MODELLING CHEMOTACTIC MOTION OF CELLS
IN BIOLOGICAL TISSUE WITH
APPLICATIONS TO EMBRYOGENESIS

THESIS SUBMITTED IN ACCORDANCE WITH THE REQUIREMENTS OF THE
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BY

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General Introduction

Motivation

Perhaps one of the most amazing events that occurs in nature, is in the emergence and growth of biological life. Emergence speaks of the well-coined phrase Primordial ooze from which the chemical building blocks of life first gave rise to the complicated molecular structure of Deoxyribonucleic acid (DNA), that has the mind boggling task of encoding every chemical and physical attribute and trait of the organism for which it is encoded. This incredible feat of nature is only equalled by the ability of single fertilized cell (zygote) to undergo a seemingly magical transformation through enlargement, growth and change to give rise to a fully formed animal (or plant). The study and body of knowledge of this latter process is called Developmental Biology, and it seeks to define and explain all of the intricate sub-stages and bio-chemical, molecular and physical processes along the time-line of this transformation, that is from fertilization to birth, hatching or germination and beyond.

One might consider, and quite reasonably, that the variety of different processes leading to the development of a complete biological organism would be so vast as to render the problem untenable. Indeed the almost inconceivable amount of genetic information contained within the nucleus of the simplest of cells would seem to corroborate this assumption. However when one takes a more holistic view, we can see that the development of any complex biological organism can be reduced to a set of five distinct processes, all of which are orchestrated to define structures from a body of cells. Viewed in this light the generation of any complex multi-cellular organism, be it small or large, must involve: cell-division, differentiation, pattern formation, change in form and growth [1].

To mediate and orchestrate these different processes during the development of the embryo are a enumerable number of bio-chemicals that are produced within the cells that can diffuse into the surrounding environment, activating (and de-activating) inter/intra-cellular signalling pathways that trigger further productions and possibly one or more of the processes suggested above. One such case of this, and which is of particular interest in this thesis, is in the role of morphogens in the growth of vertebrate embryos, where it is known that interacting morphogen gradients can give rise to spatially stable concentrations [2] that are known to be involved in organ growth [3], primitive streak formation [4] and the extension and patterning of the primary body axis [5, 6, 7].

In this thesis we are considering one such problem involving these mechanisms/processes, during the primary body axis extension in the chick embryo. During this phase of development the early brain is beginning to form and the central nervous system (CNS) is beginning to extend unilaterally in a posterior direction defining the main anteroposterior (head to tail) body axis; in simple terms one may see this as the generation of the spinal cord and surrounding structures. Extension of this axis is known to be orchestrated by a small cellular structure located at the posterior-most tip of the extension, encompassing what is known as the primary organising centre in the chick embryo: Hensen’s node. This structure
including the node is known to move independently/autonomously of the rest of the embryo and as it does so the cells in the region are growing and proliferating, and ultimately differentiating and leaving this region to literally fuel the axial extension.

This broad description leads us to the heart of our thesis, and which will preoccupy the rest of this dissertation. We postulate that the motile behaviour of the group is as a result of biochemical gradients to which the group is attracted toward areas of highest concentration or towards areas of lowest concentration of some as yet unnamed morphogen. That is we assume that the group moves as a result of a chemotaxis. Furthermore, the growth and subsequent differentiation of cells exiting the group, contributing to the growth of the CNS, are also regulated by the same morphogen. Therefore we propose that a singular bio-chemical mechanism can account for the motile and growth behaviour observed during CNS extension.

**Thesis Outline**

In the course of this dissertation we shall set out to show that a singular, elegant biochemical mechanism is sufficient to describe the complicated process of primary body axis extension during the development of the central nervous system, as orchestrated by the organizing centre, Hensen’s node. Throughout we make some large assumptions concerning the composition of the node and the nature of its migration and growth, and the bio-chemicals that we assert are responsible for the migration due to chemotaxis. However we also draw upon experimental observations and unique experiments performed during the course of this project to support any of these assumptions, but inevitably there are aspects of this work that are still open to conjecture. In any event the thesis outline is as follows:

**Chapter 1:** In this current chapter we review the mathematical and biological background of the thesis, with reference to significant contributions and developments. We will also consider the models and numerical methods we will employ and their respective derivations.

**Chapter 2:** In this chapter we abstract the problem to a one-dimensional continuous model to investigate the chemical dynamics in and around the group. We consider chemical concentrations that can either be produced within the group or by a surrounding population of cells. The cell structure of the group is also investigated in terms of homogenous and heterogeneous compositions, for example where one sub-population produces the chemical and another reacts. To aid this analysis we consider permutations using analytic methods and numerical simulations using our proprietary simulator, BioChemSim.

**Chapter 3:** Using the results of Chapter 2 we next consider extending the problem to two-dimensions with a proprietary implementation of the Cellular Potts Model (CPM) we call BioCellSim, taking a more qualitative approach to corroborate the one-dimensional findings of
Chapter 2: The distinguishing feature of the CPM is that now the group is represented as a recognisable cellular structure composed of individual cells. Thus while we seek to corroborate previous findings, we also reveal features and results not present or accounted for in one-dimension.

Chapter 4: In this chapter we introduce a experimental work in association with R. D. del-Coral, where we illustrate through numerical simulations and analytical models a plausible mechanism based on interacting morphogen gradients that can control migration of Hensen’s node and growth, proliferation and differentiation of the extending vertebrate primary body axis.

Chapter 5: In this chapter we present our conclusions and a discussion of the findings presented throughout this work, demonstrating strengths, weaknesses and outstanding areas of possible research.
Chapter 1

BACKGROUND REVIEW

1.1 Background Review

In this chapter we review both the mathematical and biological backgrounds to the problems outlined during the rest of this thesis. At a broad level we consider the field that encompasses the development of biological life, with an overview of the historical events that have construed to give us the ever growing field that is Developmental Biology. In particular we shall give a review of developmental context, or if you will the physiological setting for our project, the model organism Gallus gallus (the chicken). Mathematically the background is given in terms the development of our governing equations, described by reaction-diffusion systems, necessarily implying the emergence of steady-state or travelling wave fronts, which are assumed to give rise to pattern formation and/or positional information in the cellular structure constituting our physiological problem. In addition we shall review the numerical techniques we applied and the implementation of our proprietary software: BioChemSim and BioCellSim.

1.1.1 Developmental Biology

A Brief Historical Overview of Developmental Biology

Studying the development of biological life from the zygote to the fully fledged animal is a remarkable process, a process which has preoccupied mankind and sparked his imaginations to conjure theories that could explain such a seemingly magical process. Indeed in a time dating back several thousand years to the ancient Greeks, clearly a time without microscopy, one could be forgiven to bare any theory that could, at least in some part, explain how a seemingly invisible egg could give rise a chicken, mouse or even human.

The earliest of these theories was proposed by the great Greek philosopher Aristotle around 600 BC who postulated two, and necessarily competing theories, to explain how this transformation might occur. These two ideas, he termed epigenesis and preformation, sought to explain the formation of the early embryo. Epigenesis describes the progressive development of the different structures in the embryo in time, while preformation the idea that the embryo was a miniature version of the fully formed animal that merely grew in size. It is known that Aristotle preferred the theory of epigenesis, and clearly given our current understanding he was right. However it was preformation that was to dominate science for the two thousand years (attributed to the prevailing ideas of creationism supported through religious doctrine) until a resurgence in the 18th century, notably by the German physiologist Caspar Friedrich Wolff, attributed as one of the founders of modern embryology, who indicated that the organs of the embryo are derived from different layers of undifferentiated cells; layers which we now refer to a Germ Layers.
This cell-centred conceptualisation was to become very important, and indeed it is the prevailing theory in biology that living organisms are complex organisations of cellular structures, a theory that developed during the 19th century, notably by German botanist Matthias Schleiden and physiologist Theodor Schwann. The idea that all living organisms are seen as diverse cell populations, and more importantly the observation that the egg itself was but a single highly specialized cell, became known as Cell Theory, and it marked the death knell (at least within the scientific community) for preformationism.

While Cell Theory had given a plausible explanation for the structure of living things being entirely composed families of cell types making up the whole organism, it raised the question of how these different families of cells came about? One of the fundamental tenets of Cell Theory is that new cells are created from old ones, of course beginning with a single zygote. If this is true then how do different families of cells arise, that is by what mechanism we explain creation of bone, skin, liver or any type of cell for that matter?

An answer to this question was proposed by the German botanist August Weismann, who suggested that as cells enlarged and divided to create two daughter cells (cleavage), the contents of the parent cell, he called determinants, would be unequally divided amongst the daughter cells. Thus seen as a continual process, this would give rise to many distinct cell types in each successive cleavage, a process that was termed mosaic development. However this theory was shown to be incorrect by German biologist Hans Adolf Eduard Driesch. During experiments on sea urchin embryos he showed that at the first stage of cleavage of the zygote, if one of the first daughter cells was removed, then a still complete larva would result, albeit slightly smaller in size. This clearly implies that each daughter cell of the zygote contains within it all that is required to develop a complete organism, and indeed implies the contents of the zygote are replicated and distributed equally to its daughters, clearly contradicting Weismann’s claim.

The implication of Driesch’s experiment is that there must be some form of self-regulatory mechanism at work, which necessarily implies there must be some form of communication between cells in the embryo. Put another way, if we assume that the two original daughter cells of the zygote were left to their developmental fates, the lineages of these daughters would lead to a fully developed embryo. However if we remove one of these lineages/daughters, then the remaining lineage must reorganise to compensate for this loss. Therefore the fates of individual cells, in terms of the future body plan of the organism, are not controlled by the cells themselves but by some other extracellular control, thus implying some form of communication between cells.

Evidence of this communication was ultimately found during a famous transplant experiment performed by Hans Spemann and Hilde Mangold [8], that lead to the principle of induction, which states that tissues or cells can induce or direct the development of cells around them. To demonstrate this Spemann and Mangold had identified a small region on the newt embryo that seemed to be controlling its development. They took a small graft from this region and transplanted it into a second newt embryo, and startlingly it developed a second partial embryo. They named this region an organiser for obvious reasons,
and today it is referred to as the Spemann-Mangold Organiser and for their work they received a Nobel Prize.

Much of the work that continued through the rest of the 19th century and indeed has continued to the current day, was/is concerned with the connection between genetics and development and principally how the expression of genetic information in terms of proteins could control this. Initially it was conceived that genetics and development were separate functions of the organism. Genetics was concerned with aspects of inheritance of particular features and traits from the parents that manifested in the child, such as demonstrated by Gregor Mendel and his famous pea experiments. On the other hand development was only concerned with the development of the embryo, and differentiation of cells forming the early germ layers. It wasn’t until the finding that cells produced proteins that are expressions of genes, and that these expressions could activate or inhibit the production of proteins in other cells or tissue, that can ultimately control the development cells and ultimately the embryo. Thus the modern definition of developmental biology is the development of an organism from fertilization to birth, via the progressive development and refinement of cellular structures, controlled and governed by the expression of genetic information referred to as gene action.

**Chick Embryo As A Model Of Vertebrate Development**

The challenge in choosing any developing biological organism for analysis is driven by the requirements of the research that specifies the characteristics of an organism one is interested in. The selection of such an organism is based on several factors, such as rate of reproduction, amenability to genetic modification, physiological interference such as grafting or transplantation and/or visualisation of early of development, lifecycle behaviours and in some cases historical reasons. For example Escherichia coli (E. Coli) has long been used in molecular genetics in the study of bacteriophages in studying gene regulation and gene structure, as it has very short life-cycle implying one can see over very short time periods successive generational effects of genetic manipulation. At a larger scale, investigation of cell-level behaviour and signaling, such as in Dictyostelium discoideum, is used due its restricted number of cell types and it simplistic life-cycle. Sea urchins and amphibians such as frogs where chosen because of their ease of acquisition and robustness to experimentation over long time scales of their development. Amongst the vertebrate family, perhaps the most useful organism is the chicken (Gallus gallus). Since most of its development takes place after laying inside the egg, it can be easily visualised with lamp and microscope by careful remove of a section of egg shell. Further it is very robust during interference, where the entire embryo can be transplanted from the egg and its growth observed in-vitro. These organisms, by either historical fate or amenability to experiment in some context, are known as model organisms.
Key Developmental Stages In Chick Embryo Development

Vertebrates share very similar developmental stages after fertilisation, from initial growth, emergence of important organising centres to large scale cell migrations leading to changes in form known as morphogenesis. In the chick embryo these changes take place geographically on the surface of the egg yolk, which can be seen macroscopically as a small opaque ring in any fertilised egg (Figure 1-1).

**Cleavage.** The first stage of development is concerned with growth as the newly fertilised cell (zygote) undergoes rapid cell divisions during the stage known as cleavage, that increases the number of cells in the embryo to form the a densely packed, multi layered group of cells called the Blastodisc, containing approximately 60,000 cells. At this point a cavity appears underneath the Blastodisc, separating it from the yolk to develop a subgerminal cavity, which is translucent and is given the name the area pellucida. Surrounding this translucent area is an opaque ring of cells that define the outer border of the embryo, given the name area opaca. As development proceeds a new layer of cell develops over the surface of the yolk called the hypoblast, that together with the topmost layer (now given the name the epiblast), completely surrounds the subgerminal space. It is on this surface, the epiblast, that the embryo proper will develop.

**Gastrulation.** At this point a critical feature of development emerges, that is the first visible indicator the embryo is viable and marks the beginning of the gastrulation stage. Gastrulation marks large-scale cell migrations and morphological changes in the structure of the embryo that culminates in the formation of the main body axes, and the central nervous system giving the familiar impression of an early vertebrate embryonic foetus. At the beginning of gastrulation, at the posterior (lower edge) of the embryo between the area opaca and area pellucida, a small crescent-like formation of cells appears called Koller’s Sickle. From within the sickle a small condensation of cells appears centrally within the sickle that begins to migrate anteriorly (upward or head-wise) over the surface of the area pellucida. As it does so it produces in its wake a small invagination in the surface of the epiblast called the primitive streak, and it is this structure that defines the early anteroposterior (head to tail), and by implication, dorsoventral (back to front) body axes. Hensen’s node, together with the primitive streak, forms the primary organizing centre within avian embryos, analogous to the Spemann-Mangold organiser in amphibians. Shortly after the
formation of the primitive streak it begins to regress, together with Hensen’s node posteriorly, with cells from the lateral epiblast converging on the streak and ingressing through it into the subgerminal cavity as mesenchyme cells (loosely connected), eventually forming the future mesoderm and endoderm germ layers (see Figure 1-2).

Figure 1-2: Illustration of the early stages in vertebrate embryonic development through to the formation of the primitive streak (Wolpert 2007). In the early stages the embryo undergoes rapid cell divisions to form the Blastodisc of approximately 60,000 cells (A). As development continues a subgerminal cavity appear separating it from the yolk (B), over which a unicellular of cells forms the hypoblast (C). The hypoblast together with the newly formed upper ceiling, the epiblast, now completely enclose the subgerminal cavity (D). At the posterior marginal zone a small crescent-like group of cells appears (D) the gives rise to a small condensation of cells known as Hensen’s node that begins to migrate over the surface of the epiblast to near the centre of the embryo, and as it does it leaves in its wake an invagination in the surface of the epiblast (E). Cells on the lateral epiblast converge on the streak and ingress through it to give rise to the future mesoderm and endoderm germ layers. The primitive streak, together with Hensen’s node make up the primary organiser of the developing organism, and define the primary body axes.

Neurulation. The regression of the primitive streak is coupled with the formation of the notochord, which one can refer to as the early backbone, which defines the so-called neural plate on which the early nervous system will develop. At the anterior end the notochord cells laterally convergence at the centre of the plate and begin to fold to form the early head process (encompassing the early brain) and the neural tube. Collectively one can visualise this process as the formation of the early central nervous system, with the early brain forming at the anterior and the development of the “backbone” being fuelled and organised by the regression of Hensen’s node.

Organogenesis And Beyond. Much of the rest of the development is concerned with development of internal organs and eyes, and largely signified by growth and enlargement until hatching.
1.1.2 Mechanisms Of Cell Migration

Despite the large amount of experimental observations and data, there is still relatively little known about the forces and mechanisms that control the migrations of cells during early development of the chick embryo. The ability of a single cell to move is well understood, in terms of its physiology and biomolecular signaling that motivates such movement. However in more complicated systems, such as during gastrulation in the chick embryo during formation of the primitive streak, a number of factors could be brought to bear. In such system there are large and diverse cell populations contained within densely packed medium exhibiting cellular flows, where it is not clear which cells are actively moving and which are being moved passively [9]. That is, some cells may be actively moving in response to some signal, but in such a dense medium, there will inevitably other cells that are moving as a result of adhesive bonds with active cells, literally pushing and pulling surrounding cells. Further the forces and sources that cause such motion are also in question. However there seems to be agreement that in general, there are some fundamental mechanisms that describe cell motion/migration: cellular intercalation, cellular growth, apical constriction and chemotaxis.

Apical Constriction

In a unicellular layer of cells that are assumed to share reciprocal bilateral adhesion with their neighbours, apical constriction can be seen as the contraction on the upper or lower surface of the cells. The surface that contracts is referred to as the apical layer, and it contracts by the action of compression in the actin filaments that make up the cytoskeletal structure of the cell walls. As the apical layer contracts the upper basal layer begins to expand and a local curvature is observed.
Figure 1-3: Apical constriction. (A) Illustration of a unicellular layer of cells undergoing constriction of the apical layer by causing surrounding cells move inwards as the local curvature increases. (B) Example of the apical constriction within the sea urchin (Lytechinus variegates) from Morrill and Santos 1985; C and D after Lane et al. 1993) as seen by scanning electron microscope. (C-D) Illustration of the structure and process leading to apical constriction.

