Contents

1	Abstract	1			
2	Introduction	ntroduction 1			
3	Model 3.1 Modeling interactions and group-level properties 3.2 Individual-level fitness 3.3 Optimization methods	3			
4	Results	4 4 4 5 5			
5	Discussion	5			
6	Appendix 6.1 Linear algebra background	7 8			
7	Figures	13			

1 Abstract

Birds in flocks, like animals in many types of social groups, can use their peers to learn about the environment and update their opinions about where to move in that environment. Previous work on starlings has measured the number of other birds an individuals pays attention to and shown that this number of "neighbors" leads to a social network that is conducive to the whole flock reaching consensus. It is unclear, however, why an individual would change its behavior, i.e. the number of neighbors it learns from, to achieve this outcome. Individual birds should optimize how well they learn about the environment. There are (at least) two features of the environment that are important to an individual bird's fitness—the location of a predator and the location of resources—and it can try to learn about these from its peers. In this work, we identify the optimal strategies for birds under selection pressure dictated by these types of information and explore the relationship between the robustness measure and the correlation length of the flock, a measure commonly used to describe how close a flock's structure is to criticality.

2 Introduction

Questions:

- 1. How are the two group properties— \mathcal{H}_2 robustness and correlation length—related?
- 2. What are the equilibrium strategies of individuals seeking to optimize their own fitness?
- 3. What group properties emerge from individual-level optimization?

3 Model

3.1 Modeling interactions and group-level properties There are two ways to describe equilibrium properties of a flock sharing information over a network. The first is to write down a set of differential equations with noise to describe how each bird's opinion changes over time and then to analyze the distribution of these opinions after the dynamics have reached equilibrium [9, 8]. The second is to write down an explicitly probability distribution of the opinions at equilibrium and to analyze this distribution. With either approach, we consider a vector of opinions v where v_i a a scalar that represents the ith's bird opinion about an unspecified behavior, for example where or how fast to fly. Both approaches rely on a network of connections between the birds, which we will describe first. Let v be the adjacency matrix such that

$$A_{ij} = \begin{cases} 1 & \text{if } i \text{ pays attention to } j \\ 0 & \text{otherwise} \end{cases}$$
 (1)

If n_i is the number of neighbors node i has, then $n_i = \sum_k A_{ik}$. We assume that having more neighbors means paying less attention to each so we normalize A to find \bar{A} : $\bar{A}_{ij} = \frac{A_{ij}}{n_i}$.

The Laplacian of this matrix is given by $L = \bar{A} - I$, which is symmetric only if A is. We can symmetrize the interactions by defining $S = \frac{1}{2}(A + A^T)$ and P such that $P_{ij} = S_{ij}$ for $j \neq i$ and $P_{ii} = -\sum_k S_{ik}$. Then L and P share the properties that $L\vec{1} = P\vec{1} = \vec{0}$ and that each is negative semi-definite, with the difference being that P is necessarily symmetric. It is more common to use L in a dynamics system approach, so that

$$\dot{v} = Lv + D\xi \tag{2}$$

where D is a diagonal matrix with entires $\{\sqrt{n_i}\}$ and $E[\xi(t)] = \vec{0}$, $E[\xi(t)\xi(s)^T] = \frac{1}{2}\delta(t-s)I$, and $E[v(0)\xi(t)^T] = \vec{0}$. This describes a system in which each bird changes averages the differences between its own opinions and those of its neighbors and changes its opinion to minimize these differences. Our choice of D assumes that birds with more neighbors make noisier estimates of how to change their opinions.

The Boltzmann distribution in statistical physics is a standard way in which to write an equilibrium distribution. In physics, those states with the highest energy are least likely. In our system, those sets of opinions that are least in consensus will be less likely. Specifically, we will use the following expression to represent the "energy" of an opinion vector

$$H(v) = -v^T P v (3)$$

and the Boltzmann distribution

$$P(v) = \frac{1}{Z} \exp\left(-\frac{1}{2}H(v)\right) \tag{4}$$

where $Z = \int_{\mathbb{R}}^{N} \exp\left(-\frac{1}{2}H(v)\right) dv$.

With the dynamical approach, in the absence of noise, the system would reach consensus with each $v_i = \alpha$ for some $\alpha \in \mathbb{R}$. Similarly, with the statistical physics approach, consensus states with each $v_i = \alpha$ are most likely. However, all consensus values α are equally likely. It is therefore difficult to describe correlations between the bird's opinions at equilibrium. It becomes easier, however, to consider deviations from consensus. If all v_i are equal, then $v = \alpha \vec{1}$. To consider deviations from this consensus state, it is useful to define a matrix $Q \in \mathbb{R}^{N-1} \times \mathbb{R}^N$ such that each row of Q is orthogonal to $\vec{1}$ and the rows are pairwise orthogonal. Young et al. show in [9, 8] that this is equivalent to a Q such that $QQ^T = I_{N-1}$ and $Q^TQ = I_N - \frac{1}{N}\vec{1}\vec{1}^T$. To simply consider absolute deviations from the average opinion of the flock, we can define $z = v - \langle v \rangle \vec{1} = Q^TQv$, where $\langle v \rangle = \frac{v_i}{N}$. To consider the component of the opinion vector that is orthogonal to consensus, we can define y = Qv = Qz. Let $\Sigma_z = \lim_{t \to \infty} E[zz^T]$ and $\Sigma_y = \lim_{t \to \infty} E[yy^T]$ using the dynamical approach and $C_z = E[zz^T]$ and $C_y = E[yy^T]$ using the statistical physics approach. Since y = Qz and $z = Q^Ty$, in either case, we can translate easily between Σ_z and Σ_y or between C_z and C_y . In the Appendix, we derive equations for each of these covariance matrices and summarize the expression for the y covariance matrices in Table 1. The first thing to note

is that, if instead of using L we use P in the dynamical approach, then $C_y = 2\Sigma_y$, as shown in Claim XXX. There is therefore a close correspondence between the two approaches.

