A least microenvironmental uncertainty principle (LEUP) as a generative model of collective cell migration mechanisms

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Collective migration is commonly observed in groups of migrating cells, in the form of swarms or aggregates. Mechanistic models have proven very useful in understanding collective cell migration. Such models, either explicitly consider the forces involved in the interaction and movement of individuals or phenomenologically define rules which mimic the observed behavior of cells. However, mechanisms leading to collective migration are varied and specific to the type of cells involved. Additionally, the precise and complete dynamics of many important chemomechanical factors influencing cell movement, from signalling pathways to substrate sensing, are typically either too complex or largely unknown. The question is how to make quantitative/qualitative predictions of collective behavior without exact mechanistic knowledge. Here we propose the least microenvironmental uncertainty principle (LEUP) that may serve as a generative model of collective migration without precise incorporation of full mechanistic details. Using statistical physics tools, we show that the famous Vicsek model is a special case of LEUP. Finally, to test the biological applicability of our theory, we apply LEUP to construct a model of the collective behavior of spherical Serratia marcescens bacteria, where the underlying migration mechanisms remain elusive.

Collective movement of dense populations is observed in several biological systems at different scales, from massive migration of mammals1 to cells during embryogenesis2. In these systems, individuals which are able to propel themselves independently and interact with other nearby, start moving in a coordinated fashion once enough similar individuals are brought together. Due to the relevance of many of these processes to human activity, as well as their pervasiveness, there is a need for quantitative understanding of collective migration. Mechanistic models, in particular, incorporate the driving interactions between individuals in the specific system modeled. It is clear that different types of individuals, especially across different spatial scales, synchronize their movements through different mechanisms. This results in a variety of models specific to certain individual species1,3,4. In the specific case of biological cells, cellular migration involves numerous biophysical processes such as actin polymerization, receptor recruitment, or in bacteria flagellar motor reversal mechanisms to name a few5. However, in many cases the exact knowledge of all participating biophysical/chemical mechanisms related to a particular collective migration pattern is not trivial. Therefore, there is a need to construct models of collective migration which do not require knowledge of the exact interactions between individuals.

To address this situation, several mathematical models introduce a phenomenological short-range bias that every individual feels. In one of the most influential collective migration models6 the so-called Vicsek model, the direction of movement of particles changes towards the mean velocity of individuals in a local neighborhood, inducing long-range swarming at the population level. Such models can be further refined into mechanistic models, where individual particle dynamics are dictated by a system of Langevin equations. In Langevin equation...
models, the reorientation of individual particle velocities is brought about by the existence of a local interaction potential, which is determined by neighboring particle properties. Collective migration has been achieved, for example, through the introduction of a ferromagnetic-like interaction potential, which locally aligns particle velocities polarly, or a liquid-crystal-like interaction potential, which aligns particle velocities nematically7. Often neither biophysical nor phenomenological models are able to provide a plausible explanation or quantitative reproduction of collective migration patterns, due to the lack of complete mechanistic knowledge. Such an example is the spatiotemporal dynamics of spherical S. marcescens bacteria. Interestingly, prior modeling works8 were able to partially reproduce the experimental results, since the underlying biophysical mechanisms are still unclear. In such cases, one could rely of machine/statistical learning methods that circumvent the biophysical details9–11. However, such methods are typically of high accuracy but low interpretability, i.e. they are “black boxes” that do not offer mechanistic insights, and prone to overfitting.

Here, we view migration as an active decision-making process. Cell decision-making is the process of cells changing their phenotype according to their intrinsic programming and in response to the microenvironmental cues12–15. Cell decisions involve complex biochemical regulation in the genetic, epigenetic, translational or transcriptional level. The fundamental challenges are the (i) uncertainty of high-dimensional subcellular regulatory cell decision-making mechanisms, and the (ii) lack of knowledge in the relative contribution of intrinsic and extrinsic cell decision-making factors to multicellular spatiotemporal dynamics. Regarding cells as Bayesian decision-makers under energetic constraints, it has been proposed that cell decisions are dictated by a ‘Least microEnvironmental Uncertainty Principle (LEUP)’. This is translated into a free-energy principle, implying a statistical mechanics theory for cell decision-making. Such a statistical mechanics reduction allows for simplifying many parameters into a low-dimensional mathematical description and circumvent the uncertainty about the underlying mechanisms. Moreover, it allows to integrate heterogeneous data types as constraints of LEUP energy optimization. Applying the LEUP to collective cell migration, we aspire (i) to provide a low-dimensional statistical mechanics description, (ii) circumvent the uncertainty about the underlying biophysical mechanisms and (iii) provide a relationship to phenomenological models (e.g. the Vicsek model). Finally in Fig. 1, we illustrate how LEUP is positioned in terms of model interpretability and knowledge of biophysical details in comparison the afore-mentioned modelling/computational approaches. Interestingly, LEUP proposes a balanced solution for problems of low mechanistic knowledge and satisfactory interpretability.

