Bacterial self-organisation and computation

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1 INTRODUCTION

Researchers in the field of complex systems are often concerned with the study of collections of motile elements, such as fish, birds, insects and cells [15]. These organisms may exhibit characteristic patterns of self-organized collective behaviour such as shoaling [38], flocking [39], swarming [35] and other types of aggregation [37]. We are particularly interested in cells, as the ability to reliably predict and control the dynamical behaviour of cellular colonies has profound implications, both on the study of the biological system per se and also on the potential for their use in a diverse range of computational and engineering applications.

When millions of cells act collectively as a group (as is the case in any colony), a large variety of different patterns arise as a result of how the individual cells respond to the others in their neighbourhood and to the conditions in their environments (which will be different for different individual bacteria, even if they are relatively close to each other) [14] [28] [33] [44]. The physical appearance of these patterns is largely due to cells aggregating together in different ways at different positions. These patterns are the emergent result of local interactions and environmental conditions, and can be usefully viewed as being programmed by both the particular way the cells in question interact, and by the particular way they respond to environmental signals (and, thus, their genetic “program”). Also, these patterns are widely regarded as functional; for example, in stress conditions (such as a toxic chemical in the

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environment), the pattern formed will be ideal for protecting a maximal number of individuals (e.g. by minimising the number of cells without immediate neighbours) while promoting the colony’s “search” for a less harmful environment (by, e.g., extending “arms” of cells which explore the environment). The development of spatially-extended functional patterns purely as a result of local interactions between individuals and with their immediate environment is a form of computation which is extremely common and useful in nature, but which is poorly understood, not well-explored and underexploited in computer science”.

The main purpose of the article is to highlight chemotaxis (cellular movement) as a rich source of potential engineering applications and computational models, highlighting current research and possible future work. We first give a brief description of the biological mechanism, before describing recent work on modelling it in silico. We then propose a methodology for extending existing models and their possible application as a fundamental tool in engineering cellular pattern formation. We discuss possible engineering applications of human-defined cell patterns, as well as the potential for using abstract models of chemotaxis for generalised computation, before concluding with a brief discussion of future challenges and opportunities in this field.

2 BACTERIAL CHEMOTAXIS

The ability to sense and respond to changes in the environment is a fundamental property of living organisms. Because of the basic importance of these capabilities for survival, nature has evolved many mechanisms to facilitate reaction to environmental signals. Such behaviours are often in response to environmental stress (e.g., extremes of temperature or light, changes in pH or the presence of toxic chemicals), and may include changes in gene expression [24], immune response [36] or cellular movement [4, 14, 45]. It is the last response that particularly interests us in this article.

2.1 Chemotaxis mechanism

Chemotaxis is the movement of motile cells [8, 9, 33] in response to concentrations of certain chemicals (see [10] for a general overview). The derivation of the chemo component of the term is obvious; taxis refers to directed movement, as opposed to chemokinesis, which refers to the rate of movement. By “motile”, we mean a cell that is capable of autonomous movement; there are many mechanisms by which a cell may move itself, including “crawling” by shape deformation, the to-and-fro motion of hair-like structures on the cell
surface, or by the whipping of a molecular “propellor” (for a comprehensive overview of motility mechanisms, see [12]). This last method is used by the E. coli bacterium, as it possesses a number of thread-like flagella on its cell surface, which may rotate either clockwise or anti-clockwise under the probabilistic control of the cell. When individual flagella are rotated anti-clockwise, they form a single, coherent “bundle”, the movement of which propels the bacterium forwards. A bacterial “run” is terminated when the flagellar bundle disintegrates as a result of its rotation being reversed. In this situation, the bacterium tumbles for a short time, a situation defined as “erratic motion without net transition” [33]. When resuming directed movement after a tumble, the bacterium moves off in a new, effectively random direction. When on a run, the bacterium can therefore implement a biased random walk by manipulating the probability of either remaining running by continuing to rotate its flagella in the anti-clockwise direction, or of terminating the run by reversing the direction, causing it to tumble. In this way, a cell may explore its environment, making biased decisions about whether or not to continue moving in the current direction. In the absence of any chemical signals, individual bacteria perform a purely random walk [33]. However, a cell’s choices may be influenced by chemotactants; chemicals in the environment which may either attract or repel the bacterium. Bacteria respond to a chemotactant by sensing the chemical’s concentration gradient and then reacting appropriately (for example, a bacterium may attempt to move up the gradient of an attractant by increasing the probability of running while the detected gradient increases). One immediate question that arises from this description is how a bacterium only 2-µm long can sense a chemical gradient with a decay distance of (perhaps) many millimetres. In 1972, Macnab and Koshland [40] proposed a bacterial “temporal sensing mechanism” (i.e., a memory), which would allow bacteria to make comparisons between past and present environmental conditions. The existence of this mechanism was later confirmed by Segall, Block and Berg [42]; their work demonstrated that E. coli possess a 4-second memory that, combined with single-molecule sensitivity, allows these bacteria to respond to environmental cues in an extremely sophisticated fashion.