These covariance matrices allow us to define two group properties. The first is \mathcal{H}_2 robustness, as used in [9, 8]. This is a measure of the robustness of the consensus state in which all birds have the same opinion to noise at equilibrium. Specifically,

$$\rho = \frac{1}{\lim_{t \to \infty} E[||y||]} = \frac{1}{\lim_{t \to \infty} \text{Tr}(\Sigma_y)}.$$
 (5)

The second property is the correlation length of the flock. This is the distance over which birds' opinions tend to be positively correlated with each other. If we know the distances between the birds and the covariance matrix C_z , we can plot the correlations $\{E[z_iz_j]\}$ between each pair of birds' deviations from consensus against the distance between the birds and then identify the correlation length λ by finding the distance at which these correlations become 0.

So far, we have considered birds whose opinions only depend on the opinions of the rest of the flock. We can additionally incorporate environmental information by assuming that there is an environmental signal such that more informed birds have opinions closer to 1. If we let B be a matrix with $B_i i = 1$ if i can perceive the signal and 0 otherwise, then the dynamical system becomes

$$\dot{v} = Lv + B(\vec{1} - v) + \xi = L_s v + B\vec{1} + n_i \xi \tag{6}$$

where $L_s = L - B$. Similarly, the statistical physics model becomes

$$P(v) = \frac{1}{Z} \exp\left(-\frac{1}{2}(v-\vec{1})^T(-P+B)(v-\vec{1})\right) = \frac{1}{Z} \exp\left(\frac{1}{2}(v-\vec{1})^T(-P_s)(v-\vec{1})\right)$$
(7)

where $P_s = P - B$. Again, we can find the covariance matrices Σ_z , Σ_y , C_z , and C_y . As was the case without a signal, if we use P in the dynamical approach, then we find that $C_y = 2\Sigma_y$. We define ρ and λ as above using the matrices Σ_y and C_z in the presence of a signal.

3.2 Individual-level fitness To find the optimal strategies from the individuals' perspectives, we need a measure of individual-level fitness. We use the dynamical system to describe how well each bird learns about the environmental signal. For a given matrix L, the expected opinion q_i of each individual satisfies the equations

$$\dot{q} = L_{\rm s}q + B\vec{1}.\tag{8}$$

To generate a matrix L, we distribute the individuals randomly in space (specifically, in a unit square). has a strategy n_i that dictates how many neighbors it has, so that $n_i = \sum_{j \neq i} \mathbf{I}(A_{ij} \neq 0)$. Each individual then pays attention to its n_i nearest neighbors. One individual is chosen to be a receiver; it and all those individuals within a radius r of the receiver perceive, so that $B_{ii} = 1$ for all those birds within this radius. We do this 1000 times and find q(1) for each matrix, which we refer to as a signaling event. This is meant to simulate a situation in which such that information is present only intermittently and otherwise the birds are moving randomly through space and restructuring their network.

We consider signals that provide information about two types of content: predators and food. If the signal is information that a predator is present, we assume that whichever individual is least aware of the signal will be predated, i.e. the individual i such that q_i is lowest. Then, fitness is proportional to the probability of surviving, i.e. 1 minus the probability of being the least informed over many signaling events. If the signal is information that resources are available, we assume that whichever individual is most aware of the signal will get access to the resource, i.e. the individual i such that q_i is highest. In this case, fitness is proportional to the probability of getting the resource, i.e. the probability of being the most informed over many signaling events. The behavior that the birds can change to improve their fitness is the number of other birds they pay attention to, n_i . In particular, each individual has a strategy n_i that dictates how many neighbors it has, so that $n_i = \sum_{j \neq i} \mathbf{I}(A_{ij} \neq 0)$.

3.3 Optimization methods To understand what strategies we might expect to find, we are interested in identifying the optimal strategies. We does this in two ways. First, using the framework of adaptive dynamics in finite populations, we identify the evolutionarily stable strategy. This framework assumes that there is a homogeneous population into which a mutant individual tries to invade. If strategies change on a learning rather than an evolutionary timescale, or if there is just a lot of variation in the population, we might expect individuals to try to optimize their strategies in the context of a heterogeneous population. This is our second method of optimization. Given a random set of initial strategies over the group, we allow each bird to choose the strategy that would be best given the rest of the strategies being used and repeat this process until the birds reach an equilibrium set of strategies. We repeat this over many initial sets of strategies to find average properties of this optimization process.

4 Results

- **4.1** Robustness and correlation length are positively related. In a homogeneous flock, where every bird is using the same strategy, as both the strategy and the radius of the signal change, both \mathcal{H}_2 and the correlation length are affected. As the Figure 4 shows that there is a relationship between two group properties: the \mathcal{H}_2 norm and the correlation length of the flock.
- 4.2 There are multiple ESS, which depend on the content of the signal. Surprisingly, even without imposing costs on paying attention to more neighbors, paying attention to the whole flock is not usually the best strategy. The ESS strategy of the number of neighbors to pay attention strongly depends on which selection pressure is being applied. If a focal bird pays attention to many others, its opinion will be similar to theirs. This will prevent it from being the most informed and hence the first to locate resources. Thus, the uninvadable strategy is to pay attention to few neighbors when selection is due to knowledge about resources. On the other hand, having many neighbors will prevent the focal bird from being the least informed and thus the most likely to be eaten. Thus, the uninvadable strategy is to pay attention to many neighbors when selection is due to knowledge about predators. Figure 1a shows the ESS strategy when selection is due to knowledge about resources is always lower than the ESS strategy when selection is due to knowledge about predators (Figure 1a).