This mechanism is clearly demonstrated within the early gastrula of the sea urchin (Lytechinus variegates), where the invagination is referred to as blastopore. However while it does describe evidence on cell movements, it is not considered to be involved in larger scale migratory behaviour of cells.
Cellular Intercalation

This is the process by which two groups of cells become spatially integrated. In one case there are two layers of cells stacked on top of each other, then by some mechanism the layers breaks the adhesive bonds between the cells their respective layers and begin to merge together thereby causing a radial cell flow or migration in the plain of intercalation. This type intercalation is referred to as radial intercalation, which typically reduces three-dimensional bodies of cells into a unicellular two-dimensional layer (Figure 1-4A).

The second mechanism, Mediolateral intercalation, is similar to that of radial intercalation, however two layers appear to have a common one-dimensional interface, like sliding two pieces of paper together on a table (Figure 1-4B). This kind of intercalation occurs in many species during gastrulation, and is sometimes referred to as convergent extension, and is typical of axial extension in one-dimension such as in zebra fish development.

Chemotaxis

Chemotaxis is the behavioural response of biological organisms to exhibit a motile response in the presence of a chemical gradient, quite literally taxis meaning to move and chemo pertaining to chemical. There is an abundance of research within the literature for chemotaxis as the underlying mechanism that drives numerous processes in embryological development, too many to mention within this thesis. However there are select model cases that illustrate such chemotactic behaviour dramatically, notably the aggregation of slime-mold namely Dictyostelium discoideum, travelling bacteria Escherichia Coli and more centrally to this dissertation in the regression of Hensen’s node during gastrulation in the chick embryo (see Section 1.2.3).
Figure 1-5: Panel A illustrates both simulated dynamics (sub-labelled A,B,C,D) and actual images of aggregations and mound formation (sub-labelled E,F,G,H). In sub-panel A the ameobae (yellow) are randomly distributed on a substrate and a spiral wave of cyclic adenosine monophosphate (cAMP) is initiated with colour-coded concentrations from high (red ~0.8) to low (0.0 blue). Sub-panel A-B illustrates streak formation as the amoebae aggregate towards the centre of the spiral as they are periodically excited by the wave eventually leading to the formation of the mound illustrated in sub-panel A-D. B: Current Opinion suggests that during gastrulation the primitive streak dynamics are controlled via chemo-repulsion by FGF8 and chemo-attraction of FGF4 giving rise to observed cell flows in and around the streak.

Dictyostelium discoideum is a type of amoeba that become chemotactically active as they struggle to find nourishment in their environment. To solve this problem they begin to secrete a signaling molecule, cyclic adenosine monophosphate (cAMP), into the environment, which is picked up by surrounding amoebae that amplify and relay this chemical signal and begin to travel to its source. In time the amoebae aggregate at this source and develop into a multicellular slug in an attempt to find new feeding grounds. Bacteria, such as E. Coli being a much simpler organism, has a more fundamental mechanism to respond to reduction in nutrients and it has long been known to move freely within a substrate in search of oxygen and mineral nutrients by travelling up the gradients of these nutrients to their source. The mechanisms underlying the regression of Hensen’s node is not so easily discernible, but it is generally accepted that the production of certain fibroblast growth factors in and around the node are involved in its motility [4, 10, 9, 11].
1.2 Mathematical Modelling in Developmental Biology

1.2.1 Introduction

Much of the early work involved in developmental biology centered on observation, were given a model organism physiologists and biologists would make detailed notes and drawings of the various stages of development. From these detailed observations hypotheses were formed on the likely mechanisms or processes that drive the intricate stages of development. In the absence of genetic sequencing and/or advanced microscopy to identify regulatory networks of bio-molecules or bio-physical mechanisms, invariably these hypotheses would be investigated by interference with the normal developmental processes, such as by grafting, transplantation or even complete removal of whole sections of the embryo. Over the millennia a vast amount of data had been collected and catalogued from such experimentation, however much of the mechanisms involved in development biology still elude us to this day, primarily because of the incredible complexity that underlies even the most primitive biological organism.

It would appear that the classical approach of reductionism leads to us to an intractable problem. That is since the underlying complexity is so great, inevitably we would be lead to correlating an almost insurmountable number observations over vastly different spatiotemporal scales. This has lead to the development of fields such as System Biology that draw upon interdisciplinary research across the sub-fields of biology to draw conclusions at a more holistic level. Underpinning much of this research is the application of computational and mathematical models in a manner that captures the essential processes that give rise observable behaviour. In this respect, and quite ironically, mathematics does follow a reductionist approach, however, and as remarked by Murray [12]:

"The aim ... is not to derive a mathematical model that takes into account every single process because, even if this were possible, the resulting model would yield little or no insight on the crucial interactions within the (biological) system"

Collectively the field that encompasses the mathematical applications and tools that are used to give insight into complex biological systems is Mathematical Biology. The introduction and application of mathematics to understanding biological systems was first proposed by D’Arcy Thompson [13], who expounded that the underlying geometry of form and growth could be explained in terms of principles of mechanics and physical laws. Investigation into population dynamics and bioenergetics by Lotka [14] is regarded by many as the first text on Mathematical Biology, with robust models of predator-prey systems that are still used to this day to demonstrate the principle dynamics of such systems. In developmental biology the discovery of the Spemann-Mangold organiser in amphibians, and the consensus that cell differentiation leading to pattern formation was a function of positional information due to morphogen gradients [15], was first indicated by Turing [16] where he coined the term “gradient” in describing reaction-diffusion (or activator-inhibitor) systems that could describe pattern formation by fast and slow diffusing morphogen concentrations.
Within the remit of this thesis, we consider such systems in the context embryological development in the chicken. During early development of the central nervous system, the anteroposterior body axis extends, orchestrated and regulated by the organising centre Hensen’s node. It is known that this self-regulating behaviour allows for the node to move independently of the rest of the embryo, inducing pattern formation of the extending body axis controlled via intra-cellular signaling by the morphogen families of fibroblast growth factors (FGF) and retinoic acid (RA) [5, 7]. The central question we seek to answer, and indeed model, is what mechanism accounts for this regulation that allows the node to move? As we have suggested in the previous section there several mechanisms that can account migrations of tissue or cells within the embryo, however we propose that a single chemotactic mechanism is responsible. Thus at the heart of this project is the problem associated with understanding the origins and dynamics of experimentally observed morphogens, and the resultant dynamical behaviour of the node in terms uniform motion.

For the remainder of this section we shall consider some of these mathematical models as they contribute to the project at hand, namely embryonic development within the chick egg, with a view to understanding methods of cell migrations in around Hensen’s node leading to its regression.

### 1.2.2 Modeling Morphogen Concentrations and Pattern Formation

The term morphogen, as coined by Turing [16], speaks of a bio-chemical that can be involved in change of form and shape within developing biological systems. These changes can be associated with reorganisation of cellular structures whereby cells migrate into new positions, typically during development of the early body plan of the organism, but continue throughout the developmental process. Morphogens are also known to be involved in patterning that is not necessarily connected with reorganisation, but with molecular cell differentiation, whereby a group of cells are exposed to the distribution of one or more morphogens that trigger differentiation of cells within the group according to level of the morphogen they exposed to. The resultant type after differentiation is dependent on the developmental history of each cell, so that coupled with the exposure to the morphogen, spatial patterns in cell type will emerge.

At a broader level one can see that there is a fundamental link between cell differentiation and migration, and the spatial distribution in morphogen concentration profiles in producing spatial cell patterns. In the sections that follow we shall briefly review some of these mechanisms.
**French Flag Model Of Pattern Formation**

The interpretation and subsequent differentiation according to the spatial positioning within the morphogen concentration, was termed positional information by Wolpert [15] and is characterised in his French flag problem as illustrated in Figure 1-6.

![Figure 1-6](image)

Figure 1-6: A group of cells exposed to a morphogen concentration can differentiate according to the exposure at specific gradients or thresholds of the morphogen. Assume a homogeneous linear arrangement of cells of width \( L \). At gradients in the concentration, \( G_1 \) and \( G_2 \), differentiation can give rise to spatial patterns of different cell types corresponding to bands of the colours blue, white and red at exact position over the group, \( x_1, x_2 \) and \( L \).

In the classic setting Wolpert suggested that a simple linear passive distribution of a morphogen over the group of cells of length \( L \), that at thresholds in its concentrations ( \( T_1, T_2 \)), could signal differentiation of cells, defining differentiation threshold boundaries. Cells in the group with a concentration greater than the threshold \( T_1 \) for \( 0 < x < x_1 \) would differentiate to become blue, while below \( T_1 \) for \( x_1 < x < x_2 \) would become white and below \( T_2 \) for \( x_2 < x < L \) red, giving a simplistic model:

\[
\begin{align*}
D \frac{d^2 c}{dx^2} &= 0; & c(0) &= c_1; & c(L) &= c_2, \\
\end{align*}
\]

with \( c \) the concentration of the morphogen, \( D \) the diffusion coefficient and \( c_1 \) and \( c_2 \) the values at the boundaries representing source and sink concentrations respectively, implying a linear solution (Figure 1-6A):

\[
c(x) = \frac{c_1 - c_2}{L} x + c_1,
\]

(1.2)

While such a formulation allows for size invariance, implying we can double the size of the group, \( L \) to \( 2L \), and similar pattern would result, albeit slightly larger, such distributions do not correlate well with experimental observations. However a similar result can be achieved by assuming a non-linear distribution with active diffusion (Figure 1-6B):

\[
\begin{align*}
D \frac{d^2 c}{dx^2} - k_1 c + p(x) &= 0; & \left. \frac{dc}{dx} \right|_{x=0} &= 0; & \left. \frac{dc}{dx} \right|_{x=L} &= 0 \\
\end{align*}
\]

(1.3)

where the concentration decays proportional to its own concentration with rate \( k_1 \), and its production is given by the characteristic function:

\[
p(x) = \begin{cases}
  k_2, & 0 < x \leq a \\
  0, & a < x < L
\end{cases}
\]

(1.4)
with $k_2$ an arbitrary positive production constant defined in the domain of size $a$. In this setting it assumed that the cells within the group are sensitive not to threshold value of the morphogen, but its gradient ($G_1, G_2$). However it should be clear that this second model does not possess size invariance, as the distribution of the morphogen in this case is only dependent on the diffusion coefficient $D$ and the decay rate $k_1$. This implies that if we double the size of them medium in this case, the relative horizontal dimensions of each part of the flag would differ.

In either case the French flag problem illustrates quite elegantly how relatively simplistic mechanisms can bring about complex pattern formation, due to the dynamics of a single morphogen. In the next section we shall see that adding a secondary morphogen to the system can lead spatially stable patterns in the morphogens.

**Pattern Formation in a Two Component System**

In a landmark paper “The Chemical Basis of Morphogenesis” Alan Turing [16] proposed that spatially stable patterns could occur in a two-morphogen system if they possessed different diffusion rates. Such systems have come to be known as reaction-diffusion systems. In principle there are two morphogens, one the activator, $a$, and the second the inhibitor, $h$. The activator exhibits an autocatalytic reaction that self-enhances its own production, while simultaneously catalysing the production of the inhibitor. The inhibitor also has an autocatalytic reaction but in addition acts to down-regulate, or catalyses decay, of the activator. In its most general form, such a system can be given by the following system in one-dimension:

\[
\frac{\partial a}{\partial t} = D_a \frac{\partial^2 a}{\partial x^2} + F(a, h) \quad (1.5)
\]

\[
\frac{\partial h}{\partial t} = D_h \frac{\partial^2 a}{\partial x^2} + G(a, h) \quad (1.6)
\]

The coefficients $D_a$ and $D_h$ are constants representing the rate of diffusion, and $F(a, h)$ and $G(a, h)$ the associated kinetics terms, that assumed to be both a function of the activator and inhibitor concentrations, collectively describing the interdependent rates of production and decay.

Thus the essences of Turing’s idea was that instability would occur in an otherwise stable homogenous distribution if $D_h \gg D_a$. This seemingly simplistic idea has been shown to be able to explain a plethora of pattern forming phenomena in biological systems, depending on how one prescribes $F$ and $G$. In linear setting $F$ and $G$ can describe the mutual activation and inhibition that can create spatially stable concentration of interacting gradients, that can give rise to structures such as those illustrated in the French flag model. By introducing non-linearities in $F$ and $G$, a much richer set of behaviours can result.
For example the well-known Fitzhugh-Nagumo (a qualitative generalization of the Hodgkin-Huxley model) model involves excitable or fast variable, typically given as a non-linear cubic equation, and a slow refractory variable, typically linear as a model of squid axon induction of excitation. The form of these equations have become widespread in the modeling of reaction-diffusion system due to the variety of travelling, propagating or stationary patterns can result (Figure 1-7) such spiral waves, pulsating or stationary spots and spatially distributed patterns (See proceeding section Figure 1-6).

1.2.3 Chemotaxis

Chemotaxis is a phenomenon whereby somatic cells, bacteria and other single-cell organisms direct their movements according to certain chemicals in their environment. History of chemotaxis research is indeed well known. Migration of cells was detected from the early days of the development of microscopy but erudite description of chemotaxis was first made by TW. Engelmann and WF. Pfeifer in bacteria and H.S. Jennings in ciliates. The significance of chemotaxis in biology and clinical pathology was widely accepted in the 1930s [17]. Chemotaxis is important for bacteria to find food (for example glucose) by swimming towards the highest concentration of food molecules, or to flee from poisons (for example phenol). In multicellular organisms, chemotaxis is critical to early development (for example the movement of sperm towards the egg during fertilization) and subsequent phases of development (for example the migration of neurons and lymphocytes) as well as in normal function. In addition, it has been recognised that mechanisms that allow chemotaxis in animals can be subverted during cancer metastasis.

In eukaryotic chemotaxis, the mechanism employed is quite different from that in bacteria. However the sensing of chemical gradients is still a crucial step in the process. Due to their size, prokaryotes cannot detect effective concentration gradients, therefore these cells scan and evaluate their environment by a constant swimming (consecutive steps of straight swims and tumbles). In contrast to prokaryotes, the size of eukaryotic cells allows the possibility of detecting gradients, which results in a dynamic and polarised distribution of receptors. Induction of these receptors by chemo-attractants or chemo-repellents results in migration towards or away from the chemotactic substance [17]. Study of eukaryotic cell migration and chemotaxis are processes which are fundamental to cell growth, survival and death. Chemotaxis in particular is essential during embryonic development, immune cell function and
cancer metastasis. It has high significance in the early phases of embryogenesis as development of germ layers is guided by gradients of signal molecules [18].

The use of mathematical techniques for classification and understanding of the ever-growing amount of experimental data and possible help in designing new experiments is now more profound than ever. Recently, mathematical modelling played an growing role in biological studies in general and in particular, in developmental biology. Mathematical modelling in developmental biology has an important role in helping us discover biophysical mechanics driving the development. It is a unique tool which allows a rigorous check of hypotheses concerning these mechanisms as they emerge from the experimental observations. Several mathematical models of chemotaxis were developed depending (among which) on the type of

a) migration (for examples the basic differences of bacterial swimming, movement of unicellular eukaryotes with cilia/flagellum and amoeboid migration),

b) assay systems applied to evaluate chemotaxis (to see incubation times, development and stability of concentration gradients) and

c) other environmental effects possessing direct or indirect influence on the migration (lighting, temperature, magnetic fields, etc.).

Other publications written in genetics, biochemistry, cell physiology, pathology and clinical sciences can also incorporate data about migration or especially the chemotaxis of cells. A curiosity of migration research is that among several works investigating taxes (for examples thermotaxis, geotaxis and phototaxis), chemotaxis research shows a significantly high ratio, which point to the underlined importance of chemotaxis research both in biology and medicine [17]. For chemotaxis, a mathematical description for it requires a model to describe the cells ability to sense a gradient of ambient chemotactant and its interaction with a physical model of cell migration. It is also now appreciated that it is important to model the feedback from the evolving cell shape and the intra and extra cell signalling pathways which lead to directed cell motion. The computational challenge therefore involves the solution of partial differential equations (PDEs) on evolving surfaces where the computed solution state is used to derive movement and changes in cell shape.

Further chemotaxis is modelled by non-linear equations or systems of equations. Contrary to linear models, which have a limited range of possible solutions, non-linear models can be used to reproduce virtually any kind of known dynamics in concentration fields of morphogens. This is especially true if more than one morphogen is considered. In general these are represented by so called reaction-diffusion equations (It is used to derive the equation for the flux of cells whose motion is affected by variations in the ambient concentration of certain chemicals) [7].
An individual cell path can result in an average cell flux which is proportional to the macroscopic chemical gradient. If we let \( b(x) \) denote the density of cells centred at \( x \) and \( J(x) \) the net flux of cells per unit time in the direction of increasing \( x \). Then the dependence of cell density \( b(x; t) \) on position and time is described by the differential equation:

\[
\frac{\partial b}{\partial t} = -\nabla \cdot J(x) \tag{1.7}
\]

where the vector flux \( J \) is given by:

\[
J = -\mu \nabla b + \chi \nabla c \tag{1.8}
\]

The first term is the diffusion term, describing the non-chemotactic, random motion of cells, and the second term describes the chemotactic response.

Chemotaxis has been used in the detailed study of the developing population of Dictyostelium discoideum (Dd) amoebae. This biological organism cooperates with and shows striking social behaviour when they are deprived of food. The starving cells then communicate by means of chemical signal to synchronise their otherwise random and unorganised movement.

The molecular machinery for cell motility is currently best understood in Dd. This is because it is an organism which spends most of its life as a chemotactic amoeboid phagocyte but which also uses chemotaxis to form a multicellular organism during subsequent stages of development. In this process of transformation, there is both production of and a chemotactic response to cyclic adenosine 3’ – monophosphate (cAMP); the result is aggregation [19].