2.2 Chemotactic pattern formation

For many years, it was believed that chemotaxis in motile bacteria served only to maximise their chances of encountering nutrients. Consider the situation where a colony is inoculated at a single central point in a dish of gel containing nutrients. The colony will swarm outwards in either a single ring
or a series of concentric rings [14], responding to spatial gradients of nutrient attractants. However, it is now clear that, under conditions of stress (e.g., exposure to hazardous byproducts of respiration, such as peroxides), \textit{E. coli} themselves generate attractants, becoming moving sources of chemical signals [14, 23, 45]. This signal generation facilitates \textit{inter-cellular communication}, leading to the collective formation of spatial patterns of spots, stripes, moving bands and other complex structures [33]. Example of such structures, generated by a cousin of \textit{E. coli}, are depicted in Figure 1.

The function of each type of pattern depends very much on the external pressure on the colony. Mittal \textit{et al.} describe one scenario [33], where \textit{E. coli} form well-defined spotted clusters. Such patterns might emerge due to the presence of \textit{xenobiotics} – chemicals not usually found in the organism being studied (for example, drugs or carcinogenic compounds). In this situation, the bacteria may need to respond quickly by producing detoxifying enzymes [18], high concentrations of which may only be produced in dense cell clusters. “Clustering by chemotactic aggregation enables bacteria to generate a robust spatial structure and maintain a localized population through the temporal sensing of the concentration of a single species of molecule secreted by them...” [33].

Ben-Jacob and Levine describe a different type of pattern formation in \textit{Paenibacillus dendritiformis}, bacteria that move over hard surfaces by excreting lubricant chemicals on which they “skate” to move around [6]. A level of “wetting” sufficient for collective movement requires the contribu-
tions of a large number of individuals. In situations where food is scarce, the required population density is unsustainable, which creates a trade-off situation for the colony as a whole. This is resolved by the population forming a branching structure; in each branch there are just enough bacteria to generate sufficient lubricant without exhausting the food supply. Ben-Jacob and Levine report that individual bacteria constantly adjust their lubricant properties and production rate to generate different branch structures, according to the environmental hardness and nutrient levels. They argue that this “self-engineering” may be thought of as a form of collective constraint satisfaction, where the necessity of movement is balanced against the resources required for such migration.

3 MODELLING CHEMOTAXIS

There exist several different in silico approaches to studying collective cell behaviour. Many previous studies model the cell population and its environment at a macroscopic level, by describing the system using Partial Differential Equations. The classic model of this type is due to Keller and Segel [30] (see [29] for a comprehensive review). Numerical modelling approaches may, on the other hand, simulate chemotaxis at either the macroscopic population level, or the microscopic level of individual cells. Such models are generally based on finite difference/finite element methods (see [27] for a full list of relevant references). Lattice-Boltzmann models [27] represent bacteria and chemoattractants on a regular grid, and have been used with some success. However, such models fail to capture all aspects of real bacterial behaviour, and their application in unstructured (i.e., irregular) environments is still unclear. Finally, simulation studies that model individual cells (or groups of cells) have recently been developed [7, 22, 25, 28, 41]. It is this last approach that concerns us here, and we consider in particular the communicating walkers model, first proposed by Ben-Jacob et al. [7]. The original model considers the following biological features: (1) diffusion of nutrients, (2) bacterial movement, (3) bacterial reproduction, (4) local communication.