The ESS strategies when selection is due to knowledge about predators show two interesting features. First, there are many parameters for which there are multiple ESS strategies. It might seem that if a strategy outperforms all others when it is the resident strategy, it should also outperform other strategies as an invader. However, when there are multiple ESS strategies, if we take any two of those ESS strategies, each is uninvadable by the other. This indicates how important the social environment set by the resident strategy is. The pairwise invisibility plot in Figure 1b shows that low strategies are always costly and that high strategies are only costly when the resident strategy is high.

If a bird has the environmental signal, it will not be predated, regardless of the number of neighbors it has. The differences in performance only appear when comparing two birds, neither of whom have the signal. To understand the costs of having too few or too many neighbors, we compare the performance of two focal birds, a resident and an invader, neither of whom has the signal. There are two main cases: there can be a single informed bird or multiple informed birds. We further break down the first case depending on which of these two focal birds pays attention to the informed bird: neither, the resident, the invader, or both. For each of these cases, we find the probability that the invader is predated. An invader with one fewer neighbor than the resident always has probability of being predated greater than chance (1/N) because the resident often has an informed neighbor while the invader how no or fewer informed neighbors and thus a less informed opinion (Figure 1c). The total probability of being predated stays more or less constant regardless of the resident strategy, with a slight decrease for high resident strategies (Figure 1b and c). An invader with one more neighbor than the resident is only more likely than chance to be predated when the resident strategy is high enough (Figure 1d). This invader performs most poorly when both it and the focal resident pay attention to the informed bird(s). In this case, an invader with one more neighbor has more naive neighbors and thus a less informed opinion. This is increasingly problematic as the resident strategy increases and both the focal birds are likely

to pay attention to the informed bird(s).

The predation ESS is to pay attention to more neighbors when the signal is more public. These opposing forces caused by the costs of not paying attention to informed neighbors and the costs of paying attention to more uninformed neighbors also explains the second feature of the ESS strategies: they are non-monotonic with respect to the the radius of the signal, i.e. how public the signal is (Figure 1a). Figure 1a shows that there are essentially three categories of radii: When the radius is low, the signal is private, with only a few birds having true information. When the radius is intermediate, the signal is public, with many birds having true information. When the radius is high, the signal is universal, with essentially all birds in the flock having true information. When the signal is private, paying attention to too few birds is costly because with more neighbors it is more likely to have a truly informed neighbor, but paying attention to too many uninformed birds can also be costly. This leads to a high but not maximal ESS strategy. When the signal is public, the costs of paying attention to uninformed birds are reduced and the ESS strategy increases to become as high as possible. When the signal is universal, not paying attention to informed birds does not impose a cost. Additionally, in a group in which the residents pay attention to all of their neighbors, any mutant strategy can get just as much information by paying attention to fewer neighbors so the maximal strategy is no longer an ESS (Figure 1a).

Figure 2 shows how the ESS strategy changes as a function of the \mathcal{H}_2 robustness.

4.4 Learned strategies The equilibrium strategies when the individuals are allowed to learn are similar to the ESS strategies: learned strategies when selection is due to predation are always higher than learned strategies when selection is due to resources (Figure 3). Surprisingly, even when the group starts with heterogeneous strategies, they often reach a homogeneous equilibrium. Under selection due to predation, the \mathscr{H}_2 robustness and correlation length of the flock tend to decrease, whereas under selection due to resources, \mathscr{H}_2 robustness is relatively constant and correlation length tends to increase (Figure 3).

5 Discussion

explore vs. exploit risk aversion vs. risk tolerance

expect single ESS strategies but social information depends on the social environment set by the resident strategy expect ESS strategies to decrease as the signal becomes more public, but we find the opposite

6 Appendix

6.1 Linear algebra background

Theorem 1 Given any positive symmetric matrix Q > 0 there exists a unique positive symmetric matrix $\Sigma > 0$ satisfying $A\Sigma + \Sigma A^T = Q$ if and only if the system $\dot{x} = Ax$ is globally asymptotically stable.

Claim 2 Consider the dynamical system,

$$\dot{y} = Ay + v + Q\xi \tag{9}$$

with $E[\xi(t)] = 0$, $E[\xi(t)\xi(s)^T] = \frac{1}{2}\delta(t-s)I$, and $E[y(0)\xi^T(t)] = 0$. If all eigenvalues of A are negative and if $\Sigma_t = E[y(t)y(t)^T]$ and $\Sigma = \lim_{t \to \infty} \Sigma_t$, then Σ satisfies the Lyapunov equation

$$A\Sigma + \Sigma A^{T} = A^{-1}vv^{T} + vv^{T}(A^{-1})^{T} - QQ^{T}.$$

Further, if $C_{ij} = \lim_{t\to\infty} E[(y_i(t) - E(y_i(t)))(y_j(t) - E(y_j(t)))]$ then

$$AC + CA^T = -QQ^T$$

Proof.

$$\begin{split} & \Sigma_t = E[y(t)y(t)^T] \\ \Rightarrow & \dot{\Sigma} = E[\dot{y}(t)y(t)^T + y(t)\dot{y}(t)^T] \\ & = E[(Ay + v + Q\xi)y^T + y(y^TA^T + v^T + \xi^TQ^T)] \\ & = AE[yy^T] + E[yy^T]A^T + vE[y^T] + E[y]v^T + QE[\xi y^T] + E[y\xi^T]Q^T \\ & = A\Sigma_t + \Sigma_t A^T + vE[y^T] + E[y]v^T + QE[\xi y^T] + E[y\xi^T]Q^T. \end{split}$$

Note that we can write a solution to (9) as

$$y(t) = \exp(At)y(0) + \int_0^t \exp(A(t-s))vds + \int_0^t \exp(A(t-s))Q\xi(s)ds.$$
 (10)

Eq. (10) allows us to write

$$\begin{split} E[y(t)] &= \exp(At)y(0) + \int_0^t \exp(A(t-s))v ds \\ &= \exp(At)y(0) - A^{-1} \exp(A(t-s))v|_{s=0}^{s=t} \text{ which exists because } A \text{ has no 0 eigenvalues} \\ &= \exp(At)y(0) - A^{-1}(I - \exp(At))v. \end{split}$$