Figure 1. Applicability of different modelling/computational approaches according to their biological interpretability and the corresponding knowledge of biophysical migration mechanisms. The latter is typically encoded as the degree of interaction potential knowledge in Langevin equations (see text). In the case of extended biophysical mechanism knowledge, mechanistic models are the natural choice. When the effects of cell–cell interaction on cell migration are only partially understood, phenomenological models can be typically used. Finally, when data do not suffice to formulate an interaction potential, machine learning allows for the quantitative reproduction of experimental data. However, this has a toll in the interpretability of the resulting model, since machine learning methods are typically "black boxes". LEUP models offer a compromise that allows for quantitative predictions under lack of mechanism knowledge and satisfactory biological interpretability.
In this work, we present the simplest LEUP-driven Langevin model of swarming where individuals can sense the velocity orientations of other individuals in their surroundings. Individuals act as Bayesian inferences and change their own orientation to optimize their prior, according to environmental orientation information. Under these assumptions, individuals reorient according to the entropy gradient of the environmental information. A parameter, named the sensitivity, controls the strength and directionality of the reorientation in relation to the local gradient. We find that the system adopts a steady, polar-ordered state for negative values of the sensitivity. Conversely, the system remains out of equilibrium, but partially nematic-ordered when the sensitivity is positive.

Furthermore, we find that the qualitative behavior of the model depends on the values of the particle density, noise strength, sensitivity, and size of the interaction neighborhood. Finally, we showcase the LEUP principle by showing that our model replicates the collective behavior of spherical S. marcescens bacteria.

Materials and methods
The self-propelled particle framework. Moving and interacting cells are modeled by a two-dimensional self-propelled particle model (SPP). In this model, \( N \in \mathbb{N} \) cells move on a two-dimensional area. The \( n \)-th cell is characterized by its position, \( \vec{r}_n \in \mathbb{R}^2 \), speed, \( v_n \in [0, \infty) \subset \mathbb{R} \), and an orientation \( \theta_n \in [0, 2\pi) \subset \mathbb{R} \). Due to the small size of cells, it is assumed that viscous forces dominate. Changes in speed and orientation result from local potentials \( U_\theta(\vec{r}_m, \theta_m, v_m) \), \( U_v(\vec{r}_m, \theta_m, v_m) : \mathbb{R}^2 \times [0, 2\pi) \times [0, \infty) \mapsto \mathbb{R} \) which depend on the positions and polar velocity components of cells within a radius \( R \in \mathbb{R}_+ \). The bias of the cell to follow the potential gradients are regulated by the parameters \( \beta_\theta, \beta_v \in \mathbb{R} \), called angular and radial sensitivities, respectively. Additionally, velocity fluctuations occur due stochastic noise terms \( \xi^\alpha_n(t) \in [0, 2\pi) \), \( \alpha \in \{\theta, v\} \) where \( t \in \mathbb{R}_+ \) denotes time. The noise will be assumed to be a zero-mean, white noise term, which has the statistical properties \( \langle \xi^\alpha_n(t_1) \xi^\beta_m(t_2) \rangle = 0 \) and \( \langle \xi^\alpha_n(t_1) \xi^\alpha_n(t_2) \rangle = 2D_\alpha \delta(t_1 - t_2) \delta_{nm} \), where \( t_1 \) and \( t_2 \) are two time points, \( D_\alpha \in \mathbb{R}_+ \) is either the angular \( (\alpha = \theta) \) or radial \( (\alpha = v) \) diffusion coefficient, \( \delta(t) \) is the Dirac delta, and \( \delta_{nm} \) is the Kronecker delta. Finally, the radial acceleration will be assumed to be damped by a density dependent friction, \( \psi(\rho_n) \). In the following, it will be assumed that the density-dependent friction is given by \( \psi(\rho_n) = \rho_n - \bar{\rho}_0 \), where \( \rho_n \) is the local cell density within the \( n \)-th cell's interaction radius, and \( \bar{\rho}_0 \) is the global average cell density. Taking everything into account, the stochastic equations of motion of the \( n \)-th cell are

\[
\frac{d}{dt} \vec{r}_n = v_n \hat{v}(\theta_n) \tag{1a}
\]

\[
\frac{d}{dt} \theta_n = -\beta_\theta \frac{\partial}{\partial \theta_n} U_\theta(\vec{r}_m, \theta_m, v_m) + g(\vec{v}_n) \xi^\theta_n(t) \tag{1b}
\]

\[
\frac{d}{dt} v_n = -\beta_v \frac{\partial}{\partial v_n} U_v(\vec{r}_m, \theta_m, v_m) - \epsilon \psi(\rho_n) v_n + \xi^v_n(t). \tag{1c}
\]

where \( \hat{v}(\theta_n) \) is the normalized velocity of the cell and \( \epsilon \) is a parameter. A representation of the SPP model is shown in Fig. 2. The function \( g(\vec{v}_n) \) modulates the noise variance and allows us to model certain distributions (such as the Rayleigh distribution in Section 5).
The interaction potentials $U_a(\vec{r}_m, \theta_m, \vec{v}_m)$, which dictate the velocity dynamics of cells, need to be specified. Biophysically, the potentials should encompass steric effects, hydrodynamic interactions, chemotactic effects, and terms arising from internal cellular processes, for example, flagellar motor dynamics, actin polymerization, receptor dynamics, etc. Finding such potentials is a formidable task since not all of the mechanisms and interactions involved are known. To circumvent this problem, a variational principle of cell decision-making related to entropy maximization\textsuperscript{17}, known as the least microenvironmental uncertainty principle (LEUP), will be used\textsuperscript{18}. In the next section, we will discuss such a case.