Aggregation of Dd amoebae is an example of a phenomenon whereby the wave of excitation can change the properties of excitable media and cause the formation of spatial patterns. The monolayer of the starving amoebae is an excitable medium which conducts excitation waves of the intracellular mediator i.e. the cAMP. Since cAMP is a chemotactic attractant for the amoebae, the waves of cAMP cause motion of the amoebae. As a result of this motion amoebae are organized into streams which usually form branching radial multicellular structures. There are two major types of cAMP sources forming aggregates: a point source and a spiral wave. Figure 1-8: shows streams which were induced by a spiral wave of cAMP.

![Figure 1-8: View of aggregative structure formed by a starving population of Dictyostelium discoideum due to chemotactic response of the amoebae as illustrated in Figure 1-5 [19].](image-url)
The process of aggregation of Dd amoebae was numerically studied in a continuous model and thus the reaction-diffusion model [19] was proposed for the simulation of the process. The model is based on FitzHugh-Nagumo-type equations for cAMP waves and a continuity equation for amoebae motion. The process of aggregation induced was simulated by a periodic point source and by a spiral wave. It was shown that an aggregation pattern is formed as a result of front instabilities due to dependence of wave velocity on density of amoebae. This instability can also result in formation of wave breaks and generation of spiral waves.

For calculations, the following model was used:

\[
\begin{align*}
\frac{\partial r}{\partial t} &= \frac{g - r}{\tau} \\
\frac{\partial g}{\partial t} &= D_g \Delta g + c^a(f(g) - K_r r) \\
\frac{\partial c}{\partial t} &= D_c \Delta c - \nabla(cV(r) \Delta g)
\end{align*}
\]

(1.9)

The first two equations are a Fitzhugh-Nagumo model which describes the propagation of cAMP waves. \(g\) represents the extracellular concentration of cAMP and \(r\), the refractory period. Instead of ordinary cubic function, \(f(g)\), in the second equation, the piecewise linear function is used instead:

\[
f(g) = \begin{cases} 
-S_g, & g \leq 0 \\
K_g(g - a), & 0 < g < 1 \\
-S(g - 1), & g \geq 0
\end{cases}
\]

(1.10)

\((S\) is infinite, \(f(g)\) is only defined on the interval \(0 < g < 1\)). It was suggested that in normal conditions, the production and decay of cAMP are proportional to the cell density \(c^a(a = 1)\).

The third equation in (1.9) describes the chemotactic motion of amoebae. \(c\) is the local concentration of amoebae, and \(V(r)\) is their motility. In the model (as \(r \geq 0\)), \(V\) reaches its maximum at \(r = 0\) and decreases to 0, with increasing \(r\). Biologically, this means that cells move if they are not refractory (i.e. not able to respond to additional stimulation).

From an initially random distribution of amoebae, the formation of the aggregation pattern will occur. They will form the pattern of branching streams. Therefore a necessary condition for stream formation is non-uniformity in the initial distribution of amoebae density. If we perform a simulation, but with an initially uniform distribution of amoebae, the streams will not formed. Amoebae collect in the stimulated area and form a circular spot with high density. This mechanism of stream formation is associated with the fact that the velocity of the cAMP waves depends on the local density of the amoebae.
The reaction-diffusion model proposed describes fairly well the aggregation process in natural population of Dd. The aggregation pattern obtained by numerical simulation looks similar to the pattern in amoebae populations (Figure 1-8). In addition, the technical features which makes Dd amoebae attractive as a model includes [9]:

a) The cells exist as a homogeneous population in culture,

b) They can be induced by physiologic stimuli to undergo normal morphogenesis in vitro thus permitting direct observation of the role of chemotaxis in organogenesis,

c) Cells can be grown in suspension culture to high density to generate kilogram quantities of material for biochemical analysis,

d) Amoeboid cells are haploid and are readily manipulated by molecular genetic techniques and

e) The physiological response to chemotactic stimulation is synchronous in a cell population and can, therefore, be correlated with biochemical measurements.

1.2.4 Numerical Methods

Much of the analysis of this thesis involves the development of two numerical/computational tools: BioCellSim and BioChemSim. BioCellSim, as its name suggests is a two-dimensional cell-centered computational model based of the Cellular Potts Model of Glazier & Graner [20], which we implemented and modified to satisfy our modelling needs. BioChemSim on the other hand is an entirely novel model we derived for simulation of chemical and chemotactic dynamics. In this section we wish to briefly set out the numerical methods we employed in these models, primarily in terms of reaction-diffusion. It should be noted that we will not be discussing here the CPM per-se, rather we will only outline the numerical implementations of the chemical models we use. In all of our models we used a forward-time centered-space (FTCS) explicit Euler finite differencing scheme.

Numerical Reaction-Diffusion Equations

In all numerical modelling of chemicals we can see, both one and two-dimensions, that our diffusion equation is given as a standard parabolic equation with added kinetics expressed in (1.11) as $F(x, t)$. From the point of view of deriving a numerical model, the only difference between one and two dimensions is in the expression for the Laplacian differential operator $\Delta$, where in one dimension it takes the form $\Delta = \partial^2 / \partial x^2$ and in two dimensions $\Delta = (\partial^2 / \partial x^2 + \partial^2 / \partial y^2)$. The kinetics term can be considered equivalent in both models and as it is linear it does not affect the derivation of the numerical form, thus we can state the governing equation for both one and two dimensional models as:

$$\frac{\partial u(x, t)}{\partial t} = D\Delta u(x, t) + F(x, t); \quad x \in (-\infty, \infty), t \in [0, \infty)$$

(1.11)

Our aim is to find an explicit numerical representation of (1.11), that is, to find representations for both its spatial and temporal derivatives. The temporal derivative gives us the rate of change of chemicals in time, whereas the spatial derivative gives us the rate of diffusion of the chemicals with respect to space.
In other words the spatio-temporal derivatives can be seen as taking approximations for small perturbations in both space and time. In this way we can see that we can represent derivatives as Taylor series expansions about these small differences/perturbations, which can be visually illustrated as stencils in Figure 1-9.

![Stencils](image)

Figure 1-9: Numerical stencils for one and two-dimensional explicit Euler, forward-time centered-space differencing scheme in one dimension (A) and two dimensions (B).

Considering the spatial variation, ±Δx, in one-dimension of the chemical concentration u(x₀) due to diffusion at a discrete numerical grid point x₀, we can approximate this by the Taylor expansion:

\[
    u(x₀ ± Δx) ≈ u(x₀) + u'(x₀)Δx + \frac{1}{2} u''(x₀)(Δx)^2 + O(Δx)^3,
\]

and

\[
    u(x₀ ± Δx) ≈ u(x₀) - u'(x₀)Δx + \frac{1}{2} u''(x₀)(Δx)^2 - O(Δx)^3.
\]

If we now take the superposition of (1.12) and (1.13) and rearrange we find:

\[
    u''(x₀) ≈ \frac{u(x₀ + Δx) + u(x₀ - Δx) - 2u(x₀)}{(Δx)^2},
\]

is an approximation to the second order derivative with respect to space, x, where it should be clear that we can perform an equivalent derivation for a second spatial dimension, y, as:

\[
    u''(y₀) ≈ \frac{u(y₀ + Δy) + u(y₀ - Δy) - 2u(y₀)}{(Δy)^2}.
\]

Then by superposition of (1.14) and (1.15) we can an approximation to the Laplacian differential operator in two-dimensions as:

\[
    \Delta u(x, y) = \left( \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \right) u ≈ \left( \frac{u(x₀ + Δ, y₀) + u(x₀ - Δ, y₀) - 2u(x₀, y₀)}{Δ^2} \right)
    \]

\[
    + \left( \frac{u(x₀, y₀ + Δ) + u(x₀, y₀ - Δ) - 2u(y₀)}{Δ^2} \right)
\]

where for simplicity we have assumed that the grid is isotropic, and in one-dimension as:

\[
    \Delta u(x) = \left( \frac{\partial^2}{\partial x^2} \right) u ≈ \left( \frac{u(x₀ + Δ) + u(x₀ - Δ) - 4u(x₀)}{Δx^2} \right).
\]
The representation of the temporal derivative can be derived by equivalent means and be shown to be of the form:

\[
\frac{\partial u}{\partial t} \approx \left( \frac{u(x_0, y_0, t + \Delta t) - u(x_0, t)}{\Delta t} \right).
\]  

(1.18)

Now if we substitute (1.16) and (1.18) in (1.11), and make a symbolic change of \( u_{x,y}^t = u(x, y, t) \) then we have:

\[
\frac{u_{x,y}^{t+\Delta t} - u_{x,y}^t}{\Delta t} = \frac{u_{x+\Delta, y}^t + u_{x-\Delta, y}^t + u_{x,y+\Delta}^t + u_{x,y-\Delta}^t - 4u_{x,y}^t}{\Delta^2} + F(x, y, t)
\]  

(1.19)

which after simplification we find the FTCS finite differencing approximation to (1.11) in two dimensions:

\[
u_{x,y}^{t+\Delta t} = u_{x,y}^t + \frac{\Delta t}{\Delta^2} \left( u_{x+\Delta, y}^t + u_{x-\Delta, y}^t + u_{x,y+\Delta}^t + u_{x,y-\Delta}^t - 4u_{x,y}^t + \Delta^2 F(x, y, t) \right)
\]  

(1.20)

and in one-dimension:

\[
u_{x}^{t+\Delta t} = u_{x}^t + \frac{\Delta t}{\Delta x^2} \left( u_{x+\Delta}^t + u_{x-\Delta}^t - 2u_{x}^t + \Delta^2 F(x, t) \right)
\]  

(1.21)

**Stability of FTCS Numerical Scheme**

The stability, or instability, of the numerical schemes given by (1.20) and (1.21) is seen as a function of the numerical error associated with the scheme and is attributed to the rate at which spatial information contributes to each grid point from time \( t \) to time \( t + \Delta t \). In each time step, \( \Delta t \), in (1.20) or (1.21) the contributing information for a grid point, \( u_{x}^{t+\Delta t} \), comes from \( u_{x}^t, u_{x+\Delta}^t \) and \( u_{x-\Delta}^t \). If neglect the kinetics terms \( F(x, t) \) in (1.21) and re-write it as:

\[
u_{x}^{t+\Delta t} = (1 - 2s)u_{x}^t + su_{x+\Delta}^t + su_{x-\Delta}^t.
\]  

(1.22)

where \( s = D\Delta t/\Delta^2 \), the standard condition for stability is given by \( s \leq 1/2 \) for (1.22). Formally we can derive the condition for stability by the method of Von-Neumann stability analysis, where we assume that coefficients of the difference equation are varying so slowly in space and time as to make them approximately constant. In this way we can assume that the eigenmodes of the difference equation are all of the from

\[
u_{x}^t = \xi^t e^{ikx}\Delta
\]  

(1.23)

where \( k \) is the wave number and \( \xi = \xi(k) \) is a complex number where the importance lies in the fact that the difference between modes is linear, given by successive integer powers of the complex number \( \xi(k) \). Therefore the numerical scheme can be shown to be unstable if there are exponentially growing modes, that is \( |\xi(k)| > 1 \).


Substituting (1.23) into (1.22) we find:

\[
\frac{\xi^{t+1} e^{ikx\Delta} - \xi^t e^{ikx\Delta}}{\Delta t} = \frac{\xi^t e^{ik(x+\Delta x)\Delta} + \xi^t e^{ik(x-\Delta x)\Delta} - 2\xi^t e^{ikx\Delta}}{\Delta^2} \Delta^2
\]

\[
\xi^{t+1} - \xi^t = \frac{\Delta t}{\Delta^2} (e^{ik\Delta x} + e^{-ik\Delta x} - 2)
\]

\[
\frac{\xi^{t+1}}{\xi^t} = 1 - \frac{\Delta t}{\Delta^2} (e^{ik\Delta x} + e^{-ik\Delta x} - 2)
\]

(1.24)

now using the identities:

\[
\cos(kx) = \frac{e^{ik\Delta x} + e^{-ik\Delta x}}{2} \quad \text{and} \quad \sin^2 \left(\frac{k\Delta x}{2}\right) = \frac{1-\cos(k\Delta x)}{2}
\]

(1.25)

we can rewrite (1.25) as

\[
\xi(k) = 1 - \frac{4\Delta t}{\Delta^2} \sin^2 \left(\frac{k\Delta x}{2}\right)
\]

(1.26)

where we have written \(\xi(k) = \xi^{t+1}/\xi^t\) which is referred to as the amplification factor, and therefore to satisfy \(|\xi(k)| > 1\) we require:

\[
\left|1 - \frac{4\Delta t}{\Delta^2} \sin^2 \left(\frac{k\Delta x}{2}\right)\right| \leq 1 \quad \Rightarrow \quad \frac{\Delta t}{\Delta^2} \leq \frac{1}{2}
\]

(1.27)

for the numerical scheme to remain stable, and this condition is known as the famous Courant condition.

In essence (1.26) states that for any specification of the \(\Delta\) we are required to ensure that \(\Delta t\) is set such that the condition in (1.27) holds.

**Accuracy Of Numerical Scheme**

We shall only note here the method we have used to ensure the accuracy of our simulations and we will only be considering details that warrant a justification for our simulations. The accuracy, or more accurately, the inaccuracy of the numerical scheme, can be seen to be caused two different problems: boundary size and incorrect specification or spatio-temporal scales used.

Boundary size is important in that in a diffusive scheme, if the boundary of the domain of the problem are set too close, then the numerical simulation will not (literally) have enough space to for the solutions to saturate properly. In the context our problem, if boundary is too close to the domain of transcription, given in one-dimension as a segment and in two dimension a disk, then the chemicals will have a limited into which they can diffuse, and so the resultant chemical profiles will not be accurate. To account for this problem, we can simply ensure that the boundary size is large enough, however in practice it is not readily clear what “large enough” translates to in any given simulation. Our solutions was quite simple, when obtaining results from simulations we always sought to check the results by doubling the size of the medium to determine if the boundary was having any appreciable effect.

Spatio-Temporal scales were covered in the previous section, however when we include kinetics terms into the standard heat/diffusion equation, we must account for this contribution by rescaling both time in space, such that condition (1.27) is satisfied. A simple check we used to determine is the scales we used were accurate, was note the relation between the spatio-temporal scales: if we double the spatial step, we should reduce the time step by a factor of 4. So in our numerical results to determine accuracy we
would increase the segment/disk twice, double the spatial step and divide the temporal step by a factor of four and adjust boundaries dimensions if need be. If we found a less than 3% difference in numerical results we would accept this as being reasonably accurate.

1.3 The Cellular Potts Model

The CPM has its origins in mathematical physics as a mathematical model used to study phase transitions that occur in physical systems. While not self-evident why research into phase transitions allows us to model cellular phenomena, we shall see in the following sections how (and a common practice in science) the development of a model in one discipline can have broad applicability in others. In terms of the CPM this originates with the work of Ernst Irving, in the study of magnetic properties of ferromagnetic materials. Therefore we shall begin at the beginning, so to speak, and show the development of the Ising Model into CPM we will be using for the rest of this chapter.

1.3.1 The Ising Model

The Ising model was developed by Ernst Ising for his 1925 PhD Thesis, in an attempt to derive a mathematical model whose solution could predict spontaneous magnetization in ferromagnetic materials such as iron, nickel, cobalt or their alloys. This spontaneous change in behaviour, or “phase transition” in these materials from paramagnetic to ferromagnetic was/is of significant interest, as the production of magnets on an industrial scale clearly has large ranging applications throughout all sectors of industry and society. While there was a great deal of intuition for what phase transitions were, and even how they occur, there was no mathematical tools/models that could demonstrate this analytically. Thus Ising’s work was a first attempt to derive a closed-from analytical expression that could describe conditions, using weak assumptions, on system parameters for such phase transitions to occur. In Ising’s original work in one-dimension (referred to as one-dimensional chains, the term Ising Model was coined by Periels [21]) he failed to demonstrate such transitions which, in very broad terms, he incorrectly concluded that similar results would occur in higher dimensions. It wasn’t until nearly a decade later through the work of Periels [21] and Kramers & Wannier [22] that it was shown that a phase transition can occur in two-dimensions, and several years later Onsager [23] derived an analytical solution to the model in the absence of an external magnetic field, the so called zero-field.

Phase transitions are intuitive physical/chemical processes and they occur in numerous settings throughout nature, characterised by the change in the qualitative behaviour of some substance/material in response to a change in a parameter in the system. A common example to most of us is the transition of water into vapour or ice, dependent on a critical temperature value of the system. Changes or transitions occur abruptly and typically manifest as discontinuities in the governing mathematical system or one of its derivatives. The so-called Ehrenfest classification, named after 20th century Dutch physicist Paul Ehrenfest, classifies transitions by the order of the derivative of the system that first displays such a discontinuity. For example water freezes and boils at temperatures $T = 0^\circ\text{C}$ and $T = 100^\circ\text{C}$, that
notwithstanding variations in surrounding atmospheric pressure or temperature (an idealised system) mark precise conditions for the transition from liquid to solid or liquid to vapour. These changes manifests as a discontinuous change in the density of the water, which is given to be the first derivative of the free energy with respect to the chemical potential, and therefore a first-order transition.

For Ising’s work, magnetism, we know from classical electrodynamics that spinning electrically charged bodies will produce a magnetic dipole with poles having equal magnitude and opposite polarity. In terms of atoms the concept of “spin” is given to be a collective analogy of the angular and orbital moment of electrons about the nucleus. Clearly if two locally and equally interacting atoms have opposite spin, necessarily implying opposite polarity, then the net magnetic moment will be zero. Therefore if a material is to exhibit a useful measurable magnetic field, the configuration of atoms within the material must favour interactions between atoms with equal polarity/spin; the greater the degree of alignment, the greater the net force. Practically this is achieved by re-organising the materials microcrystalline structure, by heating the material and exposing it to a powerful magnetic field. As the magnetic field is reduced, and assuming the temperature of the material is at a critical (constant) value (less than the materials Currie point), the material will undergo a ferromagnetic phase transition to exhibit spontaneous magnetisation. In this sense the magnetisation of the material is given by a first derivative of the free energy with respect to the applied magnetic field, which is a continuously increasing function as the temperature is lowered below its Currie point. However the magnetic susceptibility of the material is given by the second derivative, which exhibits a discontinuity in the onset of spontaneous magnetisation, and therefore is described as a second-order phase transition.