Note that if we take the limit as t goes to infinity,

$$\lim_{t \to \infty} E[(y(t))] = \lim_{t \to \infty} \exp(At)y(0) - A^{-1}(I - \exp(At))v = -A^{-1}v$$

since all eigenvalues of A are negative. Eq. (10) also allows us to write

$$\begin{split} E[y\xi^T] &= \exp(At)y(0)E[\xi(t)^T] + \int_0^t \exp(A(t-s))BE[\xi(t)^T]ds + \int_0^t \exp(A(t-s))QE[\xi(s)\xi(t)]ds \\ &= 0 + 0 + \frac{1}{2}Q \\ \Rightarrow \dot{\Sigma} &= A\Sigma_t + \Sigma_t A^T + vE[y(t)^T] + E[y(t)]v^T + \frac{1}{2}QQ^T + \frac{1}{2}QQ^T \\ &= A\Sigma_t + \Sigma_t A^T + vE[y(t)^T] + E[y(t)]v^T + QQ^T \\ \Rightarrow \lim_{t \to \infty} \dot{\Sigma} &= A\Sigma + \Sigma A - vv^T(A^{-1})^T - A^{-1}vv^T + QQ^T \\ \Rightarrow A\Sigma + \Sigma A &= vv^T(A^{-1})^T + A^{-1}vv^T - QQ^T \text{ since } \dot{\Sigma} &= 0 \text{ at equilibrium.} \end{split}$$

Further, by definition of C,

$$\begin{split} C &= \Sigma - \lim_{t \to \infty} E[y(t)] E[y(t)^T] \\ &= \Sigma - (-A^{-1}v)(-A^{-1}v)^T \\ &= \Sigma - A^{-1}vv^T(A^{-1})^T \\ \Rightarrow AC + CA^T &= A\Sigma + \Sigma A^T - AA^{-1}vv^T(A^{-1})^T - A^{-1}vv^T(A^{-1})^T A^T \\ &= A\Sigma + \Sigma A^T - vv^T(A^{-1})^T - A^{-1}vv^T \\ &= -QQ^T. \end{split}$$

We define a matrix $Q \in \mathbb{R}^{N-1} \times \mathbb{R}^N$ such that each row of Q is orthogonal to $\vec{1}$ and the rows are pairwise orthogonal. Young et al. show in [9] show that this equivalent to finding Q such that $Q\vec{1} = \vec{0}$, $QQ^T = I_{N-1}$, and $Q^TQ = I_N - \frac{1}{N}\vec{1}_N\vec{1}_N^T$. Since the rows of R are orthonormal, $RR^T = I$. Further, since the rows of R are orthonormal, R is invertible, which implies that $R^T = R^{-1}$. Then we can write Again, we will use a rotation matrix $R \in \mathbb{R}^N \times \mathbb{R}^N$ such that the last row of R is $\frac{1}{\sqrt{N}}\vec{1}$ and the

other rows are normal and orthogonal to $\vec{1}$. Above, we found a $Q \in \mathbb{R}^{N-1} \times \mathbb{R}^N$ such that each row of Q was orthogonal to $\vec{1}$ and $R \in \mathbb{R}^N \times \mathbb{R}^N$ be the matrix with $\frac{1}{\sqrt{N}}\vec{1}^T$ as its last row and the first N-1 rows identical to the rows of Q. Then R is a rotation matrix, so $RR^T = R^TR = I$ and $\det(R) = 1$. This allows us to

$$\bar{y} = Rx, \ y = Qx = \bar{y}_{i=1}^{N-1} \ (\text{or} \ y = RQ^TQx), \ z = Q^TQx, \ V = \frac{1}{N} \sum_i x_i$$

6.2 Opinion dynamics with noise In our model, each bird in the flock as an opinion x_i , giving an opinion vector x. The \mathcal{H}_2 robustness of the opinion dynamics is defined by the inverse of the expected distance at equilibrium from the opinion vector to a consensus state in which all birds have the same opinion, i.e. $x_i = \alpha$ for all i and for some $\alpha \in \mathbb{R}$. That is, the dynamics are robust if, at equilibrium, the opinions are close to being in consensus. We can find this distance by rotating the opinion vector into the space orthogonal to the consensus vector, $\vec{1}$. If y = Qx, then $||y|| = y^T y$ describes how far away from consensus x is. Specifically, robustness as in [9] is given by

$$r = \lim_{t \to \infty} \frac{1}{E[||y||]} = \lim_{t \to \infty} \frac{1}{\operatorname{Tr} E[yy^T]}.$$
 (11)

We also define a full rotation matrix $R \in \mathbb{R}^N \times \mathbb{R}^N$ such that $R = \begin{pmatrix} Q \\ \frac{1}{\sqrt{N}} \vec{1}^T \end{pmatrix}$.

Young et al. [9] model opinion dynamics by assuming that each bird averages the differences between its opinion and each of the opinions of its neighbors and changes its opinion to minimize this difference and that each bird's opinion is subject to independent noise with variance proportional to the number of neighbors each bird has. The equations describing these dynamics can be written as

$$\dot{x_i} = \frac{\sum_{\text{neighbors } j \text{ of } i} (x_j - x_i)}{K_i} + \sqrt{n_i} \xi_i = \sum_j \bar{A}_{ij} x_j - x_i + \sqrt{n_i} \xi_i = \sum_j L_{ij} x_j + \sqrt{n_i} \xi_i.$$
 (12)

In vector notation this becomes

$$\dot{x} = Lx + D\xi. \tag{13}$$

The equilibrium of these dynamics is a consensus state in which all birds have the some opinion, $x = \alpha \hat{\mathbf{I}}$ for some $\alpha \in \mathbb{R}$. In the absence of any environmental signal, the consensus opinion can be any value α . In the presence of an environmental signal, some birds in the flock perceive the signal and change their opinions to be closer to 1. The environmental forcing is described by a diagonal matrix B where $B_{ii} = b$ if i perceives the environmental signal and $B_{ii} = 0$ otherwise and the dynamics then become

$$\dot{x} = Lx + B(\vec{1} - x) + D\xi = L_{s}x + v + D\xi. \tag{14}$$

where $L_s = L - B$ and $v = B\vec{1}$. If b = 0, then $L_s = L$ and $v = \vec{0}$, so the case with the signal reduces to the case without the signal.