**Least microenvironmental uncertainty principle (LEUP).** The main premise of LEUP is that cells equip Bayesian inference to decide their internal states, expressed as a combination of a sensed microenvironmental distribution (empirical/measured likelihood distribution) and a local entropy-dependent prior. From the cellular point of view, constructing an accurate microenvironmental sensing distribution is expensive, since cell sensing is an energetically costly process. Cells sense (collect information) of their surroundings by employing different processes, such as polymerizing pseudopodia, translocating receptor molecules or modifying its cytoskeleton according to mechanical signals\textsuperscript{19,20}. However, the cost can be minimized when cells build informative priors about their microenvironment. In the case of moving cells this could be achieved by, for example, promoting cell polarization through the recruitment of actin related proteins\textsuperscript{21}. In turn, this allows cells to spare the energy from building extra sensory processes. At same time, cell polarization effectively promotes the co-evolution of the cellular state and its microenvironment, where the latter becomes more predictable. The simultaneous determination of cell state and its microenvironment results in the minimization of the microenvironmental uncertainty towards a target value related to a given tissue or bacteria (e.g. biofilm). This process also entails an energy cost. It is reasonable to assume that the cell will try to optimize its prior $P(\alpha_n)$ for the sake of energetic frugality.

The prior probabilities $P(\alpha_n)$ should, consequently, fulfill certain the premises of LEUP. First and foremost, they should be normalized, i.e. \( \int P(\alpha_n = a) da = 1 \), integrating over all possible values of internal cellular states. Second, a biological cell is an imperfect sensor. Therefore, the uncertainty in sensing accuracy $S(A_n | \alpha_n)$, should reach a certain level in average, which is species-dependent. This assumption is a cornerstone of the LEUP formalism. An important feature of the LEUP formalism, and of the entropy maximization principle in general, is its applicability when every other mechanistic details of the system are unknown. Entropy is a measure of uncertainty according to information theory, therefore entropy should be maximal, in order to reflect our lack of mechanistic knowledge of the phenomenon, and to avoid introducing any artificial bias in the model arising from the specific choice of $P(\alpha_n)\textsuperscript{22}$. Given that the internal entropy is given by $S(\alpha_n) = - \int P(\alpha_n = a) \ln P(\alpha_n = a) da$, entropy maximization subjected to probability normalization and the target mean sensing accuracy translates into the optimization problem

$$
\frac{\delta}{\delta P(\alpha_n)} \left\{ - \int P(\alpha_n = a) \ln P(\alpha_n = a) da \right. \\
\left. - \beta_a \int P(\alpha_n = a) S(A_n | \alpha_n = a) da - (S(A_n | \alpha_n) - S^\text{\textbar end}) \right\} = 0,
$$

where $\frac{\delta}{\delta P(\alpha_n)}$ is the functional derivative, $S(A_n | \alpha_n) = \text{target sensing accuracy}$, and $\beta_a$ are Lagrange multipliers. Taking into account the relations among entropy and probability, Eq. (2) yields

$$
P(\alpha_n = a) = e^{-\beta_a S(A_n | \alpha_n = a)} Z,
$$

where $Z = \int e^{-\beta_a S(A_n | \alpha_n = a)} da$ is a normalization constant and $\beta_a$ is the responsiveness of the cell. Using Eq. (3), the internal entropy of the cell, defined as $S(\alpha_n) = - \int P(\alpha_n = a) \ln P(\alpha_n = a) da$, is given by

$$
S(\alpha_n) = \tilde{\beta}_a S(A_n | \alpha_n) + \ln Z.
$$

Using the relation between thermodynamic-like potentials, it is evident that the average internal energy is given by
\[ U(\alpha_n, \alpha_n) = S(A_n | \alpha_n) = \langle S(A_n | \alpha_n \neq \alpha_n) \rangle \]  

and for a particular realisation of \( \alpha_n \) we can write the internal energy as

\[ U_\alpha(\alpha_n, \alpha_n) = S(\alpha_n | \alpha_n) \]  

Now, Helmholtz-like free energy can be written as

\[ F = -\frac{1}{\tilde{\beta}_\alpha} \ln Z. \]  

Finally, we return back to the Bayesian formalism and to the main LEUP premise that cells tend to build informative microenvironmental priors \( P(\alpha_n) \). In the steady state, the latter implies that

\[ P(\alpha_n | A_n) \xrightarrow{t \to \infty} P(\alpha_n), \]  

i.e. the posterior distribution of \( P(\alpha_n | A_n) \) becomes independent of measuring/sensing the microenvironmental information \( A_n \), since the prior \( P(\alpha_n) \) includes all relevant information. Observing the Bayesian probability, this only happens when the measurement/sensing likelihood \( P(A_n | \alpha_n) \) approaches the probability distribution of the microenvironment \( P(A_n) \), i.e. the cell perfectly senses its microenvironment. Using again the Bayesian probability, we can prove the following information-theoretic equation

\[ I(\alpha_n, A_n) = S(\alpha_n) - S(\alpha_n | A_n) \xrightarrow{t \to \infty} 0. \]  