To model the problem Ising made the assumption that localised interactions between spin states can give rise to long-term correlative behaviour that could help to predict the onset of a ferromagnetic phase transition. To aid this assumption, Ising suggested that the material be discretised onto a lattice, $L$, of uniformly distributed sites, with each site representing the spin state of an individual atom. The state of each spin is given by the state variable $\sigma_i$ for which there can be $i = 1 \ldots N$ possible states, each taking on a value $\sigma_i = \pm 1$, representing the physical assumption that each site can be only one of two possible states: spin-up or spin-down. In Ising’s original one-dimensional case, $L$ can be visualised as line of equally spaced $\sigma_i$, however it can be easily extended to higher physical dimensions such as grid in two-dimensions of cube in three (Figure 1-10).
Local interactions on $L$ can now be seen as what one might term as nearest neighbour interactions, or first order interactions, that occur between sites that are at most one lattice site away from the site of interest $\sigma_i$. It is possible to consider higher order interactions, but for the rest of this work only first order interactions will be considered. If we let $D$ be the dimension of the $L$ then we can see that for a given $\sigma_i$ there will be $2D$ interactions (or local atomic bonds) forming a stencil of interacting sites as illustrated in Figure 1-11.

These interaction stencils are taken to be the smallest unit of energy computation and they form the basis of the Hamiltonian, $\mathcal{H}$, of the system. That is $\mathcal{H}$ gives us function from which we determine the current energy state of $L$. At this point we should note that this highlights Ising’s largest assumption for the model, as generally speaking a Hamiltonian gives the total energy state of a system, whereas here the assumption is that only first order interactions contribute to the final state. Therefore for $D = 1$ Ising introduced the following Hamiltonian:

$$\mathcal{H}(L) = - \sum_i \mathcal{E}(\sigma_i \sigma_{i-1} + \sigma_i \sigma_{i+1}) - \sum_i \mathcal{F}(\sigma_i - 1 + \sigma_i + \sigma_{i+1})$$

(1.28)

which expresses the summation over all lattice sites on $L = (\sigma_1, \ldots, \sigma_N)$.

The first sum in (1.28) represents the first order interaction at each site proportional to the parameter $\mathcal{E}$, which prescribes the strength of the interaction energy, and the second sum the influence of the magnetic field $\mathcal{F}$ on the interacting sites. Recalling that $\sigma_i = \pm 1$, then if we consider $\mathcal{E} < 0$ we can see that the system will favour anti-aligned or higher energy states, whereas for $\mathcal{E} > 0$ lower energy states or energy minimization, the latter being favourable for spontaneous magnetisation. Clearly $\mathcal{F} > 0$ as the
orientation of the magnetic field should tend to align states, and therefore should contribute to the minimization of energy.

As suggested earlier, an important aspect of phase transitions is that changes in the qualitative behaviour of the thermodynamic system manifest as abrupt, discontinuous changes, which in the case of ferromagnetic transitions is described as spontaneous magnetisation. We can infer from this description that transitions are not deterministic in nature, moreover they are characterised by stochastic processes, which brings us to the central object used by the Ising model, the partition function from statistical mechanics:

\[ Z(\beta, \mathcal{E}, \mathcal{F}, N) = \sum_{\pm 1} e^{-\beta \mathcal{H}(\mathcal{L})}, \quad (1.29) \]

where parameter \( \beta = 1/k_B T \) with \( T \) the absolute temperature and \( k_B \) is the Boltzmann factor that relates temperature and energy at the microscopic particle level, and the minus sign suggest that lower energy states are preferred.

In general (1.29) is the sum over all configuration microstates on \( \mathcal{L} \) which for a lattice of \( N \) sites equates to \( 2^N \) possible spin state configurations. Its use becomes apparent when we consider the probability of one such a spin state configuration, \( \mathcal{L}_0 \), that is a specific assignment of states \( \forall \sigma_i \in \mathcal{L}_0 \), then the probability of \( \mathcal{L}_0 \) occurring is given by the so-called Boltzmann probability distribution:

\[ P(\mathcal{L}_0) = \frac{e^{-\beta \mathcal{H}(\mathcal{L}_0)}}{Z}. \quad (1.30) \]

More generally (1.30) encodes statistical information about the thermodynamic features of the system, such as heat, magnetism, etc. For example the expected internal free energy is given to be the weighted probability:

\[ P(\mathcal{L}_0) = -\frac{1}{Z} \sum_{\pm 1} \mathcal{H}(\sigma) e^{-\beta \mathcal{H}(\sigma)}. \quad (1.31) \]

which we can see can be expressed as the derivative with respect to the temperature of the system \( \beta \)

\[ P(\mathcal{L}_0) = -\frac{\partial \log Z}{\partial \beta}. \quad (1.32) \]

This brings us to the central question in the Ising model: if we define the free energy per lattice site as

\[ F(\beta, \mathcal{E}, \mathcal{F}) = \lim_{N \to \infty} \log Z(\beta, \mathcal{E}, \mathcal{F}, N), \quad (1.33) \]

where \( N \to \infty \) is called the thermodynamic limit, then Ising sought a closed form analytical solution for \( F(\beta, \mathcal{E}, \mathcal{F}) \). The reasoning for this is that if phase transitions do occur, then they would manifest as discontinuities in \( F \) or one of its derivatives. However there is no guarantee that such a limit exists.

For our purposes this essentially completes the description of the Ising model in one-dimension, and we will not be considering this problem in any further detail. Next we shall consider the contributions of Potts in generalising this model.
1.3.2 The Potts (Clock) model

In this section we acknowledge the work of Renfrey Potts in generalizing the Ising model in his 1952 PhD thesis under the supervision of C. Domb. The Potts Model, as it has now come to be known, was first suggested to Potts by his supervisor Cyril Domb for his PhD thesis. The principle was that the Ising model could be generalized to more than two spin states. That is, can phase transitions be found in lattice where there are \( q \) different spin states? As stated earlier, the Ising model for \( q = 2 \) was solved for a zero field \( \mathcal{F} = 0 \) \cite{23} and indeed the Potts Model is equivalent to the Ising Model for this value of \( q \). The problem was to consider the spins as \( q \) equally spaced spins confined to the plane or as Domb suggested, \( q \) spins equally spaced around a circle, with angles

\[
\theta_n = \frac{2\pi n}{q}, \quad n = 0, 1, 2, ..., q - 1
\]  

(1.34)

which gives \( q \) symmetric spin states, which lead to the construction sometimes being referred to as the Clock Model, as one can see in Figure 1-12.

Figure 1-12: Potts’s extension of the Ising model was to suggest that spin states could be generalised so as to allow an arbitrary number of states to occur. The construction of this model assumed that the spin states can be considered as \( q \)-symmetric angular subdivisions of a two-dimensional circle, with each state given by an angle \( \theta_n = \frac{2\pi n}{q} \) where \( n = 0, 1, ..., q - 1 \). Thus this model was sometimes referred to as the clock model for its similarity to a clock face, illustrated above for \( q = 2, 3 \) and \( 4 \). One should note that \( q = 2 \) corresponds to the so-called Ising value as for this value the Potts model is equivalent to the Ising’s.

In addition, Domb suggested the following interaction Hamiltonian,

\[
\mathcal{H} = \sum_{\langle i, j \rangle} J \cos(\theta_{si} - \theta_{sj})
\]  

(1.35)

where \( (\theta_{si} - \theta_{sj}) \) is the angle between two spins at neighbouring sites, \( i \) and \( j \) on the lattice and \( J \) is the strength of the interaction. The factor under the summation, \( \langle i, j \rangle \), is taken to be shorthand for the summation over all nearest neighbours. However, in solving the problem, Potts used the following simplified Hamiltonian:

\[
\mathcal{H} = \sum_{\langle i, j \rangle} J \delta(\sigma_i, \sigma_j)
\]  

(1.36)

where \( \delta(\sigma_i, \sigma_j) \) is the Kronecker delta function defined in the usual way, \( \delta(\sigma_i, \sigma_j) = \{ 1, i = j; 0, i \neq j \} \) and again \( J \) is the strength of the interaction of the states. With these Potts was able solve the problem in 2D for \( q = 2, 3 \) & 4, that is find phase transitions for the system, but unable to find any solution for \( q > 4 \).
1.3.3 The Extended Large-\(q\) Potts Model

In the previous discussion the main efforts in the development of the Potts model were concerned with analytical techniques, to derive closed-form solutions that can describe statistical properties of the material. At this juncture the development diverges and the model's evolution is now concerned with the qualitative rather than the quantitative. To understand this let us refresh with a mental picture of what comes so far. If we consider the Ising (and Potts) model, the spontaneous magnetisation is the occurrence of a dominant set of co-aligned atomic spin states that lead to a net permanent magnetic moment in the material. However, if we were to visualise such a configuration, then we would see that rather than a uniform distribution of alignments, we would see small pockets of moments irregularly distributed over the material's microcrystalline (\(L\)) structure as illustrated in Figure 1-13.

This perceived irregularity is a function of the degeneracy of the ground/spin states in the model which is prescribed by \(q\) which clearly for low \(q\), as in the case for the Ising value \(q = 2\), is what one might expect given that random spin flips occur without regard spin orientations of its neighbours beyond the Boltzmann probability. However, the situation becomes more interesting when we consider higher degeneracy for \(q > 2\), as suggested by Anderson and Grest [24].

In their model Anderson et al were considering the domain-growth kinetics of grain structures on a lattice, where the central feature was given to be dependent on the degeneracy of the so-called ground states, or the prescription of size of \(q\), and temperature. In practical terms to achieve such grain growth in real poly-crystalline alloy, one would raise the temperature of the alloy significantly beyond its melting point \(T_m\) and then rapidly quench it to \(T \ll T_m\). At which point one can observe a growth in grain structures as domains of similarly oriented crystals increase in size. The driving force of this process can be seen as an energy minimisation process as a result of nucleation of precipitates on the boundaries between anti-aligned crystals that minimise the overall boundary length, and as a consequence reduce the number of grains. In other words the small grains will coalesce to form larger grains that grow in size reducing boundary length thereby maximising entropy in the system.
The latter observation is central to the development of the CPM, specifically Anderson et al made the distinction that energy of the system is a function of temperature, but more importantly, energy fluctuations are only significant at nucleation sites. In terms of the Potts model this is reflected in their modified Hamiltonian:

$$\mathcal{H} = \sum_{\leq i, j>} (1 - \delta(\sigma_i, \sigma_j)),$$

(1.37)

where again $\delta(\sigma_i, \sigma_j)$ is the Kroenecker delta function, $J$ is the interactions energy and $< i, j >$ is the summation over all nearest neighbours. Clearly (1.37) is zero when $\sigma_i = \sigma_j$ and $J$ when $\sigma_i \neq \sigma_j$. Like the Ising/Potts models they suggested that $L$ consists of regularly arranged spin states $\sigma_i$ where $i = 1, 2, ..., N$, however in addition each state can have $Q \in \mathbb{N}$ multiplicity such that $1 < \sigma_i < Q$ leading to a degeneracy of $\sigma_i$. This degeneracy of states on $L$, coupled with (1.37) leads to homogenised domains of like states or as they put it “…higher ground-state degeneracy had an effect on the growth kinetics, the microstructure, and the topology.”. This can clearly be seen when we consider domain-growth for increasing $Q$ (Figure 1-14), where for low $Q$ the domains are irregular and asymmetric, but as $Q$ is increased the grain structure becomes more pronounced and equiaxed.

![Figure 1-14: Simulations of grain structures growing and becoming more pronounced and equiaxed as the degeneracy, $Q$, of the spin states increases.](image)

1.3.4 **The Monte Carlo Method.**

In the discussion so far we can see that most of the early work in the development leading to the CPM was (and largely still is) concerned with analytical techniques, that attempt to derive closed-form solutions to describe qualitative features of large particle system, classically referred to as an ensemble, in the “thermodynamic limit”. However problems emerge when we try to simulate such systems that exhibit correlative behaviour over vastly different velocity or spatio-temporal scales. One historically significant example of this was outlined by Fermi & Richtmyer [25] while investigating neutron chain reactions of fissionable materials at the Los Alamos National laboratory, were thermal and fast neutrons have a signification impact on critical reactivity in a nuclear reactor. Here “thermal” is given to mean neutrons...
that are in thermodynamic equilibrium with the surrounding environment, and are much less energetic than fast neutrons, and therefore have smaller velocities; thermal neutrons 1ev and fast neutrons 1Mev.

If we consider a nuclear fission chain reaction in an atom, the net result of the reaction is the ejection of fast neutrons (and other particles such as gamma rays). If there is sufficient supply of fissile material then these neutrons will collide (absorbed) with other atoms, causing further nuclear reaction leading to a critical, and unstoppable, self-sustaining reaction such as witnessed in a nuclear bomb or nuclear power plant reactor melt-down. If power production is the goal, then clearly this reaction needs to be controlled in some manner, and to do this a moderator is added to slow the reaction by reducing the velocities of the fast neutrons. This is achieved by adding thermal neutrons into the reaction, such that collisions between thermal and fast neutrons are elastic. That is the mass of the thermal neutrons should be equal to that of the fast neutrons, ensuring kinetic energy between the collisions is conserved, in this way slowing down the fast neutrons and controlling the reaction.

If we now consider developing a model simulation of the above process, we might suggest that to capture the underlying behaviour we need to model the spatio-temporal dynamics of both types of neutron, essentially capturing the mean free paths, or the velocity and position of each neutron in successive time intervals, in what Fermi & Richtmyer [25] referred to as “census time”. The problem with this approach is clear. Since the mean free paths of both fast and thermal neutrons are considered to be of the same order of magnitude, if we allow a time interval length sufficient for a thermal neutron to make a several collisions, then a fast neutron would have made several thousand, and clearly the simulation would be become intractable. In other words, a brute-force approach will not work.

To overcome this problem Fermi & Richtmyer employed a probabilistic approach conceived by Stanislaw Ulam at the Los Alamos laboratory while studying neutron diffusion, which he later described as the problem of “...how to change processes described by certain differential equations into an equivalent form interpretable as a succession of random operations.” [26]. Ulam conveyed this idea to John von Neumann, (also at Los Alamos) who saw the importance of the idea and immediately began work to implement it. Given the secrecy surrounding their work at the time, they decided to give it the codename which has endured to this day, the Monte Carlo Method. Incidentally the idea for the name has been widely reported as von Neumann’s suggestion, however Metropolis recounted in a publication much later how he “...suggested an obvious name for the statistical method-a suggestion not unrelated to the fact that Stan had an uncle who would borrow money from relatives because he ‘just had to go to Monte Carlo’” [27]. Be that as it may, the approach can be summarised by the following algorithm:

1. Define a domain of possible inputs.
2. Generate inputs randomly from a probability distribution over the domain.
3. Perform a deterministic computation on the inputs.
4. Aggregate the results.

Algorithm 1
We should note at this point, that this algorithm/approach was by no means new, and indeed an early variant of this approach was given in the 18th century by Georges-Louis Leclerc, Comte de Buffon’s needle experiment to approximate the value of $\pi$. Since Buffon’s experiment is probably the most intuitive description of how the so-called Monte Carlo algorithm works, so we shall briefly describe it here. Let us take a unit square and inscribe it with a unit circle (I). Now sprinkle grains of sand uniformly over the square and circle (II). Now take the count of the total number of grains $N_T$ and the count of those contained exclusively within the circle $N_C$ (III). Finally take the ratio of the two counts and equate it to the ratio of the area of the square to the area of the circle, that is $N_C/N_T \approx 1/\pi$ which implies $\pi \approx N_T/N_C$ (IV). We can see that if we iterate this approach, adding more and more grains of sand, then we should expect this process to converge, for all practical purposes, on the true value of $\pi$.

The true power of the approach however, is given by its applicability to computational processing by computer, first sketched in 1949 by Metropolis & Ulam [28] in what is considered to be paper that first alluded to this method, where they themselves point out that “...modern computing machines are extremely well suited to performing the procedures described [the Monte Carlo Method]”. However it was until 1953 in a subsequent paper by Metropolis et al [29] that provided the first complete general Monte Carlo technique of which they propose “... a general method, suitable for fast electronic computing machines, of calculating the properties of any substance which may be considered as composed of interacting individual molecules.”.

