic 428 flow to optimize their energetic effort irrespectively of favorable or unfavorable unknown wind 429 430 conditions while being robust to ground disturbances such as relief.

431 Authors' contributions

JRS, FR, TJE and AH developed the modelling; JRS ran the models on Matlab software; TJE and SÅ tagged the gulls and collected the data; JRS, TJE and FR analysed the modelling results; JSB provided the tracking system; FR and AH supervised the collaboration; FR drew figs 1 and 9; TJE drew figs 2 and 3; JRS drew figs 5, 6, 7, 8 and S1-S19; TJE, JRS and FR drew fig 4; JRS wrote the first draft of the paper; all authors prepared and revised the manuscript.

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