Substituting the above in Eq. (4), the mutual information reads

\[ I(\alpha_n, A_n) = \tilde{\beta}_\alpha S(A_n | \alpha_n) + \ln Z - S(\alpha_n | A_n). \]  

Two cases have to be considered, depending on the sign of \( \tilde{\beta}_\alpha \):

- When \( \tilde{\beta}_\alpha < 0 \) the only positive term is \( \ln Z \) in Eq. (10). Thus, it is reasonable to expect the mutual information to decrease to a minimum value.
- When \( \tilde{\beta}_\alpha \geq 0 \) one can show that the system, for a large enough sensing radius, goes to an ordered state at the steady state (see “Collective cell migration patterns for different parameter regimes”). This implies that the microenvironmental entropy \( S(A_n | \alpha_n) \) tends to zero. In this case, the Eq. (10) becomes

\[ I(\alpha_n, A_n) = \ln Z - S(\alpha_n | A_n). \]

This implies that the mutual information has the potential to reach zero. Interestingly, if this happens using Eq. (9), we can recover a similar definition to Boltzmann entropy for the internal cell state entropy, i.e.

\[ S(\alpha_n) = \ln Z. \]

**LEUP-based dynamics.** The internal energy depends on the internal states of the cell, as well as the internal states of other cells in the surroundings. Such internal states can be a vector of physical quantities (e.g. velocity, acceleration) and/or chemical variables such as intracellular proteins, genes and so on. Here, we focus on the former to define an interaction potential that models the equations of motion.

By doing so, it is evident that the responsiveness of the cells to LEUP can be quantified by the sensitivity \( \beta_\alpha = -\tilde{\beta}_\alpha \). Analogously, we can write the equations of motion of the model similar to Eq. (1c) as

\[ \frac{d}{dt} r_n = v_n \tilde{v}(\theta_n) \]  

\[ \frac{d}{dt} \theta_n = \beta_\nu \frac{\partial}{\partial \theta_n} S(\Theta_n | \theta_n) + g(\tilde{v}_n) \xi_\nu^\theta(t) \]  

\[ \frac{d}{dt} v_n = \beta_\nu \frac{\partial}{\partial v_n} S(V_n | v_n) - \varepsilon \psi(\rho_n) v_n + \xi_\nu^v(t). \]

To illustrate entropy calculation, it will be assumed that the orientations of cells within the interaction neighborhood are distributed according to

\[ P(\theta \in \Theta_n | \theta_n) = \frac{\sinh \gamma}{2\pi [\cosh(\gamma) - \cos(\bar{\theta} - \mu)]}, \]  

where \( \mu \) is the mean of the distribution and \( \gamma \) is a parameter related to the variance. This is a wrapped Cauchy distribution, periodic over the interval \([0, 2\pi]\). Similarly, cell speeds will be assumed to be distributed half-normally

\[ P(v \in V_n | v_n) = \sqrt{\frac{2}{\sigma^2 \pi}} \exp \left( -\frac{v^2}{2\sigma^2} \right), \]
where $\sigma^2$ is proportional to the variance of the distribution. Accordingly, the angular entropy is

$$S(\theta_n \mid \theta_0) = \ln(2\pi) + \ln(1 - e^{-2\gamma}), \quad (14)$$

while the speed entropy is

$$S(V_n \mid v_n) = \frac{1}{2} \ln \left( \frac{\pi \sigma^2}{2} \right) + \frac{1}{2}, \quad (15)$$

The parameter $\sigma$ can be determined from the local speed variance, while the parameter $\gamma$ depends on the local polar order (i.e. the degree of parallel alignment) of cell velocities in the neighborhood. It should be noted that the qualitative behavior of the model is independent of the particular choice of distributions, and the distributions considered here are suggested only for ease of calculation. Before defining $\gamma$, we will first define the observables characterizing the order of the velocity field.

**Collective migration observables.** Let us define the normalized complex velocity of the $n$-th cell, $z_n \in \mathbb{C}$ as $z_n = e^{\theta_n}$, where $i$ is the imaginary unit. The $k$-th moment of the velocity over an area $A$ is given by

$$\langle z^k \rangle_A = \frac{1}{N_A} \sum_{m \in A} z_{m,k},$$

where the sum is over all cells in area $A$, and $N_A$ is the total number of cells in $A$. The polar order parameter in the area $A$ is given by

$$S_A^1 = |\langle z \rangle_A|, \quad (16)$$

which is the modulus of the first moment of the complex velocity in $A$, while the nematic order parameter in the area $A$ is given by

$$S_A^2 = |\langle z^2 \rangle_A|, \quad (17)$$

where $\langle z^2 \rangle_A$ is the second moment of the complex velocity in $A$. The order parameters are bounded, i.e.

$$0 \leq S_A^1, S_A^2 \leq 1, \quad (18)$$

due to the complex velocities $z_n$ being normalized. The parameter $\gamma$ for the distribution of orientations in the neighborhood of the $n$-th cell is given by

$$\gamma = -\ln \left( S_{C_{RN}}^1 \right), \quad (19)$$

where the subindices $C_{RN}$ indicate a circular area of radius $R$ centered at $\hat{r}_n$. The former directly stems from the properties of the wrapped Cauchy distribution.