In their paper they considered a system of $N$ identical moving particles with spherically symmetric potentials, arranged on a regular two-dimensional lattice $\mathcal{L}$ (here we can take $\mathcal{L}$ to be equivalent to the two-dimensional Ising lattice), and assumed the system is given as a canonical ensemble, implying that the system is in a state of thermodynamic equilibrium with its environment. Generally speaking, if the positions of the particles are known, it is possible to calculate the potential energy of the system by

$$E = \sum_{i=1}^{N} \sum_{j=1}^{N} V(d_{ij}), \quad i \neq j$$  \hspace{1cm} (1.38)

where $V$ is the potential energy between particles and $d_{ij}$ is given as the minimum distance between particles $i$ and $j$. To calculate a specific property, $F$, of the system one integrates the weighted probability of that quantity over the sum of all possible microstates in $2N$ configuration space:

$$\langle F \rangle = \int \mathcal{F} e^{-\frac{E}{kT}} d^{2N} p \, d^{2N} p / \int e^{-\frac{E}{kT}} d^{2N} p,$$ \hspace{1cm} (1.39)

where $d^{2N} p$ is taken to be the volume element (note the comparisons between (1.38) and (1.39) with Ising’s model (1.29) and (1.30)). However, and equally with Fermi & Richtmyer, (1.39) quickly becomes intractable when $N$ becomes large, say several hundred particles. To solve this problem they of course suggested a Monte Carlo algorithm, however rather than randomly sampling a configuration and weighting it against its Boltzmann probability (as suggested in (II) in the above algorithm), they picked configurations with Boltzmann probability and weighted them evenly.
To demonstrate this, consider a single particle $p$ on a two-dimensional initial lattice configuration, $L_0$, at a starting positions $p_0 = [x_0, y_0]^T$, and assume that it attempts to move to a new position such that

$$p_1 = p_0 + \alpha \begin{bmatrix} \xi_1 \\ \xi_2 \end{bmatrix},$$

(1.40)

where $\alpha$ is the lattice constant that represents the maximum arbitrary distance the particle is allowed to travel in one step of the simulation, and $\xi_1$ and $\xi_2$ are numbers generated at random such that $\xi_k$ is between 1 and $-1$; the particle will attempt to move in a purely random direction in each step a distance no greater than $\pm 2\alpha$ in any direction from its original position. If we now calculate the difference in the energy, $(E_0 - E_1) = \Delta E$, of this new configuration, $L_1$, then the change will be accepted or rejected according to the Boltzmann probability:

$$P = \begin{cases} 
1, & \Delta E < 0 \\
1, & \Delta E > 0 \text{ and } \xi_3 < e^{-\frac{\Delta E}{k_BT}} \\
0, & \text{otherwise}
\end{cases}$$

(1.41)

were $\xi_3$ is a random number between 0 and 1. If we iterate this process for each of the $N$ particles, recording the state of the configuration in each move attempt (regardless of the outcome), then one can see that the properties of the system can now be given by a statistical average:

$$\langle F \rangle = \frac{1}{M} \sum_{j=0}^{M} F_j$$

(1.42)

where $F_j$ is the value of the property $F$ of the system after the $j^{th}$ iteration. We can assume this if we appeal to the law of large numbers, which implies that the configuration will conform to the Boltzmann distribution within a finite number of iterations, and therefore so should any statistical measure we make on it. Put another way, if we assume that $E_0 > E_1$ then the relative probability of the system being in these two energy states is given by

$$\frac{P(E_0)}{P(E_1)} = e^{-\frac{(E_0-E_1)}{k_BT}},$$

(1.43)

then the probability outcome can be seen to be a function of the temperature of the system. That is for very high temperatures if $k_BT \gg | - (E_0 - E_1)|$, then the states $E_0$ and $E_1$ are equally likely to exist, since in a thermally agitated state the positions are random. Conversely for low temperatures if $k_BT \ll | - (E_0 - E_1)|$ it is more probable that the system will tend to lower energy states, and therefore, as long as we specify an appropriate temperature in any simulation, one can see the system will conform to the canonical distribution.
1.3.5 The Cellular Potts Model

Through the work of Ising and Potts and the development of sophisticated computational machines to model neutron chain reactions in the development of “the bomb”, and tractable methods to solve such problems, namely the Monte Carlo method, we arrive at the last key stage that marks the introduction of the these models into the world of biological cell simulations. In this regard it was through the work of Graner & Glazier in researching biological cell sorting [20, 30], that showed how large scale spatial cell relocation could be simulated using the extended Potts model of Anderson and Grest.

The fundamental difference between the models of Anderson & Grest and Graner & Glazier, is that in the former domain growth in the granular structure is a function of grain boundary growth, that is metallic grains grow by increasing their boundary length by coalescing of smaller grains and absorbing boundary segments of other equally sized neighbouring grains. In the latter however biological cells do not appreciably changes their volumes during their life-cycles. The implication of this difference is in the effective manipulation of system energy. Both will attempt to maximise the entropy of their respective systems (given their Ising/Potts derivation), however one achieves this by domain coarsening of a granular structure (Figure 1-14) and thereby reducing the boundary length, while the other by spatial reorganisation of cells into a configuration that minimises contact energy between dissimilar cell types, they termed as differential adhesivity. Thus the advancements to the Potts model made by Graner & Glazier, were firstly to impose a volume constraint on what were previously termed grains to what are now termed cells, and that each cell can have a “type” with each type having the ability to define its own attributes.

To put this in perspective of the large-Q Potts Model, let us assume again we have a two-dimensional Ising lattice, L, and that lattice sites are given by the state variable \( \sigma_{ij} \) having \( Q \in \mathbb{N} \) possible states such that \( 1 < \sigma_{ij} < Q \) where \( i,j \) refers to the lattice coordinates in two-dimensions and are arranged such that \( \sigma_{0,0} \) refers to the site at the top-left-most coordinate and \( \sigma_{n,n} \) the bottom-right-most. Again \( \sigma_{ij} \) are not unique on \( L \), that is, there is a degeneracy of states and this degeneracy implies identity with respect to a specific grain or cell. That is the \( k^{th} \) cell has a number of \( \sigma_{ij} = k \in Q \), collectively defined as \( \sigma^k \), such that \( L = \sum_{k=1}^{Q} \sigma^k = \sum_{i=0}^{n} \sum_{j=0}^{n} \sigma_{ij} \). Put another way, each cell has an area defined as the sum of all \( \sigma_{ij} = k \).
The calculation of the current energy state of $\mathcal{L}$ can be defined similarly to (1.37), but now we take into account that energy calculations on the boundaries between cells are now a function of type,

$$
H = \sum_{i \neq j} J(\sigma_{i,j}, \sigma_{i,j}) \left( 1 - \delta(t(\sigma_{i,j}), t(\sigma_{i,j})) \right)
$$

(1.44)

were we see that the contribution between like $\sigma_{i,j}$ are zero, and $J(\sigma_{i,j}, \sigma_{i,j})$ otherwise, implying as suggested that the contribution between dissimilar $\sigma_{i,j}$ are not constant, but a function of boundary interactions between cells. More specifically this expresses that for any given $\sigma_{i,j}$ on $\mathcal{L}$, the energy contribution to the system at this site is the sum of the energies of all its nearest neighbours $\sigma_{i,j}$ such that $\sigma_{i,j} \neq \sigma_{i,j}$.

**Differential Adhesion**

The exact contribution of each neighbour satisfying this inequality is given by a triangular adhesion matrix that describes the adhesive relationships, or contact energies, between the cell types on the lattice, which implies that adhesion forces between cells types are symmetric, and indeed this is true of all forces in the CPM; energetic interactions between cells of differing types on $\mathcal{L}$ conform to Newton’s third law. Considering Glazier & Graner [20] they defined three cell types: Substrate (S), Dark (D) and Light (L), to demonstrate how a random configuration of these cell types would sort themselves into homogenous, homotypic domains. That is, Dark cells would preferentially adhere to Dark, and Light would equally adhere to Light, with the Substrate acting as a special medium on which the Light and Dark cells interact as illustrated in Figure 1-16.

![Figure 1-16: Results of a cell simulation for Dark and Light cells from [20] that demonstrates how differential adhesion between the differing cell types can lead to cell sorting.](image)

To achieve this result Glazier & Graner specified the following adhesion matrix, were the entries in the first matrix of the form #,# demonstrate the interactions between cell types. In this example the adhesive bonds between Dark and Dark cells are given as $(D, D)$ and between Dark and Light by $(D, L)$ with the second matrix giving the actual integer adhesive bonding values used in [20].

$$
J(\sigma_{i,j}, \sigma_{i,j}) = \begin{bmatrix}
S, S & S, D & S, L \\
S, D & D, D & D, L \\
S, L & D, L & L, L
\end{bmatrix} = \begin{bmatrix}
16 & 16 & 16 \\
16 & 2 & 11 \\
16 & 11 & 14
\end{bmatrix}
$$

(1.45)
Considering these values, and the fact that the CPM will attempt to minimise contact energy, we can see directly the following relationships are prescribed: Dark cells will adhere to Dark cells preferentially over all interactions since $D, D < D, L < L, L < S, S$. Light cells will prefer to adhere to Dark cells rather than themselves as $L, L > D, L$. However Dark and Light cells will avoid contacts with Substrate cells as $L, L < S, S > D, D$. But since Dark to Dark cells have the smallest contact energy, Light cells will be forced to interact with the Substrate $(D, D < L, L)$ which can be seen in Figure 1-16B as the Light cells engulfing the Dark.

**Incompressibility and Volume Constraint**

Affecting this influence of adhesive bonding, or contact energies, is the other addition by Glazier & Graner, namely a volume constraint. In the absence of such a constraint, the cell structure on the lattice would coarsen \([24]\), eventually to a single large cell with area equal to sum of the areas of the initial cells. This is because the CPM evolves to maximise the entropy of the system, implemented as boundary level interactions whereby cells contract or expand said boundaries, such that changes would cause a lowering of system energy. In terms of the adhesion matrix in equation \((1.45)\), we can see that this would imply that we would be left with a single large Dark cell, as all intracellular interactions would occur preferentially for Dark cells and thus the Light cells would be consumed and disappear.

Thus the aim of the volume constraint is to maintain, on average, cells at a constant area, or in other words we impose an incompressibility constraint. This was achieved by exacting an energetic penalty for any boundary level interaction that would cause a cell’s area to deviate from a specific target value. Glazier & Graner defined this penalty by first defining the area of the \(k^{th}\) cell as \(a(\sigma^k)\) and giving it a type dependent target area \(A(\tau(\sigma^k))\), and then suggesting that the penalty grows quadratically as the \(a(\sigma^k)\) deviates from \(A(\tau(\sigma^k))\). However this penalty should not be imposed for the surrounding substrate, referred to as \(S\) in the definition of the adhesion matrix \((1.45)\). Biologically the substrate can be thought of as a surrounding fluid medium, for example a growth culture as found in petri dishes, or a surrounding extracellular matrix. Therefore they did not impose a volume constraint for this special cell type. Thus the total penalty exacted for one energy computation is expressed as,

\[
\mathcal{H}_\nu = \lambda \sum_{<k>} [a(\sigma^k) - A(\tau(\sigma^k))]^2 \theta(\tau(\sigma^k))
\]  

\[(1.46)\]

were \(< k >\) is a shorthand notation for the summation of all interacting cells in the current computation, and \(\theta(\tau) = \{0: \tau = S; 1: \tau \neq S\}\) and \(\lambda\) is a model parameter specifying the strength of the area constraint. The effect of the area constraint can be seen as the elasticity or an incompressibility factor of cell boundaries. For example when for very small \(\lambda \rightarrow 0\) the cells would grow unbounded, effectively turning off the area constraint and reducing the model to that of Anderson & Grest. For very large \(\lambda\) the constraint would freeze cells on the lattice at their current area, pinning the cells and revealing the underlying discretization of the lattice structure. Thus there is an intermediate \(\lambda\) that allows for cell
boundaries to fluctuate in a manner similar to their biological counterparts. Lastly the current energy state of \(L_0\) is given by the sum of local nearest neighbour interactions \(\forall \sigma_{ij} \in L_0\) and is defined as follows:

\[
\mathcal{H}(\sigma_{ij}) = \sum_{<ij>} J(\sigma_{ij}, \sigma_{ij}) \left(1 - \delta(\tau(\sigma_{ij}), \tau(\sigma_{ij}))\right) + \lambda \sum_{<k>} \left[ a(\sigma^k) - A(\tau(\sigma^k)) \right]^2 \theta(\tau(\sigma^k)).
\] (1.47)

### 1.3.6 Implementation Of The CPM

What remains is of an algorithmic nature in describing how one would implement (1.47) within a software program. In general equation (1.47) prescribes exactly how we should calculate the energy associated with single lattice point \(\sigma_{ij}\) from the sum of contributions from nearest neighbours \(\sigma_{ij}\). Given an arbitrary lattice such as that illustrated in Figure 1-15, we follow a Monte-Carlo scheme, whereby we make random changes on cell boundaries, that is we pick at random a \(\sigma_{ij}\) then attempt to change its identity to that of another adjacent cell. That is we pick a nearest neighbour of \(\sigma_{ij}\) at random say, \(\sigma^R_{ij}\), and for both of the sites we compute the energy before and after the attempted change in identity, that is

\[
\Delta E = \mathcal{H}(\sigma_{ij}) - \mathcal{H}(\sigma^R_{ij}),
\] (1.48)

and accept this change with Boltzmann probability as in (1.41), reproduced here for convenience:

\[
P = \begin{cases} 
1, & \Delta E < 0 \\
1, & \Delta E > 0 \text{ and } \xi < e^{\frac{\Delta E}{k_B T}} \\
0, & \text{otherwise.}
\end{cases}
\]

This process continues until all of the sites on the lattice have attempted such a change, at such time we say a Monte-Carlo step has been computed, and we repeat the whole process again.

As suggested previously, it might take tens of thousands of such steps to converge on an outcome that sufficiently resembles any observable behavior we are modelling. In terms of our problems, biological cell dynamics behaviour, this can be readily seen by direct observation, however we should note that herein lies the largest problem with the CPM. There has been no formal analysis carried out on it to determine an optimal configuration of model parameters that have a consensus amongst its adopters. In the literature it is usual to find justification by morphological characteristics (as we do here), or by association with other applications by different adopters, therefore the CPM can only be considered as a qualitative model at best.
Chapter 2

1D Continuous Models

for Chemotactically Moving Cells

2.1 Introduction

In this chapter we describe the definition and analysis of a one-dimensional continuous model, designed for the study of motile dynamics of a group/population of cells and to analyse travelling solutions. In deriving the model we do not assume any biological factors pertaining to cellular dynamics, such as intracellular adhesion, cellular density or include any experimental observations. In particular we are only interested in deriving conditions for the group to move given an applied force, which results in the group moving (potentially) with constant, uniform velocity. The applied force, termed chemotaxis, is the reaction of the group to move along (attract) or against (repel) the gradient of a locally produced chemical, which we define to be produced either within the group or one of its sub-populations (internal), or by the surrounding environment (external). Further we make the distinction that the composition of the group may vary in terms of how sub-populations of the group react to the chemical. That is, we assume that the group may be composed of two sub-populations, with one population reacting to the chemical, while the other does not. Or in the simpler case, the group is comprised of a single population all reacting equivalently. We will use homogenous to represent a group composed of a single population of equivalently reacting cells, and heterogeneous to represent the composition of two unequally reacting sub-populations.

For the remainder of this chapter we shall investigate the construction and analysis of these models, with the primary focus of understanding the behaviour of dynamical systems that result, to analyse the stability and existence of travelling solutions. The mathematical model we derive here is based on reaction diffusion systems implemented as a system of linear piecewise partial-differential equations, the solutions of which describes the chemical dynamics, and the associated chemotaxis, or if you will, velocity function. We shall proceed by dividing modification of the model, in terms of heterogeneous and homogenous dynamics, defining the respective equations (and solutions) and analysing the dynamical systems that result.
2.2 Homogenous Model of a Migrating Group Of Cells

As suggested the homogenous model is composed of a group of homotypic cells (of the same type), all of which share an equivalent motile response to the chemotactic chemical. The model describes the concentration \( u(x, t) \) of a chemical (morphogen, chemotactic agent) which is produced within a domain of constant size, \( a \), moving with constant speed \( c(t) \). This chemical can diffuse outside the production domain and degrade. This implies that in the homogenous model, the group is reacting chemotactically to a chemical it produces, resulting in a closed autonomous chemotactic system, an auto-chemotaxis if you will. As one would expect the emergence of travelling solution within such a system would require special initial conditions, such as a perturbation or agitation of the group from the ‘dominant’ stationary equilibrium. Thus the challenge of the homogeneous system, and something not in the heterogeneous model, is what chemotactic force is necessary for the group to move uniformly with constant speed, given an initial perturbation.

2.2.1 Concentration profile of Internally Produced Chemotactic Agent

From a biological point of view we can consider the production domain as a domain of transcription (DoT) expressing intracellular mRNA, which in general is held at a constant concentration within cells and does not permeate through cell membranes. Thus the production domain can be considered as a source for the diffusing chemical, \( u(x, t) \), defined in the interval \( a(t) < x < \beta(t) \) (see Appendix A), representing the width of the activating domain of \( u(x, t) \). We assume a constant concentration within \( a(t) < x < \beta(t) \) (grey shaded region in Figure 2-1A), and is zero concentration everywhere else. Therefore we can model the diffusing chemical \( u(x, t) \) as:

\[
\frac{\partial u}{\partial t} = D \Delta u - k_1 u + p(x, t), \quad k_1, D \in \mathbb{R}^+ \tag{2.1}
\]

where \( x \in (-\infty, \infty) \) and \( t \in [0, \infty) \), \( k_1 \) is the constant rate of decay proportional to the concentration of \( u \), \( D \) is the coefficient of diffusion, \( \Delta \) the Laplacian differential operator in one dimension and \( p(x, t) \) is defined as:

\[
p(x, t) = \begin{cases} k_2, & a(t) < x < \beta(t) \\ 0, & \text{otherwise} \end{cases} \tag{2.2}
\]

where \( k_2 \) is a constant describing production rate within production domain and zero everywhere else.

Further if the size of the production domain is \( a \), and moves with constant speed \( c \), we have symmetrically posed about the origin by setting \( a = -a/2 + ct \) and \( \beta = a/2 + ct \). We shall derive models in their general form in terms of \( a \) or \( \beta \) and express solutions, when it simplifies the result, in terms of \( a \). Thus the chemical, \( u(x, t) \), diffuses from the production domain, \( p(x, t) \), into the surrounding environment and acts as a chemotactic agent.