If $\bar{y} = Rx$, then the first N-1 coordinates of \bar{y} are y = Qx and the last is $\sum_i x_i/\sqrt{N}$. The dynamics of \bar{y} are described by

$$\begin{split} \dot{\bar{y}} &= R\dot{x} \\ &= RL_{\rm s}x + Rv + RD\xi. \\ &= RL_{\rm s}R^T\bar{y} + Rv + RD\xi \\ &= \tilde{L}_{\rm s}\bar{y} + \tilde{v} + \tilde{D}\xi \end{split}$$

where $\tilde{L}_{\rm s}=RL_{\rm s}R^T$, $\tilde{v}=Rv=RB\vec{1}$, and $\tilde{D}=RD$. If $b\neq 0$, $L_{\rm s}$ and therefore $\tilde{L}_{\rm s}$ have all negative eigenvalues and $\bar{y}=(0,0,\dots 0,\sqrt{N})^T$ is a globally asymptotically stable equilibrium. Therefore, by Claim 2 and Theorem 1, $\lim_{t\to\infty} E[\bar{y}(t)\bar{y}(t)^T]=\Sigma_{\bar{y}}$ where $\tilde{L}_{\rm s}\Sigma_{\bar{y}}+\Sigma_{\bar{y}}\tilde{L}_{\rm s}^T=-RD^2R^T$. We can find $\Sigma_y=\lim_{t\to\infty} E[y(t)y(t)^T]$ by removing the last row and column of $\Sigma_{\bar{y}}$.

If b=0, L has a 0 eigenvalue corresponding to the eigenvector $\vec{1}$. Therefore, RLR^T also has a 0 eigenvalue, so we can therefore no longer find the desired $\Sigma_{\bar{y}}$. Since the last row of $R=\frac{1}{\sqrt{N}}\vec{1}$, all entries of the last column of LR^T are 0. Therefore, all entries of the last column of RLR^T are 0. In other words, the dynamics of the first N-1 elements of \bar{y} , i.e. y, do not depend on the last. This lets us write

$$\dot{y} = QLQ^T y + Q\xi = \bar{L} + Q\xi$$

where $\bar{L} = QLQ^T$. Another way to see this is to write

$$\begin{split} \dot{y} &= Q\dot{x} \\ &= QLx + Q\xi \\ &= QL(x - \langle x \rangle \vec{1}) + Q\xi \text{ since } L\vec{1} = \vec{0} \\ &= QLQ^TQx + Q\xi \\ &= \bar{L}y + Q\xi. \end{split}$$

Again, by Claim 2 and Theorem 1, we can now find $\lim_{t\to\infty} E[y(t)y(t)^T] = \Sigma_y$ by finding a solution to $\bar{L}\Sigma_y + \Sigma_y\bar{L} = -QD^2Q^T$.

To summarize, in the absence of an environmental signal, Σ_y satisfies $\bar{L}\Sigma_y + \Sigma_y\bar{L}^T = -QD^2Q^T$ where $\bar{L} = QLQ^T$ and in the presence of an environmental signal, Σ_y can be found in the first N-1 rows and columns of $\Sigma_{\bar{y}}$ where $\Sigma_{\bar{y}}$ satisfies $\tilde{L}_s\Sigma_{\bar{y}} + \Sigma_{\bar{y}}\tilde{L}_s^T = -RD^2R^T$ where $\tilde{L}_s = RL_sR^T$.

6.3 Probability distribution at equilibrium A quite different way of modeling social interactions the method used by Bialek et al. in [3], which is to assume the opinions come to equilibrium and write down an expression for the probability of finding the opinion vector in every state. In statistical physics, a Hamiltonian describes the energy of a given state, H(v), and the Boltzmann distribution describes a probability distribution in which the most energetic states are least likely. Analogously, we can write a function H(v) that gives a high energy when the opinions are not in consensus and a low energy when the opinions are in consensus and a probability distribution of opinion vectors that favors less energetic states. In order to do this, we must use a symmetric interaction matrix, rather than the possibly asymmetric \bar{A} we used above. Bialek et al. study a model in which opinions are a three-dimensional velocity vector and the birds can weight the opinions of their nearest neighbors with any value J. In order to simplify this model and make it directly comparable to the opinion dynamics model we described above, we only consider a scalar-valued opinion x_i and we impose the condition that the weights given to the neighbors' opinions are inversely proportional to the number of neighbors. In particular, we define $S = \frac{1}{2}(A + A^T)$ and P such that $P_{ij} = S_{ij}$ for $j \neq i$ and $P_{ii} = -\sum_k S_{ik}$. This allows us to write down a Hamiltonian

$$H(v) = -v^T P v (15)$$

where the negative sign makes opinions far from consensus give high energies and conversely. The probability distribution over opinion vectors v then becomes

$$P(v) = \frac{1}{Z} \exp\left(-\frac{1}{2}v^T(-P)v\right),\tag{16}$$

where $Z = \int_{\mathbb{R}}^{N} \exp\left(-\frac{1}{2}H(v)\right) dv$. We can now use this probability distribution to find the covariances and correlations between the birds' fluctuations around consensus. If we let $V = \sum_{i} v_i/N = \langle v \rangle$ and $z = v - V\vec{1}$ so that z represents the deviations in each bird's opinion from the consensus opinion. Then

$$P(z, V) = \frac{1}{Z} \exp\left(-\frac{1}{2}(z + V\vec{1})^T(-P)(z + V\vec{1})\right)$$
$$= \frac{1}{Z} \exp\left(-\frac{1}{2}z^T(-P)z\right) \text{ since } P\vec{1} = P^T\vec{1} = \vec{0}.$$

Since P has a 0 eigenvalue, it is not invertible. However, z is orthogonal to the eigenvector of P with eigenvalue 0. Therefore, the covariance of z is $C_z = -P^{\dagger}$. By dividing each entry C_{zij} of the covariance matrix by $C_{zii}C_{zjj}$, the covariance matrix can be used to find the correlation matrix.