While global polar and/or nematic order are characteristic of steady flows, rotating flow fields are commonly observed in out-of-equilibrium systems. The vorticity is an observable which is equal to twice the local angular velocity, and is thus a measure of the local strength and direction of rotation of the field. The vorticity $\omega$ is defined as

$$\omega(\hat{r}) = [\nabla \times \vec{v}_{\text{mean}}(\hat{r})] \cdot \hat{k}, \quad (20)$$

where $\vec{v}_{\text{mean}}(\hat{r})$ is the mean velocity field at point $\hat{r}$, and $\hat{k}$ is vector normal to the plane where cells move.

**Statistical evaluation of experiments and model predictions.** For the statistical evaluation of the results we have used the $\chi^2$-test. The testing hypothesis is that the experimental data are explained by the model predictions. To test it, we construct

$$\chi^2_i = \sum_{j=1}^{N} \left( \frac{\hat{O}_i^{(j)} - O_i^{(j)}}{\sigma_i^{(j)}} \right)^2, \quad (21)$$

where $O_i^{(j)}$ is the experimental values of a certain observable $j$, being either speed or vorticity, and the $\hat{O}_i^{(j)}$ and $\sigma_i^{(j)}$ is the corresponding mean value of the stochastic model predictions, based on an ensemble of 50 simulations for each density point $i = 1, \ldots, N$, $N = 11$. The quantity $\frac{\hat{O}_i^{(j)} - O_i^{(j)}}{\sigma_i^{(j)}}$ can be viewed as a z-score for each $\hat{O}_i^{(j)}$ and for large enough simulation ensemble should converge to a normal distribution. The total degrees of freedom for both observables is $2N = 22$. The we calculate the reduced $\chi^2$-statistic, or $\chi^2$ per degree of freedom, which is defined as $\chi^2_{2N} = (2N)^{-1} \sum_i \chi^2_i = 1.97$ being close enough to 1. This suggests that our fitting is satisfactory, since values $\chi^2_{2N} \gg 1$ indicate a bad fit to the experimental data.

**Results**

**From LEUP to phenomenological models of collective migration: the relationship to the Vicsek model.** By using the LEUP, we have modeled interaction as a change in velocity dictated by the local entropy gradient. The modulation of $\beta_\alpha$ parameters modulates the response of cells to the local entropy gradient and gives rise to relationships with known phenomenological models, such as the Vicsek model. The absolute value $|\beta_\alpha|$ is proportional to the likelihood of the cell to change its velocity according to a given entropy gradient. If $\beta_\alpha < 0$, cells tend to go against the local entropy gradient towards the entropy minimum. In the specific
case of $\alpha = \theta$, a negative sensitivity would restrict the distribution of angles to a narrow selection. Conversely, $\beta_\theta > 0$ forces cells to follow the entropy gradient towards the entropy maximum, broadening the distribution. From here on, we will assume that the effect of cell interactions will be averaging the radial component, therefore $\beta_\theta < 0$.

To evaluate the effect of these two opposite migration strategies, we analyze the angular steady states in the two parameter regimes. Without loss of generality, we assume that, in the steady state, the mean velocity is $\bar{v} = 1$. By expanding $S_n$ using Eq. (16), defining the components of the mean neighborhood velocity as $\bar{v}_{x,n} = \sum_{m,d} C_{n,m} \sin \theta_{m,n}$ and $\bar{v}_{y,n} = \sum_{m,d} C_{n,m} \cos \theta_{m,n}$, and differentiating Eq. (14), we find that the orientation of $\theta_n$ at the entropy extrema must be such that (see Supporting Information)

$$\tan \theta_n = \frac{\bar{v}_{y,n}}{\bar{v}_{x,n}},$$

but $\frac{\bar{v}_{x,n}}{\bar{v}_{y,n}} = \tan \bar{\theta}$, the tangent of the mean orientation of the neighbors, excluding the n-th cell. This results in two extremum points $\theta_n = \bar{\theta}$ and $\theta_n = \bar{\theta} + \pi$, one where the velocity of the n-th cell is parallel to the average velocity of its neighbors, and one when it is antiparallel. In the first case

$$\sin \theta_n \propto \bar{v}_{y,n} \text{ and}$$

$$\cos \theta_n \propto \bar{v}_{x,n},$$

while in the second case

$$\sin \theta_n \propto -\bar{v}_{y,n} \text{ and}$$

$$\cos \theta_n \propto -\bar{v}_{x,n}.$$

It can be shown (see Supporting information) that $\theta_n = \bar{\theta}$ corresponds to an entropy minimum, while $\theta_n = \bar{\theta} + \pi$ corresponds to an entropy maximum. Consequently, the behavior of the regime $\beta_\theta < 0$ is analogous to that of the Vicsek model\(^4\). Conversely, the regime $\beta_\theta > 0$ corresponds to an anti-ferromagnetic analog of the Vicsek model.