We can, through a suitable change of variables, repose our problem if we assume that solutions to (2.1) are given as travelling waves. Then in a co-moving frame of reference the profiles of such waves do not change in time and therefore (2.1) becomes independent of time. Thus if we introduce the variable \( \xi = x - ct \) and substitute into (2.1) and (2.2) our problem becomes,
\[ D \frac{d^2 u(\xi)}{d\xi^2} + c \frac{du(\xi)}{d\xi} - k_1 u(\xi) + p(\xi) = 0, \quad (2.3) \]

and our problem is now independent of time and has gained an advective term proportional to \( c \).

\[ p(\xi) = \{k_2, \; \alpha < \xi < \beta \} \quad (2.4) \]

However we can make a symbolic change for convenience and assume that \( \xi = x \) and the piecewise solution to (2.3)-(2.4) can be stated as:

\[
0 = \begin{cases} 
D\Delta u + cu_x - k_1u, & x < \alpha \\
D\Delta u + cu_x - k_1u + k_2, & [x] \leq [\beta - \alpha] \\
D\Delta u + cu_x - k_1u, & x > \beta
\end{cases} \quad (2.5)
\]

where \( c \) is the speed of the frame and we have substituted directly the definition (2.4). Solutions of the system (2.5) can be shown to be of the form (See Appendix A):

\[
u = \begin{cases} 
Ae^{\lambda_1 x} + Be^{\lambda_2 x}, & x \leq \alpha \\
Ce^{\lambda_1 x} + De^{\lambda_2 x} + \frac{k_2}{k_1}, & \alpha \leq x \leq \beta \\
Ee^{\lambda_1 x} + Fe^{\lambda_2 x}, & x \geq \beta
\end{cases} \quad (2.6)
\]

where

\[
\lambda_{1,2} = \frac{-c \pm \sqrt{c^2 + 4k_1D}}{2D} \quad (2.7)
\]

and the unknown coefficients: \( A, B, C, D, E \) and \( F \) can be determined by application of boundary conditions. For clarity we will use the subscript, \( i \in [1,2,3] \), to identify the intervals defined in (2.5) ordered from left to right on the \( x \)-axis. Using this subscript we can prescribe the boundary condition at the extremes of the spatial domain as,

\[
u_1(-\infty) = u_3(\infty) = 0, \quad (2.8)
\]

and at the boundaries of the intervals, which by necessity of continuity we require \( u = C^1(-\infty, \infty) \), giving the conditions on \( u \) as:

\[
u_1(\alpha) = u_2(\alpha) \quad \nu_2(\beta) = u_3(\beta) \quad (2.9)
\]

and

\[
\begin{align*}
\left. \frac{du_1}{dx} \right|_{x=\alpha} &= \left. \frac{du_2}{dx} \right|_{x=\alpha} \\
\left. \frac{du_2}{dx} \right|_{x=\beta} &= \left. \frac{du_3}{dx} \right|_{x=\beta}
\end{align*} \quad (2.10)
\]

Therefore we can solve for the coefficients in (2.6) by employing the boundary conditions ((2.8)-(2.10), and derive the piecewise solutions for the system (2.5) as (See Appendix A for solution):

\[
u = \begin{cases} 
k_2 & \lambda_2 \left( e^{\lambda_1(x-\beta)} - e^{\lambda_1(x-\alpha)} \right), \quad x \leq \alpha \\
k_2 & \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \left( \lambda_2 e^{\lambda_1(x-\beta)} - \lambda_1 e^{\lambda_2(x-\alpha)} + (\lambda_1 - \lambda_2) \right), \quad \alpha \leq x \leq \beta \\
k_2 & \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \lambda_1 \left( e^{\lambda_2(x-\beta)} - e^{\lambda_2(x-\alpha)} \right), \quad x \geq \beta
\end{cases} \quad (2.11)
\]
for which we can illustrate both stationary (Figure 2-1A) and travelling segments (Figure 2-1B).

Figure 2-1: Plots of stationary and travelling segments for the homogenous model. A: For a group/segment of width \( a \) on the spatial \( x \)-axis we show a stationary (\( c = 0 \)) solution profile of the \( u \) (solid black profile) illustrating a symmetric distribution. B: For the same group we can show a travelling solution (\( c \neq 0 \)) illustrating an asymmetric distribution with the maximum concentration lags the mid-point and the concentration on the back of the domain is above the concentration on the front.

Considering (2.11) the first observation we can make is that it is a symmetric function when \( c = 0 \), the location of maximum concentration, \( x_{\text{max}} \), is located at the origin. However in a travelling solution, \( x_{\text{max}} \) shifts away from the origin in either the positive or negative \( x \)-direction depending on whether the domain moves to the right (\( c > 0 \)) or to the left (\( c < 0 \)) The magnitude of the shift increases with the speed of the group. The exact location can be found by equating the spatial derivative of \( u_2 \) to zero, giving

\[
\frac{d u_2(x)}{dx} = \frac{d}{dx} \left[ \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} (\lambda_2 \lambda_1 e^{\lambda_1 (x-\beta)} - \lambda_1 e^{\lambda_2 (x-\alpha)} + (\lambda_1 - \lambda_2)) \right]
\]

\[
= \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} (\lambda_1 \lambda_2 e^{\lambda_1 (x-\beta)} - \lambda_2 \lambda_1 e^{\lambda_2 (x-\alpha)})
\]

\[
= \frac{k_2 \lambda_1 \lambda_2}{k_1 (\lambda_1 - \lambda_2)} (e^{\lambda_1 (x-\beta)} - e^{\lambda_2 (x-\alpha)}). \tag{2.12}
\]

then the condition for (2.12) to be equal to zero can only be true when

\[
e^{\lambda_1 (x-\beta)} - e^{\lambda_2 (x-\alpha)} = 0
\]

\[
e^{\lambda_1 (x-\beta)} = e^{\lambda_2 (x-\alpha)}
\]

\[
\lambda_1 (x-\beta) = \lambda_2 (x-\alpha)
\]

\[
x (\lambda_1 - \lambda_2) = \beta \lambda_1 - \alpha \lambda_2
\]

\[
x = \frac{\beta \lambda_1 - \alpha \lambda_2}{(\lambda_1 - \lambda_2)}
\]

which gives, after putting the expression for \( \lambda_1 \) and \( \lambda_2 \), the location of the maximum concentration of \( u \),

\[
x_{\text{max}} = -\frac{ca}{2\sqrt{c^2 + 4k_1 \beta}} \tag{2.13}
\]

Considering (2.13) we can see that for \( c > 0 \) the location of \( x_{\text{max}} \) shifts to the left and for \( c < 0 \) shifts to the right, that is the sign of \( c \) describes the group’s direction of motion, and trivially for \( c = 0 \) implies \( x_{\text{max}} = 0 \) indicating a stationary solution. In addition we should note that for any speed the
definition (2.13) ensures the location of the maximum is always located within domain, \(-a/2 \leq x_{\text{max}} \leq a/2\) with \(x_{\text{max}} \rightarrow -a/2\) when \(c \rightarrow -\infty\) and \(x_{\text{max}} \rightarrow a/2\) when \(c \rightarrow \infty\).

2.2.2 Motion due to chemotaxis

Chemotaxis as a chemo-mechanical process is observed throughout nature in both plants and animals, in many diverse situations such as in the probing behaviour of plant meristems, aggregation of the amoebae Dictyostelium discoideum, travelling bands of bacteria and in particular importance in this work, axial migration of cells during embryogenesis.

Due to chemotaxis, the organism will attempt to move towards areas of higher concentrations of the chemotactic agent, termed attraction, or towards areas of lower concentrations, termed chemorepulsion. The mechanisms by which they determine the gradient varies from organism to organism. For example, certain amoebae are of insufficient size and/or complexity to determine the gradient of the substance, so their response is to perform random walks, literally sampling the substance periodically. Conversely larger cells, such as eukaryotic cells, can determine the differential concentration over the length of their bodies, and thus can determine the gradient of the substance at a specific point. Characteristically the implication is that organisms have a motile response to the local gradient of a substance, regardless of the mechanism they employ, and this is the premise of our chemotaxis model. Thus the question becomes how do we implement, or describe this in terms of solution ((2.11))? We could take the gradient at some specific point within the group:

\[
c = c_0 \frac{du}{dx}
\]

where \(\alpha \leq x_0 \leq \beta\)

where \(c_0\) is the strength of chemotaxis, or if you will, the sensitivity of the group to the chemical gradient. This raises the question of which point is most appropriate within the group, and of course we could arbitrarily pick a point, say \(x_0 = \beta\), and use this consistently throughout the analysis. However we take a more “representative” approach, and consider the total gradient to be calculated as an average in (2.14) over the group for which we can show:

\[
c = \bar{c} = \frac{c_0}{a} \int_{a}^{\beta} u_x dx
\]

\[
= \frac{c_0}{a} (u(\beta) - u(\alpha)).
\]

Since the concentrations on the boundaries of the group are equal at the boundaries of ((2.11)), the choice of which to substitute depends only on where \(u_1\) is defined. Substituting the expression for \(u_2\) from ((2.11)) into (2.15) we find:

\[
c = c_0 \frac{u_2(\beta) - u_2(\alpha)}{|\beta - \alpha|}
\]

\[
= c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \frac{\lambda_1 e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)}}{\lambda_1 e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)}}
\]

\[
= c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \frac{\lambda_1 (e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)})}{\lambda_1 e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)}}
\]

\[
= c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \frac{\lambda_1 (e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)})}{\lambda_1 e^{\lambda_1(\beta - \alpha)} - e^{\lambda_1(\beta - \alpha)}}
\]
\[ c = \frac{k_2 c_0}{\beta - \alpha |k_1(\lambda_1 - \lambda_2)|} \left( \lambda_1 (1 - e^{\lambda_2(\beta - \alpha)}) - \lambda_2 (e^{\lambda_1(\alpha - \beta)} - 1) \right) \]

Which, after substituting in terms of a production domain of width \( a = \beta - \alpha \), leads to the definition of the chemotaxis function:

\[ c = \frac{k_2 c_0}{ak_1(\lambda_1 - \lambda_2)} \left( \lambda_1 (1 - e^{a\lambda_2}) + \lambda_2 (1 - e^{-a\lambda_1}) \right). \tag{2.16} \]

Thus the dynamical behaviour of the group, specifically in terms of travelling solutions, is given by the equation (2.16). However the analysis of this is non-trivial since \( \lambda_{1,2} \) are both functions of \( c \), which not only implies (2.16) is a non-linear function, but is also implicit in \( c \). Therefore we have a non-linear dynamical system where concentration \( u(x) \) given by ((2.11) depends on speed \( c \) and is defined by the current \( u \) has on changes to (2.16), which we can write as:

\[ c = f(c). \tag{2.17} \]

The consequence of the implicit and non-linear nature of the system (2.3)-(2.16), as is the case with many non-linear dynamical systems, is that determining analytical solutions is either at best non-trivial and/or intractable or at worst insoluble. In the case of (2.16) we can see by direct observation, that there is at least one solution when the right-hand side of (2.16) \( c = 0 \) that corresponds to a stationary solution, and another is that \( \text{RHS} \to 0 \). However beyond these superficial observations, we cannot determine (analytically) the existence of any further solution(s). Therefore if we are to determine any further solutions, and indeed the dynamical behavior governed by (2.16), we will need to employ qualitative techniques.

As stated above there is one stationary solution when \( c = 0 \), and we can show this to be true for the entire parameter space of the model,

\[ c = 0 \Rightarrow \frac{dk_2 c_0}{ak_1 \sqrt{k_1} d} \left( \frac{k_1}{d} (1 - e^{-a \frac{k_1}{d}}) - \frac{k_1}{d} (1 - e^{-a \frac{k_1}{d}}) \right) = 0 \tag{2.18} \]

whereas on the other hand travelling solutions correspond to \( c \neq 0 \). We can determine if such solutions exist by graphical considerations if we plot functions of the left and right-hand sides of (2.16) as independent functions:

\[ g(c) = c \tag{2.19} \]

and

\[ f(c) = \frac{k_2 c_0}{ak_1(\lambda_1 - \lambda_2)} \left( \lambda_1 (1 - e^{a\lambda_2}) + \lambda_2 (1 - e^{-a\lambda_1}) \right). \tag{2.20} \]

against each other on the same axis. Travelling solutions will correspond to non-zero graphical points of intersection between these two functions as illustrated Figure 2-2A. These solutions represent a segment moving with constant and uniform speed.
Figure 2-2: Illustration of travelling solutions for the chemotactically moving segment. A: Plots of $f(c)$ and $g(c)$ for chemo-repulsion ($c_0 < 0$, solid black profiles) and chemo-attraction ($c_0 > 0$, dotted lines). For both chemo-repulsion and attraction there always exists a travelling solution when $c = 0$ located at the origin of the graph. However for chemo-attraction ($c_0 < 0$) two further travelling solutions occur with the condition that $f'(c = 0) > 1$. This is illustrated as points of intersection between $f(c)$ and $g(c)$, describing two symmetric solutions interpreted as the segment travelling right-wise (B) or left-wise (C) on the x-axis with equal but opposite velocity. The asymptotic behaviour for $c \to \infty$, can be seen by noting $\lambda_1 \to 0$ and $\lambda_2 \to \infty$ as $c \to \infty$ and $\lambda_1 \to \infty$ and $\lambda_2 \to 0$ as $c \to -\infty$. Substituting these expressions into (2.16) we can see directly that $f(c) \to 0$ as $c \to \pm \infty$. Model parameters where chosen such that: $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$ and $a = 50$.

An important observation we can make from the plots illustrated in Figure 2-2A, is that the chemotaxis function is odd, and therefore we can expect not one but two solutions, that are equal in magnitude and opposite in sign. The physical interpretation of these solutions can be seen as the segment moving in opposite directions. If there is a travelling solution to the right, then there will also be a travelling solution to the left, illustrated respectively in Figure 2-2B and Figure 2-2C. We can determine whether such solutions exist as condition on $c_0$ from the observation that travelling solutions exist only at intersections between $f(c)$ and $g(c)$, and further, this can be true only when $\frac{df(c)}{dc} |_{c=0} > 1$, that is:

$$\frac{df(c)}{dc} = \frac{d}{dc} \left( \frac{k_2 c_0}{ak_1(\lambda_1 - \lambda_2)} \left( \lambda_1(1 - e^{a\lambda_2}) + \lambda_2(1 - e^{-a\lambda_1}) \right) \right)$$

$$= \frac{k_2 c_0}{ak_1} \frac{d}{dc} \left( \frac{\lambda_1(1 - e^{a\lambda_2}) + \lambda_2(1 - e^{-a\lambda_1})}{(\lambda_1 - \lambda_2)} \right)$$

$$= \frac{k_2 c_0}{ak_1} \left( \frac{d}{dc} \left( \frac{\lambda_1(1 - e^{a\lambda_2})}{(\lambda_1 - \lambda_2)} \right) + \frac{d}{dc} \left( \frac{\lambda_2(1 - e^{-a\lambda_1})}{(\lambda_1 - \lambda_2)} \right) \right)$$

$$= \frac{k_2 c_0}{ak_1} \frac{d\lambda_1}{dc} \left( \frac{1 - e^{a\lambda_2}}{(\lambda_1 - \lambda_2)} \right) \frac{\lambda_1 e^{a\lambda_2}}{(\lambda_1 - \lambda_2)}$$

$$- \frac{\lambda_1(1 - e^{a\lambda_2})}{(\lambda_1 - \lambda_2)^2} \frac{d\lambda_1}{dc} + \frac{d\lambda_2(1 - e^{-a\lambda_1})}{(\lambda_1 - \lambda_2)} + \frac{d\lambda_1}{dc} \frac{\lambda_2 e^{-a\lambda_1}}{(\lambda_1 - \lambda_2)}$$

$$- \frac{\lambda_2(1 - e^{-a\lambda_1})}{(\lambda_1 - \lambda_2)^2} \frac{d\lambda_2}{dc} = \frac{1}{1}$$
evaluating \( \lambda_{1,2} \), and their derivatives at \( c = 0 \) we can show that,
\[
\lambda_{1,2} \bigg|_{c=0} = \pm \frac{k_1}{\sqrt{d}} \quad \text{and} \quad \frac{d\lambda_{1,2}}{dc} \bigg|_{c=0} = -\frac{1}{2d},
\] (2.21)
and substituting these values back into \( \frac{df(c)}{dc} \) gives:
\[
\frac{df(c)}{dc} \bigg|_{c=0} = \frac{k_2 c_0}{ak_1} \left( -\frac{1 - e^{-a \sqrt{d}}}{4d \sqrt{k_1}} + \frac{k_1}{4d \sqrt{d}} e^{-a \sqrt{d}} - \frac{1 - e^{-a \sqrt{d}}}{4d \sqrt{k_1} \sqrt{d}} - \frac{k_1}{4d \sqrt{d}} e^{-a \sqrt{d}} \right),
\]
which after simplification we have
\[
\frac{df(c)}{dc} \bigg|_{c=0} > 1 \Rightarrow \frac{k_2 c_0}{2ak_1 \sqrt{d} k_1} \left( e^{-a \sqrt{d}} \left( \frac{k_1}{\sqrt{d}} + 1 \right) - 1 \right) > 1.
\] (2.22)

However while (2.22) is a necessary condition for travelling solutions to exist, it is not sufficient to enable a stationary group to become motile. Considering Figure 2-1A we can conclude that this is a consequence of the symmetric distribution of the chemotactic agent about the group, that from definition (2.16) we can see clearly results in a zero net force/velocity. Therefore if travelling solutions are to exist then we need to break this symmetry, and this can be achieved in several ways. In numerical simulations we gave the group an initial push or perturbation from its stationary equilibrium for an initial time period, essentially given as an initial velocity of \( c = 0.015 \), we used as a fixed parameter in (2.16) for sufficient time to allow it to saturate to a travelling concentration profile. Subsequently the initial velocity was removed the chemotactic force applied, which if condition (2.22) was upheld, resulted in travelling solutions. Alternatively we considered that asymmetry could be a result of diffusion driven molecular fluctuations in the chemotactic agent, essentially manifesting as noise in the agent’s concentration, implemented numerically as Gaussian noise.