The Hamiltonian framework, like the opinion dynamics framework, can also accommodate an environmental signal. In this case, the appropriate Hamiltonian is

$$H(v) = -(v - \vec{1})^T P_{s}(v - \vec{1})$$
(17)

where $P_s = P - B$. It is clear that H is minimized when $v = \vec{1}$. Again, we are interested in the deviations from the consensus opinion. As above, let $V = \sum_i v_i/N$ and $z = v - V\vec{1}$ so that $v = z + V\vec{1}$.

$$P(v) = \frac{1}{Z} \exp\left(-\frac{1}{2}(v - \vec{1})^T(-P_s)(v - \vec{1})\right)$$

= $\frac{1}{Z} \exp\left(-\frac{1}{2}(z - (1 - V)\vec{1})^T(-P_s)(z - (1 - V)\vec{1})\right)$

Since the subspace of $z \in \mathbb{R}^N$ such that $\sum_i z_i = 0$ is N-1 dimensional, it will be easier to rotate z into \mathbb{R}^{N-1} and integrate over all of \mathbb{R}^{N-1} .

$$\begin{split} P(v) &= \frac{1}{Z} \exp \left(-\frac{1}{2} (R^T y - (1 - V) \sqrt{N} R^T e_N)^T (-P_{\rm s}) (R^T y - (1 - V) \sqrt{N} R^T e_N) \right) \\ &= \frac{1}{Z} \exp \left(-\frac{1}{2} (y - (1 - V) \sqrt{N} e_N)^T (-R P_{\rm s} R^T) (y - (1 - V) \sqrt{N} e_N) \right) \\ &= \frac{1}{Z} \exp \left(-\frac{1}{2} (y + \sqrt{N} V e_N - \sqrt{N} e_N)^T \tilde{P}_{\rm s} (y + \sqrt{N} V e_N - \sqrt{N} e_N) \right) \end{split}$$

where $\tilde{P}_s = -RP_sR^T$. To find P(z) we can integrate out V. However, to consider this as a probability density function over y and V, we have to consider the determinant of the transformation from v to y and V. Combining the transformations from v to (z, V) and from z to (y, V),

$$y = RQ^T Qv = Qv$$
 and $V = \frac{1}{N}\vec{1}^T v = \frac{1}{\sqrt{N}}R_N.v$

so that the entire transformation can be represented by $\bar{R} = \begin{pmatrix} Q \\ 1/\sqrt{N}R_N \end{pmatrix}$, so that $\det(\bar{R}) = \frac{1}{\sqrt{N}}\det(R) = \frac{1}{\sqrt{N}}$. Therefore, $P(y,V) = \sqrt{N}P(v)$ is the appropriate density function for y and V. Finally, note that $Z = \sqrt{(2\pi)^N \det(-P_{\rm s}^{-1})}$. This gives

$$\begin{split} P(y,V) &= \sqrt{N}P(v) \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}((V-1)\sqrt{N}e_N + y)^T \tilde{P}_s((V-1)\sqrt{N}e_N + y)\right) \\ &\Rightarrow P(y) = \int_{\mathbb{R}} P(y,V)dV \\ &= \int_{\mathbb{R}} \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}((V-1)\sqrt{N}e_N + y)^T \tilde{P}_s((V-1)\sqrt{N}e_N + y)\right) dV \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}y^T \tilde{P}_s y\right) \int_{\mathbb{R}} \exp\left(-\frac{1}{2}\left(\tilde{P}_{sNN}N\left(V-1\right)^2 + (V-1)\sqrt{N}\tilde{P}_{sN\cdot}y + (V-1)\sqrt{N}y^T \tilde{P}_{s\cdot}N\right)\right) dV \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}y^T \tilde{P}_s y\right) \int_{\mathbb{R}} \exp\left(-\frac{1}{2}\left(\tilde{P}_{sNN}N\left(V-1\right)^2 + 2(V-1)\sqrt{N}\tilde{P}_{sN\cdot}y\right)\right) dV \text{ since } \tilde{P}_s \text{ is symmetric} \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}\left(y^T \tilde{P}_s y - \left(\frac{\tilde{P}_{sN\cdot}y}{\sqrt{\tilde{P}_{s}NN}}\right)^2\right)\right) \times \\ &\int_{\mathbb{R}} \exp\left(-\frac{1}{2}\left(\tilde{P}_{sNN}N\left(V-1\right)^2 + 2(V-1)\sqrt{N}\tilde{P}_{sN\cdot}y + \left(\frac{\tilde{P}_{sN\cdot}y}{\sqrt{\tilde{P}_{sNN}}}\right)^2\right)\right) dV \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}\left(y^T \left(\tilde{P}_s - \frac{1}{\tilde{P}_{sNN}}\tilde{P}_{s\cdot}N\tilde{P}_{sN\cdot}\right)y\right)\right) \int_{\mathbb{R}} \exp\left(-\frac{1}{2}\left(\sqrt{\tilde{P}_{sNN}N}\left(V-1\right) + \frac{\tilde{P}_{sN\cdot}y}{\sqrt{\tilde{P}_{sNN}}}\right)^2\right) dV \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}\left(y^T (\tilde{P}_s - \frac{1}{\tilde{P}_{sNN}}\tilde{P}_{s\cdot}N\tilde{P}_{s\cdot})y\right)\right) \int_{\mathbb{R}} \exp\left(-\frac{1}{2}\tilde{P}_{sNN}N \left(V-1 + \frac{\sum_i \tilde{P}_{sNi}y_i}{\tilde{P}_{sNN}\sqrt{N}}\right)^2\right) dV \\ &= \frac{\sqrt{N}}{Z} \exp\left(-\frac{1}{2}\left(y^T (\tilde{P}_s - \frac{1}{\tilde{P}_{sNN}}\tilde{P}_{s\cdot}N\tilde{P}_{s\cdot})y\right)\right) \times \sqrt{\frac{2\pi}{\tilde{P}_{sNN}N}} \\ &= \frac{1}{\sqrt{(2\pi)^{N-1}\det(-\tilde{P}_s^{-1})\tilde{P}_{s\cdot}N}}} \exp\left(-\frac{1}{2}\left(y^T (\tilde{P}_s - \frac{1}{\tilde{P}_{sNN}}\tilde{P}_{s\cdot}N\tilde{P}_{s\cdot})y\right)\right) \end{split}$$