Next, let us assume that the model has a steady state, where the Helmholtz free energy per cell is given by Eq. (7). Due to its extensivity, the Helmholtz free energy of complete, non-interacting, steady state system is

$$F_T \approx -\frac{1}{\beta_\theta} \sum_{n=1}^{N} \ln Z_n = -\frac{1}{\beta_\theta} \ln \left( \prod_{n=1}^{N} Z_n \right),$$

where $Z_n$ is the normalization constant of Eq. (3) for the n-th cell. For a weakly interacting system, the mean-field effective normalization constant $Z_T := \prod_{n=1}^{N} Z_n$ is given by

$$Z_T = \int e^{-\beta_\theta \sum_{n=1}^{N} [\ln(2\pi) + \ln(1 - e^{-\gamma_n})+\ln(1-\beta_\theta)]} d\gamma_n.$$

Note that this is only valid in the limit $\beta_\theta \to 0$. Integrating and substituting the resulting $Z_T$ into Eq. (7) (see Supporting information), yields the Helmholtz-like free energy

$$F = N \left[ \left( 1 - \frac{1}{\beta_\theta} \right) \ln(\gamma_n) + \ln(4\pi) + \frac{\ln(1-\beta_\theta)}{\beta_\theta} \right].$$

Eq. (25) is well-defined only for $\beta_\theta < 1$. This indicates that no steady state exists for $\beta_\theta \geq 1$, hinting at an out-of-equilibrium regime\(^3\). The present model belongs to the class of models with logarithmic potentials (see Eqs. (6) and (14)). The existence of a non-normalizable state in certain parameter regimes is a staple of systems with logarithmic potentials\(^2\).

**Collective cell migration patterns for different parameter regimes.** The model was implemented computationally to characterize the model and the effects of the different parameters on the resulting macroscopic behavior. The general qualitative behavior of the model can be observed in Fig. 3. In the regime $\beta_\theta < 0$, cells tend to travel in a single direction after some time has elapsed, similar to the Vicsek model. Conversely, in the $\beta_\theta > 0$ regime, cells are seen to move collectively in transient vortex-like structures, even after long times have elapsed. Qualitatively, the patterns resulting from different parameter combinations are summarized in Table 1. Analyzing simulations, two important phenomena are observed. First, there is a critical parametric regime $\Omega_C := \{ (\beta_\theta, R) : S_3 > 0 \}$ where patterns emerge. Specifically, for low values of interaction radius $R$ no structures can be formed. This indicates that medium-to-long range spread of information is necessary for ordering. On the other hand, for $\beta_\theta > 0$ for large values of $R$, outside of $\Omega_C$, again no patterns occur. This implies that for large interaction radii there is a destructive interference of the travelling information. A second important observation is that patterns do not depend on the choice of $\beta_\theta$ when this is different than zero. For $\beta_\theta \neq 0$, LEUP dynamics divide the population into fast and slow cells. While fast cells are useful for spreading information (and therefore, increasing the effective interaction range), slow cells are necessary for maintaining local ordering. On the other hand, if we fix the initial speed distribution and assume $\beta_\theta = 0$, then we find different patterns emerging as shown in Table 1 (also see in SI). Furthermore, we quantitatively characterized global ordering at long times. The global polar order parameter, given by Eq. (16), for the complete simulation domain,
Figure 3. Simulation snapshots of the velocity field at long times. Arrows show the direction and magnitude of the velocity field. The snapshots were taken after 1000 time steps. 1000 particles were simulated, with an interaction radius of 3, and noise standard deviation of angles and speeds equal to 0.01. Here $g = 1$, $\beta_\nu = -5$ and $\varepsilon = 0$. In (a) the value of the angular sensitivity was equal to 18 while in (b) the angular sensitivity was equal to $-0.25$. Periodic boundaries were employed.

Table 1. Qualitative description of the observed patterns for different angular sensitivity and interaction radius regimes, as well as radial sensitivity. The patterning regime $\Omega_C$ is the blue area in Fig. 4a,b.

| $\beta_\theta$ | Radial sensitivity ($\beta_\nu$) | $R \notin \Omega_C$ | $R \in \Omega_C$ |
|----------------|-------------------------------|---------------------|------------------|
| $< 0$          | $\beta_\nu \neq 0$            | Polar aligned streets of cells | Scattered polar aligned cells |
|                | $\beta_\nu = 0$ (for uniform distribution) | Compact polar aligned cluster | Compact polar aligned cluster |
| $> 0$          | $\beta_\nu \neq 0$            | No order or patterns | Vortices |
|                | $\beta_\nu = 0$ (for uniform distribution) | No order or patterns | Nematic streaming and vorticles |

Figure 4. Pattern formation regimes $\Omega_C$ in the interaction radius—sensitivity plane for positive and negative values of angular sensitivity. In (a) there exists an optimal regime where we can find pattern formation. But in (b) for negative values of beta we can see patterns at smaller values of interaction radius and at smaller values of angular sensitivity.
measures the global degree of polar alignment, or polarization. The global nematic order parameter, given by Eq. (17) for the complete simulation domain, measures the tendency of all cells to align nematically, or along a single axis. These order parameters take a value of one when there is global order, while taking a value of zero when the system is completely disordered. It should be noted that polar order implies nematic order, but the reverse is not true.