2.2.3 Existence of Travelling Solutions
In our model of chemotactic motion we defined the velocity function to be a function of the concentrations on the boundaries of the segment given as \( c = c_0 \Delta u \), where \( \Delta u \) is taken as the difference over the segment. The free parameter \( c_0 \) models the strength of segments motile reaction to the chemotactic gradient, and in this sense it can be seen as a parameter that determines the existence of travelling solutions. Put more simply the sensitivity of the group to the chemotactic agent has to be greater than some minimum threshold in order for the group to move. Thus let us re-write (2.22) to account for this and re-state the condition on which travelling solutions will emerge as:
\[ c_0 < \frac{2a k_1 \sqrt{d k_1}}{k_2 \left( e^{-\sqrt{\frac{k_1}{d}}} \left( \sqrt{\frac{k_1}{d} + 1} \right) - 1 \right)} \]  

(2.23)

As previously suggested we cannot determine non-trivial solutions of (2.16) because of its implicit and non-linear nature, however we can draw some interesting conclusions from the observation that it appears to be cubic function, as illustrated in Figure 2-2A. Further we can take advantage of this fact and take a cubic approximation to (2.16), and note that since by definition of being odd then even derivatives of an odd function are identically zero,

\[ -f(c) = f(-c) \implies \frac{d^{2n} f(c)}{dc^{2n}} \bigg|_{c=0} = 0, \quad \forall n \in \mathbb{N} \]  

(2.24)

so when we take the cubic Taylor approximation:

\[ f(c) \approx \sum_{n=0}^{3} \frac{f^{(n)}(0)}{n!} (c - 0)^n = g(c) = \frac{g^{(0)}(0)}{0!} (c - 0)^0 + \frac{g^{(1)}(0)}{1!} (c - 0)^1 + \frac{g^{(2)}(0)}{2!} (c - 0)^2 + \frac{g^{(3)}(0)}{3!} (c - 0)^3 \]

and we can show that \( f(0) = f^{(2)}(0) = 0 \) then,

\[ \frac{dg(c)}{dc} \bigg|_{c=0} c + \frac{1}{6} \frac{d^3 g(c)}{dc^3} \bigg|_{c=0} c^3 \]

and we see that quadratic term does not appear, (see Appendix B for derivation)

\[ g(c) = \frac{c_0 k_2}{2a k_1 \sqrt{d k_2}} \left( e^{-\sqrt{\frac{k_1}{d}}} \left( a \sqrt{\frac{k_2}{d} + 1} \right) - 1 \right) \left( 1 - e^{-\sqrt{\frac{k_1}{d}}} \left( a \sqrt{\frac{k_2}{d} (3d - a^2 k_1)} + 1 \right) \right) c^3, \]  

(2.25)

and clearly this has led to a much simpler approximation and further analysis of this in terms of analytical (and numerical) solutions is straightforward. If we now let

\[ A = \frac{c_0 k_2}{2 a k_1 \sqrt{d k_1}}, \quad B = \frac{c_0 k_2}{16 a d k_1 \sqrt{d k_1}} \left( 1 - e^{-\sqrt{\frac{k_1}{d}}} \left( a \sqrt{\frac{k_1}{d} (3 - a^2 k_1)} + 1 \right) \right) \]

and substitute into (2.16) we have:

\[ c \approx -(A + Bc^2)c, \]  

(2.26)

solutions are then given by

\[ (1 - A - Bc^2)c = 0, \]  

(2.27)

or equivalently

\[ B \mu (c^2 - c) = 0; \quad \gamma = \frac{1 - A}{B}, \]  

(2.28)

which gives

\[ c = 0; \quad c = \pm \sqrt{\gamma}; \quad \text{iff } \gamma > 0. \]  

(2.29)
This suggests there are only three possible solutions: travelling symmetric pair or non-travelling/stationary. If this is the case then there must exist a transition or bifurcation point. That is, there exists a point at which a qualitative change in solutions occurs, namely from stationary to travelling (or vice versa), and this point can be found from (2.23) as the condition that $y = 0 \Rightarrow c_0^* = 1/A^*$. This leads us to the conclusion that the behavior being described is prescriptive of a Pitchfork bifurcation.

Pitchfork bifurcations are found in dynamical systems that contain symmetry and where the bifurcation/transition point is a symmetry breaking event, for example the buckling of a axially constrained beam that under a critical load will cause the beam to buckle in one direction or another with equal displacement. In our setting this symmetry breaking event has the physical interpretation of the segment moving positively or negatively on the x-axis. We illustrate this in Figure 2-3A where we plot (2.16) and (2.26) in the parameter plane $c_0$ vs $c$, and we see that for $c_0 > c^*$ there are only stationary solutions, $c = 0$, and when $c_0 < c^*$ travelling solutions emerge in pairs, which corroborates the observations made in the previous section.

![Figure 2-3: Plots illustrating the presence of a pitchfork bifurcation in the chemotaxis equation.](image)

We have found that for certain values of parameter $c_0$ we have 3 travelling wave solutions and for others only one and transition between these two cases is described by pitch-fork bifurcation. Therefore concerning the stability of all these solutions there are only two possibilities: the bifurcation occurs sub-critically, i.e. single solution $c = 0$ is unstable before bifurcation and stable afterwards and two other traveling wave solutions are unstable; or it occurs super-critically when the single solution $c = 0$ is stable before bifurcation and unstable afterwards while two other traveling wave solutions are stable (see Figure 2-3B-C). The important consequence of this is that if we can determine the stability of stationary solution $c = 0$ for a certain value of $c_0$ we can deduce on stability of all traveling wave solutions for any value of $c_0$. It is intuitively clear that if the protein produced in the segment is an attractor ($c_0 > 0$) then the segment should be stationary, since any shift of segment with respect to the $u$-profile will results to formation of forces pushing it backward. Therefore stationary solution $c = 0$ is stable when $c_0 > 0$ meaning that it will be stable for all values of $c_0$ satisfying inequality (2.23). Therefore it becomes unstable.
when (2.23) is not satisfied while two traveling solutions which appear through pitch-fork bifurcation are stable (supercritical bifurcation).

### 2.2.4 Stability of a Chemotactically Moving Group

Let us address the stability of traveling wave solutions in a more thorough way. When we speak of stability we are referring to how solutions are affected upon small perturbations in the location of the segment and/or \( \mathbf{u} \) profile of traveling wave solutions. In our problems stability of traveling wave solutions is a question of whether the velocity of chemotactically moving segment will restore after perturbations.

We begin by making the observation from our simulations with a segment moving with prescribed velocity \( c \) (non-chemotactic motion): no matter what was initial \( \mathbf{u} \) profile in simulations it saturates to the profile described by the equation (2.11). This could be stated as the stability of the \( \mathbf{u} \) profile, described by the equation ((2.11)). Furthermore if the \( \mathbf{u} \) profile of the travelling wave solution for a chemotactically moving segment \( \mathcal{C} = f(c) \) is perturbed then at any moment of time the \( \mathbf{u} \) profile will tend to the profile described by the equation (2.11) that is \( c(t) \rightarrow f(c(t)) \), where the velocity is given by instantaneous velocity of the moving segment. We should note that at the same time the velocity of the moving segment will change according \( c = c_0 \Delta \mathbf{u}(t) \). From Figure 2-2 and Figure 2-3 we know that the velocity of traveling wave solution \( c \) is defined by the value of \( c_0 \). However we cannot state that instantaneous velocity \( c \) satisfies (2.16), since we have already shown that it is the value of the parameter \( c_0 \) that uniquely determines \( c \), and not the contrary. In other words if we prescribe a value of \( c \) that does not satisfy (2.16) then by extension it does not correspond to any value of \( c_0 \). The logical conclusion to this argument is that the profile of the chemotactic agent is not a travelling wave (that is non-steady state solution) and the profile is changing with time implying:

\[
\begin{align*}
    c & \neq f(c), \\
    c & \rightarrow f(c),
\end{align*}
\]

and further:

\[
\begin{align*}
    c & \neq f(c), \\
    c & \rightarrow f(c),
\end{align*}
\]

so that it will correspond to the current value of the parameter \( c_0 \).

This conclusion is borne out in numerical simulations, where we considered a steady state solution that had speed \( c^* \) and then perturbed the segment by a small amount resulting in a new speed \( c_{new} \) proportional to the change in \( \Delta \mathbf{u} \). However the segment would always return to its starting velocity \( c^* \) as illustrated in Figure 2-4. This result demonstrates two things: a) clearly the solution is asymptotically stable and b) we can assume the pitchfork bifurcation is super-critical (Figure 2-3C), for if it were not then we should not have any stable travelling solutions.
Figure 2-4: Perturbation of a segment for a steady state solution. If we assume that we have a solution for a segment of width $a$ travelling with speed $c^*$, and then we perturb the segment $\pm 2$ grid units (translates to $1/25$th of the segment’s size) then the segment will attain a new speed $c_{new}$ for a short period of time proportional to the change in $\Delta u$ on its boundary, but then return back to it initial speed $c^*$.

For formal analysis of stability of traveling wave solution which moves with velocity $c^*$ let us assume that after perturbation it moves with speed $c = c^* + \delta c$ and that the rate of its change is some function of the velocity $c$:

$$\frac{dc}{dt} = F(c). \quad (2.32)$$

If we can show that $c \to c^*$ then the traveling wave solution is stable and unstable otherwise. Let us consider the linear approximation of the above equation:

$$\frac{dc}{dt} = A + Bc$$

Since $\frac{dc}{dt} = 0$ when $c = c^*$ we conclude that $A = -Bc^*$ and therefore:

$$\frac{dc}{dt} = B(c - c^*)$$

Now we shall find out what is a sign of factor $B$ (note that negative $B$ would mean that $c \to c^*$ and therefore the traveling wave solution is stable). To do so we note that $c \to f(c)$ (see (2.31)) and therefore:

$$c = c^* + \delta c \Rightarrow f(c^* + \delta c) \Rightarrow f(c^*) + f'(c^*)\delta c,$$

$$\Rightarrow c^* + \delta c \to c^* + f'(c^*)\delta c$$

$$\Rightarrow \delta c \to f'(c^*)\delta c.$$  

And so for $\delta c > 0$ (or $c > c^*$) and $f'(c^*) > 1$, $\delta c$ is increasing (RHS is above LHS) from which we conclude that $B$ is positive if $f'(c^*) > 1$ and negative if $f'(c^*) < 1$. Therefore the traveling wave solution is stable if $f'(c^*) < 1$. If we now consider Figure 2-2A, we stated that intersection between the dashed line and profiles $f(c)$ (solid black lines) are travelling wave solutions corresponding to $c^* = f(c^*)$, which clearly show that at these intersection points $f'(c^*) < 1$.

### 2.2.5 Concentration profile of an Externally Produced Chemotactic Agent

In the previous section we considered a homogenous group that is attracted or repelled by a diffusing chemotactic agent it produces, and showed that the dynamical behaviour, in terms of travelling solutions and governed by (2.16), is prescriptive of a Pitchfork bifurcation as illustrated in Figure 2-3A. In this section we will consider the same homogenous group, however we shall consider that the chemotactic agent is produced by a surrounding population of cells, rather than the group itself. The analysis that
follows will mirror the previous sections for an internally produced chemical, beginning with the production of the diffusing agent that is equivalent to (2.2), re-produced here for convenience,

$$\frac{\partial u}{\partial t} = D \Delta u + c(\delta u; c_0) \frac{\partial u}{\partial x} - k_1 u + p(x); \quad k_1, k_2, D \in \mathbb{R}, \ \Delta = \text{laplacian} \quad (2.34)$$

Where $c(\delta u; c_0)$ is the velocity of motion depending on $\delta u$ which is the difference in $u$ on the left and right boundaries of the segment and $c_0$ is chemotactic forcing parameter measuring the sensitivity of the group to the chemotactic gradient. However the kinetics in (2.2) prescribed by (2.1) are changed to describe external production:

$$p(x) = \begin{cases} 0 & \alpha(t) < x < \beta(t) \\ k_2 & \text{otherwise} \end{cases} \quad (2.35)$$

To accommodate this change in the production characteristics we can redefine the piecewise system given by (2.5) and describing travelling wave solutions in the following way,

$$0 = \begin{cases} D \Delta u + cu_x - k_1 u + k_2, & x \leq \alpha \\ D \Delta u + cu_x - k_1 u, & \alpha \leq x \leq \beta \\ D \Delta u + cu_x - k_1 u + k_2, & x \geq \beta \end{cases} \quad (2.36)$$

where the production term in the diffusion equation, $k_2$, is now expressed outside of the group $|x| > |\beta - \alpha|$, and zero within, and whose general solution can be determined equivalently to (2.6) as:

$$u = \begin{cases} Ae^{\lambda_1 x} + Be^{\lambda_2 x} + \frac{k_2}{k_1}, & x < \alpha \\ Ce^{\lambda_1 x} + De^{\lambda_2 x}, & \alpha \leq x \leq \beta \\ Fe^{\lambda_1 x} + Ge^{\lambda_2 x} + \frac{k_2}{k_1}, & x \geq \beta \end{cases} \quad (2.37)$$

Again to determine the unknown coefficients we apply the same boundary and continuity conditions defined by (2.8)-(2.10)), which leads us to the solution of (2.37) for a homogeneous group with an externally produced chemotactic agent:

$$u = \begin{cases} -\frac{k_2}{k_1(\lambda_1 - \lambda_2)}(\lambda_2 e^{\lambda_1(x-\beta)} - e^{\lambda_1(x-\alpha)}) - (\lambda_1 - \lambda_2), & x \leq \alpha \\ -\frac{k_2}{k_1(\lambda_1 - \lambda_2)}(\lambda_2 e^{\lambda_1(x-\beta)} - \lambda_1 e^{\lambda_2(x-\alpha)}), & \alpha \leq x \leq \beta \\ -\frac{k_2}{k_1(\lambda_1 - \lambda_2)}(\lambda_1 e^{\lambda_2(x-\beta)} - e^{\lambda_2(x-\alpha)}) - (\lambda_1 - \lambda_2), & x \geq \beta \end{cases} \quad (2.38)$$

for which we can illustrate both stationary (Figure 2-5A) and travelling (Figure 2-5B) frames.

Figure 2-5: Plots of stationary and travelling solutions for a homogenous group with an externally produced chemotactic agent. The group is represented by the grey shaded segment of width $a$ that reacts to the production of an externally produced chemotactic agent, $u$, that in a stationary solution (A) displays a symmetric profile with respect to the group. In a travelling solution (B) the group moves to the right on the spatial $x$-axis mirroring the behaviour of an internally produced chemical in Figure 2-2.
As with (2.7) the system (2.38) displays a symmetric distribution with respect to the group in a stationary solution, with its minimum concentration located centrally within the group. And further for a travelling solution the minimum concentration lags behind this central position proportional to the velocity of the frame given by \( c \). Again we can determine the location of the minimum by evaluating the spatial derivative within the group in exactly the same manner for (2.13):

\[
\frac{du(x)}{dx} = \frac{d}{dx} \left( \frac{k_2}{k_1(\lambda_1-\lambda_2)} \left( \lambda_1 e^{\lambda_2(x-\alpha)} - \lambda_2 e^{\lambda_1(x-\beta)} \right) \right) \\
= \frac{k_2}{k_1(\lambda_1-\lambda_2)} \left( \lambda_2 \lambda_1 e^{\lambda_2(x-\alpha)} - \lambda_1 \lambda_2 e^{\lambda_1(x-\beta)} \right)
\]

(2.39)

where again \( u(x) \) is the concentration profile inside the group and the condition for (2.39) to be identically zero can only be true when \( e^{\lambda_2(x+\frac{a}{2})} - e^{\lambda_1(x-\frac{a}{2})} = 0 \), which is identical to the condition required for the derivation of (2.13). Thus the derivation and expression of the location of the maximum, \( x_{max} \), for (2.38) is also identical.

### 2.2.6 Motion Due To Chemotaxis

Let us now derive the chemotaxis function for the solution (2.38). We will again follow the same derivation as for (2.8). As before the boundary conditions are such that the expressions in (2.38) we employ to take the difference in concentration over the entire group can be chosen arbitrarily, so again we follow the derivation for (2.16):

\[
c(c; c_0) = c_0 \frac{u(\beta) - u(\alpha)}{a} \\
= \frac{k_2}{k_1(\lambda_1-\lambda_2)} \left( \lambda_1 e^{\lambda_2(\frac{a}{2} - \frac{\alpha}{2})} - \lambda_2 e^{\lambda_2(\frac{\alpha}{2} + \frac{a}{2})} \right) + \frac{k_2}{k_1(\lambda_1-\lambda_2)} \left( \lambda_2 e^{\lambda_1(\frac{\alpha}{2} - \frac{a}{2})} - \lambda_1 e^{\lambda_1(\frac{\alpha}{2} + \frac{a}{2})} \right) \\
= \frac{c_0 k_2}{a k_1(\lambda_1-\lambda_2)} \left\{ \lambda_1 e^{\lambda_2(\frac{a}{2} + \frac{a}{2})} - \lambda_1 e^{\lambda_2(\frac{\alpha}{2} + \frac{a}{2})} - \lambda_2 e^{\lambda_1(\frac{\alpha}{2} + \frac{a}{2})} + \lambda_2 e^{\lambda_1(\frac{\alpha}{2} + \frac{a}{2})} \right\} \\
= \frac{c_0 k_2}{a k_1(\lambda_1-\lambda_2)} \left\{ \lambda_1 - \lambda_1 e^{a\lambda_2} + \lambda_2 e^{-a\lambda_1} + \lambda_2 \right\}
\]

leading to the chemotaxis function for a homogenous group with an externally produced chemotactic agent:

\[
c(c; c_0) = \frac{k_2 c_0}{a k_1(\lambda_1-\lambda_2)} \left( \lambda_1 (1 - e^{a\lambda_2}) + \lambda_2 (1 - e^{-a\lambda_1}) \right)
\]

(2.40)

Clearly (2.40) is identical to (2.16) and so we conclude that while the production dynamics of the chemotactic agent have changed from internal to external (or vice versa), the dynamical behaviour of the group of cells in terms of travelling solutions are equivalent.