Therefore, $E[y_i] = 0$ for all i and $E[yy^T] = (\tilde{P}_s - \frac{1}{\tilde{P}_{sNN}} \tilde{P}_{sN} \tilde{P}_{sN})^{-1}$. Then,

$$E[zz^T] = R^T \left(\tilde{P}_{\mathrm{s}} - \frac{1}{\tilde{P}_{\mathrm{s}NN}} \tilde{P}_{\mathrm{s}N} \tilde{P}_{\mathrm{s}N} \right)^{-1} R = \left(-P_{\mathrm{s}} - \frac{1}{\tilde{P}_{\mathrm{s}NN} N} B \vec{1} \vec{1}^T B \right)^{-1}$$

Let $N_{\rm I}$ be the number of birds that can perceive the signal, i.e. $N_{\rm I}$ is the number of non-zero elements along the diagonal of B. Then, $\tilde{P}_{{\rm s}NN}=\frac{b}{N}N_{\rm I}$. Finally, we find that

$$C_z = E[zz^T] = \left(-P_s - \frac{1}{bN_I}B\vec{1}\vec{1}^TB\right)^{-1}$$
 (18)

Now, let $P' = P_{\rm s} + \frac{1}{bN_{\rm I}}B\vec{1}\vec{1}^TB$. Then $P'\vec{1} = -B\vec{1} - \frac{1}{bN_{\rm I}}B\vec{1}\vec{1}^TB\vec{1} = -B\vec{1} - \frac{bN_{\rm I}}{bN_{\rm I}}B\vec{1} = \vec{0}$, so we cannot invert -P' to find the covariance matrix. However, as above, we can discard the 0 eigenvector since we know that $z \cdot \vec{1} = 0$. This gives us $E[(v_i - \frac{\sum_k v_k}{N})(v_j - \frac{\sum_k v_k}{N})]$.

6.4 Connecting two frameworks To find \mathcal{H} , we found the matrix Σ_y such that

$$QLQ^T \Sigma_y + \Sigma_y QLQ^T = -QD^2 Q^T. (19)$$

Table 1: P^{\dagger} is the inverse of P with the 0 eigenvector / value removed.

	Without signal	With signal
From noisy dynamics	$\bar{L}\Sigma_y + \Sigma_y \bar{L}^T = -QD^2Q^T$	$\tilde{L}_{\mathrm{s}}\Sigma_{\bar{y}} + \Sigma_{\bar{y}}\tilde{L}_{\mathrm{s}}^T = -RD^2R^T, \ \Sigma_y = (\Sigma_{\bar{y}})_{i,j=1}^{N-1}$
From Boltzmann equilibrium	$C_y = -QP^{\dagger}Q^T$	$C_y = (\tilde{P}_{\mathrm{s}} - \frac{1}{\tilde{P}_{\mathrm{s}NN}} \tilde{P}_{\mathrm{s}N} \tilde{P}_{\mathrm{s}N.})^{\dagger}$
		or $C_y = Q(-P_s - \frac{1}{bN_I}B\vec{1}\vec{1}^TB)^{\dagger}Q^T$

If the vector v is described by the dynamics $\dot{v} = Dv$, with $H(v) = -v^T Pv$, then

$$\begin{split} \dot{H} &= -\dot{v}^T P v - v^T P \dot{v} \\ &= -v^T D^T P v - v^T P D v \\ &= -v^T (D^T P + P D) v. \end{split}$$

Therefore, H(v) decreases at all v if and only if $D^TP + PD$ is a positive matrix.

Table 1 shows how to find the appropriate covariance matrices depending on the framework being used and whether or not a signal is present. If we use P instead of L with the noisy dynamics framework and all birds have the same number of neighbors, then $C_y = 2\Sigma_y$.

Table 2: Table of matrices and variables used in the text.