Similarly to other velocity alignment models, the model shows an order-disorder transition with increasing noise amplitude and decreasing density (SI figure Fig. 1). More importantly, we observe that in the regime $\beta_0 < 0$ the system also undergoes a transition towards polar order with decreasing $\beta_0$. After the transition, most particles have a similar orientation (Fig. 5a,c). In the regime $\beta_0 > 0$, a phase transition is also observed towards nematic order with increasing $\beta_0$. In this case, however, the nematic ordering is not perfect, as evidenced by the nematic order parameter reaching values of around 0.35 after transition (b) compared to the value of 0.9 of the polar order parameter after transition in the $\beta_0 < 0$ regime. This is further evidenced by the bimodal distribution of orientations with peak separation of approximately $\pi$ radians (Fig. 5d). These simulation results further corroborate our previous theoretical results.

In turn, we study the effect of speed sensitivity $\beta_v$ in terms of phase transitions. We fix the angular sensitivity $\beta_0$, either positive or negative, thus the speed distribution will only depend on $\beta_v$ values. When $\beta_v$ is positive then speed distribution become bimodal and for $\beta_v < 0$ the speed distribution becomes unimodal (see Fig. 10 in supplementary material). Moreover, we show that if the radial sensitivity $\beta_r < 0$ decreases then the average speed increases. On the other hand, for any value of $\beta_r > 0$, the first and the second
moments of the speed distribution cannot be defined, since this is bimodal. Finally, for increasing cell densities, the average speed increases as well (see supplementary Fig. 9).

**A collective migration example of restricted mechanistic knowledge: the spherical bacteria case.** Collective motion of bacteria has been extensively studied and modeled. Most studies have focused on the collective properties of *S. enterica*, *E. coli*, and *M. xanthus*. These species of bacteria are similar since they have a high aspect ratio. It has been shown that volume exclusion, coupled with a high aspect ratio, is sufficient to induce velocity alignment in the system6, and accordingly, ordered clusters of bacteria are observed at high densities.

However, it has been recently shown25 that even spherical *S. marcescens* bacteria do display collective migration (for experimental details please see SI section). The biophysical mechanism whereby spherical bacteria interact with one another must be different from the high body aspect ratio volume exclusion mechanism proposed for elongated bacterial species.

Recently, a combination of biophysical agent-based and hydrodynamics model has been proposed to describe these experiments. In this study the experimental observations were only partially reproduced. Therefore, the biophysical mechanisms underlying collective migration in spherical bacteria are still not well understood. An important aspect to consider is the bacterial speed $v_n$. It was found experimentally27 that bacterial speed followed a Rayleigh distribution, dependent on bacterial density. Collective effects on cell orientations, on the other hand, were studied by observing the vortical behavior of the population28.

To reproduce the experimentally observed Rayleigh distribution for cell speed, we chose the function $g(v_n) = v_n^{-4}$ as shown in34. It is important to note that this term is not impacting the qualitative behavior of the average bacteria speed but only its variance (see SI Fig. 11). Moreover, the interested reader could see the impact of the friction term in the average cell velocity in SI Fig. 12. As shown in Fig. 6, our model qualitatively and quantitatively reproduces both the speed distribution and vorticity behavior of the experimental system.

Interestingly, the behavior of the experimental system was replicated for high values of the sensitivities and quantitively reproduces both the speed distribution and vorticity behavior of the experimental system. Interrestingly, the behavior of the experimental system was replicated for high values of the sensitivities $\beta_1$ and $\beta_2$, and large interaction radii $R$. Our LEUP model not only allows for a quantitative reproduction of the experiments, but also provides insight into the potential biophysical mechanisms. Such values of the sensitivities and interaction radii indicate far-reaching, strong tendencies of bacteria to average their speeds while reorienting and traveling differently from their neighbors. Spherical, rear-propelled particles have been shown to destroy polar order as a result of hydrodynamic interactions26, similarly to our model. Considering that *S. marcescens* is an example of a spherical, rear-propelled particle27, our results agree with previous findings indicating that *S. marcescens* interacts through long-range hydrodynamics28. The long range interaction radius suggests the existence of hydrodynamically induced interaction (which has been suggested by Ariel et. al as well as by other studies27,28) or self avoiding interaction29.

**Discussion**

In this work, we have introduced an off-lattice model of LEUP-induced collective migration, based on the self-propelled particles modeling framework. It was assumed that individuals changed their radial and angular velocity components independently through LEUP. Reorientation is governed by a stochastic differential equation depending on a white noise term and a force arising from an interaction potential.

The exact form of the interaction potential can be very complex, and its specific form is dependent on particular mechnochemical details of the modeled system. While it has been shown that, in general, interactions among individuals can effectively drive the entropy of the entire system towards an extremum point30,31, here we do the opposite. Instead of modeling the interaction potential biophysically, it was assumed that particles followed the LEUP, which dictates that cells change their internal states in order to minimize the uncertainty of the internal states of cells in their surroundings. Although LEUP has been conceptualized to deal with high-dimensional internal states involved in cell decision-making, here we restrict on physical internal states such as speed and orientation. While cell speed was assumed to always minimize uncertainty, there was no assumption made on the cell orientation. Particles are therefore free to reorient either towards or against the gradient of entropy of the orientational distribution of particles in their neighborhood, depending on the sign of the sensitivity parameter, which also dictates the strength of the interaction. The orientational distribution in the neighborhood was assumed to be wrapped Cauchy distributed. Such a distribution facilitates the mathematical analysis of the model. However, the usage of other wrapped distributions do not qualitatively change the general behavior of the model (see SI). Please note that non-parametric methods for estimating entropies without assuming any underlying parametric distributions exist. For instance, such methods employ kernel density estimation, $k$—nearest neighbours or regression methods32.