Diffusion of nutrients is handled by solving the diffusion equation for a particular nutrient concentration on a triangular lattice. Bacteria are represented by “walkers”, each of which is a coarse-grained sub-population of the colony. Each walker is an agent [48] described by its location and an internal degree of freedom (or “internal energy”), which affects its activity. Walkers lose energy at some constant rate, and may consume nutrients where available. When the walker’s internal energy reaches zero it becomes stationary.
When food is sufficient, a walker’s internal energy increases, and when it reaches some threshold it divides (reproduction). Stationary walkers produce a communication chemical at some fixed rate (in an attempt to drive other bacteria away) and each of the active walkers consumes the chemical at some fixed rate. The movement of the active walkers is therefore a random walk with a bias along the gradient of the communication field. Results obtained using this simulation framework have been encouraging, generating many of the patterning phenomena observed in the laboratory [5, 16, 17].

3.1 Extensions to the model

We now propose certain extensions to the basic communicating walkers model. These are intended to improve the realism of the model, by incorporating consideration of such factors as generation of nutrient chemoattractant gradient by nutrient consumption, release and consumption of chemoattractant, release and consumption of chemorepellant, rheology of medium matrix and bacterial reproduction. In detail, the additional features we propose are (1) Substrate structure. Standard agar gel is made up of aggregates of helices that give it a porous structure, which may produce effects that are unrelated to inter or intracellular processes. The substitution of agar with materials such as carboxymethyl cellulose, removes this structure, giving a more homogeneous environment and removing the contribution of gel structure to pattern formation. The facility to vary this parameter would allow the possibility of studying the effect of both gel strength and structure. (2) Nutrient spectrum and distribution. Previous work has assumed uniform nutrient distribution. It is clear that non-uniform distribution will affect pattern formation in complex non-linear ways, so we must consider the possibility of “doping” the media surface with regions of nutrient of arbitrary complexity. (3) Response to toxins. The Budrene and Berg patterns [14, 23] have been shown to be due to the accumulation of respiration by-products such as hydrogen peroxide when growing on organic acids as sole carbon and energy source. The formation of micro-colonies aids in the more rapid destruction of hydrogen peroxide by endogenous catalase. We can vary the speed of accumulation and destruction of hydrogen peroxide by doping and incorporation of exogenous catalase in the medium, and examine the effect on pattern formation. (4) Genetic components. There are many mutants available in E. coli, that affect chemotaxis, nutrient use and toxin degradation. The effect of these mutations on pattern formation can be assessed within the model we propose, and would allow us to vary parameters as dictated by the discrete in silico modelling results.
3.2 Benefits of this approach

There are now many compelling reasons for considering this individual-based approach to the study of bacterial pattern formation. If we require additional biological realism, adding extra components to a system of differential equations will quickly render it intractable if we require a numerical solution. However, using the discrete approach, the addition of an extra parameter or feature adds, at worst, a constant time factor to the time required by the algorithm. The biological realism is central, as it may well be that case that a novel component of the system that has not previously been considered proves to be significant to the success or failure of the model. Traditional approaches generate biologically feasible patterns, but we seek a degree of resolution that may not be achievable without using a more fine-grained model. In addition, it is not clear that these models are capable of generating regular patterns of both “rings” and “dots”, which may be required by the anticipated applications of the patterns (see Section 4). Differential equation models do not treat each cell or subset of cells as individuals behaving in a stochastic fashion. Our proposed approach uses a discrete approach that models the behaviour of a relatively fine-grained subset of cells which will yield the group-level pattern that might emerge from an actual experiment. The degree of granularity of the simulation can be easily modified by altering the number of bacteria represented by each walker. In addition, bacteria such as *E. coli* move at a constant rate, so it is easy to “clock” the simulation and ensure global synchronisation. By making the simulation spatially explicit and discrete (and therefore relatively easy to parallelise), we can easily take advantage of the computational power offered by cluster machines and the grid.

Support for this cell-centred view of biological modelling has gathered momentum in the past few years. As Merks and Glazier argue in a recent article [32], “...in many aspects of biological development, cells’ inner workings are irrelevant: what matters are the cell’s biophysical properties, the signals it emits and its response to extracellular signals.” While we would caution that their argument is from a developmental perspective, and that we may not be able to fully treat the cell as a “black box” for our purposes, we believe that the fundamental message is sound – that cell-centred models will play an increasingly significant role in both generating hypotheses about and explaining collective cellular behaviours.
3.3 Fundamental issues

Some of the fundamental questions that we seek to address in the future are:

1. What level of complexity at the individual (bacterial) level is required to
generate the observed complexity at the collective level? Additionally, how
much of the observed complexity at the collective level is a reflection of envi-
ronmental rather than individual complexity?