A	adjacency matrix	$A_{ij} = \begin{cases} 1 & \text{if } i \text{ pays attention to } j \\ 0 & \text{otherwise} \end{cases}$ $\bar{A}_{ij} = \begin{cases} \frac{1}{K_i} & \text{if } i \text{ pays attention to } j, \text{ where } K_i = \# \text{ of bird of } j \\ 0 & \text{otherwise} \end{cases}$ $B_{ij} = \begin{cases} b & \text{if } i = j \text{ and } i \text{ perceives the environmental signs } j \\ 0 & \text{otherwise} \end{cases}$
		0 otherwise $\frac{1}{2}$ if i pays attention to i where $K_i = \#$ of hir
\bar{A}	normalized adjacency matrix	$\bar{A}_{ij} = \begin{cases} K_i & \text{if } i \text{ pays attention to } j, \text{ where } K_i = \# \text{ or bird} \\ 0 & \text{otherwise} \end{cases}$
D	1. 1. 1	b if $i = j$ and i perceives the environmental signal
В	diagonal signal matrix	$B_{ij} = \begin{cases} 0 & \text{otherwise} \end{cases}$
λ	correlation length	
Λ	diagonal matrix of eigenvalues of P	
L	opinion dynamics w/o environmental signal	$L = \bar{A} - I$
$L_{ m s}$	opinion dynamics with environmental signal	$L_{\rm s} = L - B$
$ar{L}$	dynamics of deviations from consensus w/o signal	
$ ilde{L}_{ m s}$	dynamics of deviations from consensus w/ signal	$\tilde{L} = RL_{\mathrm{s}}R^T$
S	symmetrized interaction matrix	$S = \frac{1}{2}(\bar{A} + \bar{A}^T)$
P	energy interactions w/o environmental signal	$P_{ij} = \begin{cases} S_{ij} & \text{for } j \neq i \\ -\sum_k S_{ik} & \text{for } j = i \end{cases}$
D	onergy interactions with environmental signal	•

$$\tilde{P}_{\rm s}$$

$$Q \quad \mbox{rotates opinion vector away from consensus}$$

energy interactions with environmental signal

$$R$$
 full rotation matrix whose last row is $\frac{1}{\sqrt{N}}\vec{\mathbf{1}}^T$

$$\rho$$
 \mathscr{H}_2 robustness

 $P_{\rm s}$

$$W$$
 matrix of eigenvectors of P

$$Q \in \mathbb{R}^{N-1} \times \mathbb{R}^N$$
 such that $Q\vec{1} = \vec{0}$, $QQ^T = I_{N-1}$, and $Q^TQ = I_{N-1}$

$$\lim_{t\to\infty} \frac{1}{E[||y||]}$$
 where $y=Qx$

 $P_{\rm s} = P - B$

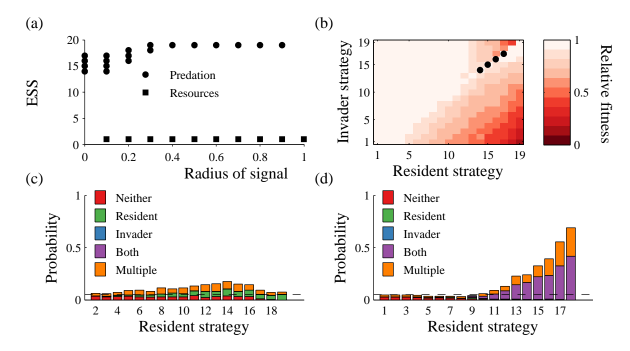


Figure 1: The ESS number of neighbors is always higher if selection is due to predation than due to resources. When selection is due to predation, there can be multiple ESS strategies, which are a non-monotonic function of the radius of the signal. In (a), we show the (possibly multiple) ESS strategies as a function of the radius of the signal. In (b) we show the relative fitness of an invader as a function of the invader and resident strategies when r = 0.1. The dots show the multiple ESSs. For (c) and (d), we choose a focal resident bird and consider the situation where neither the focal bird nor the invader has the environmental signal. The total of the stacked bars is the total probability that the invader will be predated when neither the focal bird nor the invader have the signal. We further break down these cases into five cases: there is a single informed bird and neither focal bird, only the focal resident, only the invader, or both pay attention to it or there are multiple informed birds. The horizontal dashed line shows 1/N, the probability of being predated if the strategies are neutral. In (c), the invader has one fewer neighbor than the resident. In (d), the invader has one more neighbor than the resident. Parameters: in all panels N = 20, $\beta = 1$, T = 1, in (b)-(d) r = 0.1.

7 Figures

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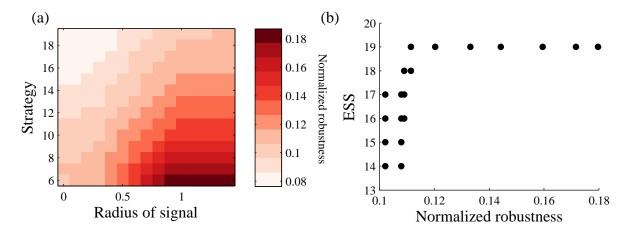


Figure 2

Figure 3: When the birds choose the optimal number of neighbors, given the strategies the rest of the flock are using, they settle on higher strategies when selection is due to predation than when selection is due to resources. Under selection due to predation, the \mathcal{H}_2 robustness and correlation length of the flock tend to decrease, whereas under selection due to resources, \mathcal{H}_2 robustness is relatively constant and correlation length tends to increase. The upper row shows results from implementing selection based to predation and the lower row shows results from implementing selection based on resources. In each panel, the horizontal axis represents the number of times the birds are allowed to choose optimal strategies. The first column shows one example of how the birds' strategies change over time. The second column shows, for many initial conditions, how the \mathcal{H}_2 robustness changes over time. The third column shows, for many initial conditions, how the correlation length changes over time. Parameters: N=20, $\beta=1$, r=.1, T=1.

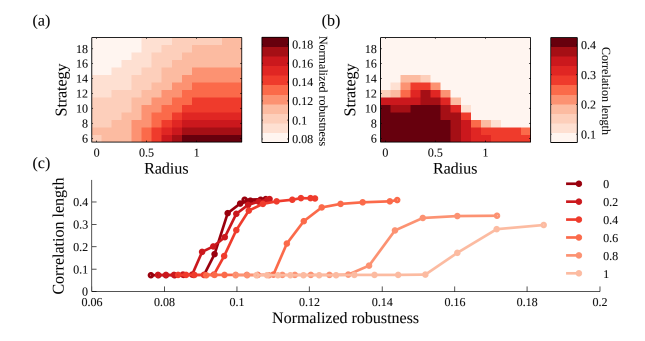


Figure 4

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