We show that, when the parameter $\beta_0$ is negative, the model produces steady-state polar alignment patterns. Interestingly, we showed that the classical formulation of Vicsek model is a special case of LEUP. Conversely, when the parameter $\beta_0$ is positive, particles tend to reorient against the mean velocity of their neighborhood. In this regime, the free energy diverges, indicating an out-of-equilibrium parameter regime. This kind of parameter-dependent dichotomy is similarly observed in systems with logarithmic potentials33, involved in processes such as long range-interacting gases34, optical lattices35, and DNA denaturation36. The dichotomy arises from the logarithmic form of the entropy driving interaction in our model. It has been shown that, due to the non-normalizability of the steady state solution, such systems require a time-dependent expression for their analysis34. Therefore, an in-depth theoretical analysis of our model would require a similar multiparticle, time-dependent expression of the angular probability densities.
However, our LEUP migration model may go beyond the observed patterns in past Viscek-type models. In particular, in Barua et al.\textsuperscript{37} we have developed a discrete speed version of our LEUP migration model, where cells can have only zero or a finite speed. This model exhibits Turing patterns, i.e. dynamics clusters of non-motile cells of specific characteristic wavelength, where previously published Viscek-like models cannot may produce moving clusters of swirling cells (e.g. the milling Viscek model) but never static ones.

As a proof of principle, we show that our model replicates the collective vortical behavior of spherical motile particles. Recently, the collective behavior of spherical particles have been modeled as a combination of steric repulsion and hydrodynamic interactions\textsuperscript{38}. Our study has shown that hydrodynamics and steric interactions induce long-range microenvironmental entropy maximization, which coincides with the $\beta \theta > 0$ LEUP regime. This generalizes the type of biophysical mechanisms required to produce vortical patterns.

It should be noted that, while spherical \textit{S. marcescens} bacteria have been modeled biophysically, their collective behavior was partially reproduced\textsuperscript{8}. This hints at an additional biological and/or biochemical interaction between cells. While our LEUP-based model is coarse-grained in terms of specific biophysical/biochemical interactions, it allows for a plausible reproduction of the experimentally observed collective velocity behavior by fitting a only few parameters. The application to spherical bacteria allows us to showcase the potential of the LEUP principle when the precise interaction mechanisms are not known.

As already mentioned, we have made some assumptions to simplify the model. Our model assumes a Gaussian, white noise term in the SDEs. This results in normal diffusive behavior in the absence of interactions. It has been observed experimentally, however, that in some conditions, cells perform Lévy walks resulting in...
superdiffusive behavior\textsuperscript{39}. By changing the distribution or time correlations of the noise\textsuperscript{9,40}, it would be possible to both replicate the non-Gaussian dynamics of single cells, and investigate the effect of single anomalous dynamics on collective behavior.

We have also assumed that particle velocities are the only internal states relevant for reorientation, for simplicity and as a proof of concept of the LEUP principle. However, it is reasonable to think that other states, such as relative position or adhesive state, may be relevant to include when modeling specific systems. This reveals an interesting point in the application of LEUP-driven models which is the selection of the most relevant/dominant internal variables. Although experimental intuition could be the easiest approach, we are currently developing a spatial principle component analysis method that would allow to select the most relevant internal variables using spatial data such as multiplexing biopsies or spatial RNA sequencing.

As stated above, LEUP circumvents the biophysical details of cell migration. The need to model systems of interacting agents without previous knowledge of the biophysical mechanisms involved has sparked at least another agent based model\textsuperscript{41}. In this model, similarly to ours, agents act without a mechanistic rule. Rather, they consider every possible action and penalize those which are not favorable to their internal standards. While both the aforementioned model and LEUP are defined in a similar spirit, modeling under LEUP consists in correctly identifying the relevant internal cellular states for entropy optimization, while in\textsuperscript{42} modeling is concerned with defining suitable penalizations for each possible decision scenario.

LEUP has additional appealing features. For instance, LEUP allows for replicating a plethora of collective migration patterns. In this particular case, we have analytically derived the polar and nematic alignment Vicsek models for LEUP arguments. In this sense, LEUP acts as a generative model for collective migration mechanisms. This is particularly useful upon limited knowledge of such mechanisms, a problem called structural model uncertainty. Another advantage of LEUP is the mapping of biophysical mechanism combination to the migration patterns. In this particular case, we have analytically derived the polar and nematic alignment Vicsek models for LEUP arguments. Although experimental intuition could be the easiest approach, we are currently developing a spatial principle component analysis method that would allow to select the most relevant internal variables using spatial data such as multiplexing biopsies or spatial RNA sequencing.

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**Author contributions**

H.H. and A.B. formulated the mathematical model and conceptualized the project. A.B. and J.M.N.S. performed the analysis and simulations. All authors interpreted the results. J.M.N.S. wrote the manuscript with contributions from all authors. All authors read and approved the final manuscript.

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