2. Is it possible to simulate the
growth and movement of bacterial populations such that the complex patterns
they generate in vitro can be specified in advance in computationally feasible
times?

3. If achieved, how may the associated abstract computational architectures,
as well as the directed self-assembly of bacteria in vitro be applied
in various domains? The first question is of fundamental importance to re-
searchers in the field of biocomplexity. The second and third questions are
important if the great commercial potential for this work can be realised. We
propose using the model not only to further our understanding of collective
cellular behaviour and chemotaxis, but as a framework for directed pattern
formation. The fundamental idea is to develop a biologically valid simula-
tion of bacterial self-organisation, via continual and rigorous testing against
biological reality. This laboratory work is absolutely crucial to the develop-
ment of a reasonable model. Without this feedback from the biologists,
the model will be nothing more than a very crude approximation of reality.

We may then allow a user to specify a desired final pattern of bacteria, which
may be useful for a particular application such as bio-sensing. Creating
this pattern by top-down placement of individual bacteria is possible, but
may be unfeasible in the environments of practical interest, so we seek to set
up the system such that the pattern self-organizes from an initial configura-
tion or parameters (cell density, initial placement of nutrients, gel structure,
etc.). The parameter space in which suitable configurations are located is
huge and multi-dimensional, so we may use an evolutionary algorithm (EA)
metheuristic to search it more efficiently. EAs are well-known for highly
effective performance in large combinatorial, numeric and mixed combinator-
ial/numeric spaces (as in the case of the configuration space here), and recent
work has already showed their efficacy in searching a space of in vitro bacte-
rial colony compositions. We could use the EA to negotiate the space of
initial configurations and parameter settings, and the fitness function (which
will be multiobjective) will simulate the model and then calculate a vector
of similarity measures between the desired and final pattern. This forward
simulation approach is vital, because the desired pattern formation can take
several days in the laboratory. By running several million simulations in this
time, each with different parameter settings, we can efficiently identify ap-
propriate values for parameters that interact in complex non-linear ways and minimise the amount of experimental laboratory work required. We now consider possible practical implications of the extended model that we have just described.

4 POSSIBLE APPLICATIONS

Models of chemotaxis have already been successfully applied to the training of artificial neural networks [13, 19], multimodal function optimisation [34], the design of aerofoils [34], the control of robots for environmental monitoring [20], and the modelling of pedestrian evacuation processes [41]. It is clear that this phenomenon has inspired a powerful optimisation strategy that is competitive with existing nature-inspired methods. However, what is lacking from the majority of these chemotactic models is a consideration of how the highly-localised interactions of the bacteria generate long-range biologically-advantageous patterns. By including such mechanisms, we may further enrich the possibilities for distributed bacterial optimisation.

Perhaps more significantly, the physical patterns of bacteria will find many applications in the biological and engineering domains. Self-assembly will clearly play a strong role in determining the future direction of (bio) nanotechnology [47]. Our model would have implications for any application that requires spatially registered cellular engineering. These may include the development of novel biomaterials, the design of biosensors, and tissue engineering [2]. We also anticipate its eventual application to the construction of structured microbial consortia [1], whose functioning in applications such as wastewater treatment and bioremediation is dependent on their precise structural ordering. Recent work has achieved small-scale, local directed pattern formation, using genetically-modified bacteria [3]. By incorporating genetic factors with bacterial movement, our simulation framework would provide a powerful tool for the emerging field of synthetic biology [21, 26].

5 CONCLUSIONS

In this position paper we have outlined an ambitious vision of harnessing the complexity of bacterial communication and self-organisation for the purposes of human-defined computation and engineering. Rather than simply observing and modelling complex pattern formation, we seek to control and utilise it. Quite apart from the massive potential benefits to biology, we wish to understand a real and complex biological system so thoroughly that we can
control it and harness its computational power to provide a wide range of potential applications. In doing so we expect to establish a highly complex computational architecture for local-and-environmental-interaction based pattern generation, from which we may extract several simplifications and extensions with varieties of combinatorial properties. All of these we expect to find applicable in many of the several high-impact sciences and technologies emerging across the board which involve (or will benefit from) the need for functional, adaptable and useful self-organisation of collections of locally interacting and environmentally responsive units (e.g., softbots in a network topology, self-organising and responsive biosensors, biofilters, nanodevices, MEMs, smart skin technologies, functional and programmable materials, cellular electronics, environmental control, bioremediation). We now call for a concerted research effort in this direction.

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