### 2.3 Model for the Heterogeneous Migrating Group

In this section we consider the case of a migrating group that is composed of two cell types. We assume that one of these types is attracted or repelled by the gradient of a locally produced, diffusing
chemotactic agent, and another type within the same group that is not. As before the dynamics of the diffusing chemotactic agent is described by (2.1), however we will need to redefine the domain of production as given by (2.2) to account for the inhomogeneity of the group. Let us assume once again that the group is defined in the interval \((\alpha, \beta)\). Further for simplicity let us assume that the group is divided equally down the vertical axis, composing two halves defined symmetrically about the origin. We consider again that production can be internal or external to the group, however we make the distinction that one half of the group reacts chemotactically while the other half does not. As in the preceding section, we shall proceed by dividing the analysis into scenarios of external and internal production and investigate the permutations of the reacting half of the group and the dynamical systems that result.

In the cases we have analysed so far we have assumed the group is composed of a single population of cells, all reacting equivalently to the chemotactic agent. However in this variation of the model we have two sub-populations with one part reacting actively to the chemotactic agent and the other part is producing. For clarity, let us term the part that is producing the chemotactic agent as \(\mathcal{P}\) and the part that is reacting to the agent as \(\mathcal{R}\). Clearly then \(\mathcal{P}\) or \(\mathcal{R}\) can be either to the left or right of the origin. Therefore let us redefine the definition of \(\mathcal{P}\) and \(\mathcal{R}\) to take into account spatial positioning by defining:

\[
\mathcal{P} = \begin{cases} 
\mathcal{P}_l & \alpha < x < (\beta + \alpha)/2 \\
\mathcal{P}_r & (\beta + \alpha)/2 < x < \beta, 
\end{cases} 
\tag{2.41}
\]

and

\[
\mathcal{R} = \begin{cases} 
\mathcal{R}_l & \alpha < x < (\beta + \alpha)/2 \\
\mathcal{R}_r & (\beta + \alpha)/2 < x < \beta. 
\end{cases} 
\tag{2.42}
\]

2.3.1 Concentration profile of an Internally Produced Chemotactic Agent

For now we shall only consider \(\mathcal{P}\) and delay consideration of \(\mathcal{R}\) until the next section when we consider chemotactic motion. In terms of concentration profiles then, there are two configurations given by (2.41): one profile when \(\mathcal{P} = \mathcal{P}_l\) and another profile when \(\mathcal{P} = \mathcal{P}_r\). These two configurations give rise to two reaction-diffusion systems, which for \(\mathcal{P} = \mathcal{P}_l\):

\[
p(x) = \begin{cases} 
k_2 & \alpha < x < \frac{\beta + \alpha}{2} \\
0 & \alpha > x > \frac{\beta + \alpha}{2}.
\end{cases} 
\tag{2.43}
\]

\[
u = \begin{cases} 
k_2 \lambda_2 \left( e^{\lambda_1 (x-((\beta + \alpha)/2))} - e^{\lambda_2 (x-\alpha)} \right) + k_1 (\lambda_1 - \lambda_2), & x \leq \alpha \\
\frac{k_2}{k_1 (\lambda_1 - \lambda_2)} \left( \lambda_2 e^{\lambda_1 (x-((\beta + \alpha)/2))} - \lambda_1 e^{\lambda_2 (x-\alpha)} + (\lambda_1 - \lambda_2) \right), & \alpha \leq x \leq (\beta + \alpha)/2 \\
\frac{k_2 \lambda_2 \left( e^{\lambda_2 (x-((\beta + \alpha)/2))} - e^{\lambda_2 (x-\alpha)} \right)}{k_1 (\lambda_1 - \lambda_2)}, & x \geq (\beta + \alpha)/2
\end{cases} 
\tag{2.44}
\]

and for \(\mathcal{P} = \mathcal{P}_r\):

\[
p(x) = \begin{cases} 
1 & \frac{(\beta + \alpha)}{2} < x < \beta \\
0 & \beta < x < \frac{(\beta + \alpha)}{2}.
\end{cases} 
\tag{2.45}
\]
While it is not necessarily clear that the systems form a symmetric pair, given by rotating either about the vertical axis, which can be seen more readily if consider plots as illustrated in Figure 2-6. Beyond this symmetry the concentration profiles are equivalent to those illustrated in Figure 2-1A for the homogenous system \((2.11)\), with the only difference being how the concentrations are distributed about the mid-point of the group. However this will play an important role when we consider chemotactic motion in the next section when we consider the permutations of \(P_{l,r}\) and \(R_{l,r}\).

\[
\begin{aligned}
    u &= \begin{cases}
        \frac{k_2 \lambda_2 \left(e^{\lambda_1(x-\beta)} - e^{\lambda_1(x-(\beta+\alpha)/2)}\right)}{k_1(\lambda_1 - \lambda_2)}, & x < (\beta + \alpha)/2 \\
        \frac{k_2 \lambda_2 \left(e^{\lambda_1(x-\beta)} - \lambda_1 e^{\lambda_2(x-(\beta+\alpha)/2)} + (\lambda_1 - \lambda_2)\right)}{k_1(\lambda_1 - \lambda_2)}, & (\beta + \alpha)/2 < x < \beta \\
        \frac{k_2 \lambda_1 \left(e^{\lambda_2(x-\beta)} - e^{\lambda_2(x-(\beta+\alpha)/2)}\right)}{k_1(\lambda_1 - \lambda_2)}, & x > \beta
    \end{cases} \tag{2.46}
\end{aligned}
\]

\(\text{Figure 2-6: Concentration profiles for an heterogeneous group. We divide the group evenly down the vertical axis and assume that half of the group is producing the chemotactic agent } P. \text{ In terms of } P \text{ there are two symmetric variations: the left-hand part of the group is producing the agent we call } P_{l}(A) \text{ or the right-hand part of the group is producing the agent } P_{r}(B).\)

\subsection*{2.3.2 Motion Due To Chemotaxis}

In line with the previous analysis we will derive the chemotaxis function from \((2.15)\) and substitute from the heterogeneous systems \((2.43)-(2.44)\) and \((2.45)-(2.46)\). Unlike the previous models of chemotaxis we have considered so far, the part of the group that is reacting to the chemotactic agent is not necessarily the same as the part that is producing it. To differentiate these scenarios we introduced in the last section the quantities \(P\) and \(R\), defined respectively in \((2.41)\) and \((2.42)\), with the rather obvious assignment of \(P\) for the part that produces the chemotactic agent and \(R\) for the part that reacts to it. If we now recall that the chemotactic force can either act to repel or attract \(R\), and noting that we can either have \(P = R\) or \(P \neq R\), then there are eight cases we can consider.

Consider now the case when \(P \neq R\) we have four permutations: two when \(P = P_{l}\) and \(R = R_{r}\) and two when \(P = P_{r}\) and \(R = R_{l}\) for chemo-repulsion or chemo-attraction, illustrated in Figure 2-7. Clearly the distinguishing factor in this case, is that the part of the group that reacts to the chemotactic agent is different from the part that produces it. However unlike the previous analysis there is an implied asymmetry in the chemotaxis function in this case, since \(R\) will always be situated within an asymmetric
distribution of the agent. Practically this implies that for \( c_0 \neq 0 \) we will always have travelling solutions, or in other words, there will always be one trivial stationary solution when \( c_0 = 0 \) or there will always be travelling solutions when \( c_0 \neq 0 \). This is in direct contrast to the dynamical behaviour of previous chemotaxis functions, were dual travelling solutions emerged at a bifurcation point when \( 0 > c_0 < c_0^* \), were \( c_0^* \) is the expressed as the right-hand side of inequality (2.23).

Figure 2-7: Concentration profiles of travelling wave solutions for an heterogeneous system when \( \mathcal{P} \neq \mathcal{R} \). When the group is divided equally into two parts about the origin with one part producing the chemotactic agent, \( \mathcal{P} \) (grey shaded rectangle), and another being chemo-repelled or chemo-attracted to the gradient of the agent, \( \mathcal{R} \), then we have four permutations when \( \mathcal{P} \neq \mathcal{R} \): two when \( \mathcal{P} = \mathcal{P}_1 \) and \( \mathcal{R} = \mathcal{R}_r \) and two when \( \mathcal{P} = \mathcal{P}_r \) and \( \mathcal{R} = \mathcal{R}_1 \) for attraction or repulsion. However it can be shown that there only exists two distinct cases that, since \( (\mathcal{A}, \mathcal{B}) \) and \( (\mathcal{C}, \mathcal{D}) \) are merely symmetric variations of the same problem and thus \( (\mathcal{A}, \mathcal{B}) \) is qualitatively and quantitatively equivalent to \( (\mathcal{C}, \mathcal{D}) \).

Since in this case \( \mathcal{P} \neq \mathcal{R} \) we have the choice to substitute from either (2.43) and (2.44) or (2.46), that is the group is composed of either \( \mathcal{P}_1 \) and \( \mathcal{R}_r \) or \( \mathcal{P}_r \) and \( \mathcal{R}_1 \). Let us first consider the case of \( \mathcal{P}_1 \) and \( \mathcal{R}_r \), and derive the associated chemotaxis function, noting that we can only substitute for \( u_3 \) from (2.44), then:

\[

c = c_0 \left( \frac{u(\beta)-u((\beta+\alpha)/2)}{[\beta-(\beta+\alpha)/2]} \right) = \frac{c_0 k_2 \lambda_1}{k_1 (\lambda_1 - \lambda_2) [\beta - (\beta + \alpha)/2]} \left( e^{\lambda_2 (\beta - (\beta + \alpha)/2)} - e^{\lambda_2 (\beta - \alpha)} - e^{\lambda_2 (\beta + (\beta + \alpha)/2)} + e^{\lambda_2 (\beta + \alpha)} \right)

\]

and when the group is symmetric with respect to the vertical axis:

\[

c = \frac{2c_0 k_2 \lambda_1}{ak_1 (\lambda_1 - \lambda_2)} \left( e^{\lambda_2 (\frac{\alpha}{2} - 0)} - e^{\lambda_2 (\frac{\alpha}{2} - (\frac{\alpha}{2})^2)} - e^{\lambda_2 (0 - \frac{\alpha}{2})} + e^{\lambda_2 (0 - (\frac{\alpha}{2})^2)} \right) = \frac{2c_0 k_2 \lambda_1}{ak_1 (\lambda_1 - \lambda_2)} \left( 2 e^{\frac{\alpha^2}{4}} - e^{\lambda_2} - 1 \right)

\]

and we find the chemotaxis function:

\[

c = - \frac{2c_0 k_2 \lambda_1}{ak_1 (\lambda_1 - \lambda_2)} \left( e^{\frac{\alpha^2}{4}} - 1 \right)^2.
\] (2.47)

Clearly the sign of \( c(a; c_0) \) is given by difference in the concentration over \( \mathcal{R} \). If we neglect the contribution from \( c_0 \) this difference will be negative implying the group will have the tendency to move left-wise (Figure 2-8B). More generally it can be shown that \( \mathcal{R} \) is attracted by \( \mathcal{P} \) when \( \mathcal{P} \neq \mathcal{R} \). It should also be clear that \( c_0 \neq 0 \Rightarrow c(c; c_0) \neq 0 \) and conversely \( c_0 = 0 \Rightarrow c(c; c_0) = 0 \), which we can show by noting that \( \lambda_1, (\lambda_1 - \lambda_2) \) and the model parameters \( a, d, k_1 \) and \( k_2 \) are strictly positive and clearly
Therefore $c$ only changes sign when $c_0$ changes sign, and further it follows that (2.47) admits travelling solutions whenever $c_0 \neq 0$, with direction of motion given by the sign of $c_0$, and stationary solutions when $c_0 = 0$. We can illustrate these points by considering Figure 2-8.

Figure 2-8: Motile dynamics of a heterogeneous group when $\mathcal{P} \neq \mathcal{R}$. Concentration profiles in a travelling frame for repulsion (A) and attraction (B) illustrate the group moving either positively or negatively on the x-axis given by the sign of $c_0$. Unlike the homogenous system, the heterogeneous system will always exhibit travelling solutions when $c_0 \neq 0$ and these will appear in pairs, with a trivial solution when $c_0 = 0$. However the implied asymmetry brought about by the difference in concentration profiles over $\mathcal{R}$, manifest as asymmetries in the velocity of the group for attraction (solid black line) and repulsion (dashed black line) (C). We can summarise this behaviour by plotting solutions in the parametric plane $c_0$ vs $c$ (D) were repelling solutions (dashed black line) exist when $c_0 < 0$ and attracting solutions (solid black line) when $c_0 > 0$, and a trivial stationary solution when $c_0 = 0$. Model parameters where chosen such that: $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$ and $\alpha = 5.0$.

Figure 2-8(A-B) shows the concentration profiles for the system (2.44) in a travelling frame ($c_0 \neq 0$) for chemo-repulsion, $c_0 < 0$, (A) and chemo-attraction, $c_0 > 0$, (B). Travelling solutions can be seen to exist by using the same argument as for (2.19) and (2.20), by plotting the left and right-hand sides of (2.47) on the same axis, and again solutions are given by intersections that appear in pairs (Figure 2-8C). Unlike the homogenous case we observe that the solutions are asymmetric about the vertical, that is the velocity of the group, $f(c)$, is greater for repulsion than for attraction. The observed asymmetry is a result of the definition of the chemotaxis function, which is given as the difference in concentration over $\mathcal{R}$. Depending on the direction of motion, i.e. attraction or repulsion, we can see that the difference is either taken at the front or the rear of what can be seen as a travelling wave. Therefore one would expect the difference to be more precipitous at the front of the wave than at the rear, thus implying a differential in the gradients of the pairs of travelling solutions.

However it is possible to show that the asymmetry in (2.47) can be overcome, and further that attraction or repulsion can be maximised by scaling the size of the group. We can see this more clearly if re-write (2.47) to express the dependence of travelling solutions on the group’s size,

\[
c = -\frac{2c_0 k_2 \lambda_1}{ak_1(\lambda_1 - \lambda_2)} \left( e^{\frac{a\lambda_2}{2}} - 1 \right)^2,
\]  

(2.48)
and plot solutions in plane $c$ vs $a$. Such solutions, as illustrated in Figure 2-9A, are shown as profiles for attraction (solid) and repulsion (dashed), demonstrating how the velocity of the group varies as the size of the group is increased symmetrically.

![Figure 2-9: Velocity characteristics of the group.](image)

The chemotaxis equation (2.48) for an heterogeneous group when $\mathcal{P} \neq \mathcal{R}$ exhibits asymmetry in its velocity between the forces of attraction and repulsion. However we can show that if we consider the velocity to be dependent on the size of the group and plot solutions to (2.48) for symmetric variations in its size $a$ (A), then we can show that it is possible for these force to be not only equal, $a_2$, but we can also be maximised for attraction, $a_1$, or repulsion $a_2$. Further if we vary the size of the group asymmetrically, that is vary $\mathcal{P}$ while holding $\mathcal{R}$ (or vice versa) by moving the boundary between them (B), then the velocity of the group changes equivalently for a change of $\mathcal{P}$ or $\mathcal{R}$, illustrating a symmetric distribution. Model parameters where chosen such that: $k_1 = 0.00025, k_2 = 2k_1, D = 0.5$ and $a = 50$.

When the size of the group is $a < a_2$ then the force of repulsion is greater than that of attraction and maximises for $a = a_1$ until $a = a_2$ when both forces become equal. Then as $a > a_2$ the force attraction is greater than that of repulsion until it reaches its maximum at $a = a_3$ beyond which both forces decay. Therefore we can show qualitatively that we can maximize (2.47) for attraction or repulsion and further the two forces can be equal. Lastly, if we vary the size of the group asymmetrically, that is if we vary the size of $\mathcal{P}$ while holding $\mathcal{R}$ constant (or vice versa) (Figure 2-9B) by moving the boundary between $\mathcal{P}$ and $\mathcal{R}$, then the velocity of the group varies symmetrically.

### 2.3.3 Concentration profile of an Externally Produced Chemotactic Agent

In this section we again consider a heterogeneous group of cells with only one part reacting chemotactically and $\mathcal{P} \neq \mathcal{R}$. However we now assume that the agent is produced by a surrounding population of cells and not the cells in the group, and so $\mathcal{P}$ in this case can be defined as $\mathcal{P} = \mathcal{P}_e = \{x: x < a, x > \beta\}$ where the subscript $e$ denotes the external population of cells. The distribution of the agent, $u$, for this configuration can be seen to be distributed symmetrically about the stationary group, implying the concentrations at the left ($\alpha$) and right ($\beta$) boundaries of group are equal and will have a minimum concentration located at the group's centre with coordinate $x_c = (\beta + \alpha)/2$, illustrated in Figure 2-10.
Figure 2-10: Concentration profiles of an externally produced chemotactic agent for a heterogeneous group. The production domain of the chemotactic agent is produced by a surrounding population of cells, $P_e$, with a profile distributed symmetrically about the group with equal concentrations at the left and right boundaries given respectively as $\alpha$ and $\beta$ with a minimum concentration located at the centre of the group with coordinate $x = (\alpha + \beta)/2$. The inhomogeneity of the group is once again given as a bissection with one part reacting to the agent, which can either be on the right-hand $\mathcal{R} = \mathcal{R}_r$ or left-hand.

The composition of the group, in terms of inhomogeneity, is once again given by the bissection of the group about the centre point of the group. This bissection defines two equally sized halves with one part chemotactically sensitive to the gradient of the chemotactic agent, while the other part is not, identified by the term $\mathcal{R}$.

2.3.4 Motion Due To Chemotaxis.

As before we define $\mathcal{R} = \mathcal{R}_{lr}$ to define whether the left or right part of the group is reacting to the chemotactic agent described by (2.38), and analyse the motile dynamics by deriving the chemotaxis equation from (2.44) for $\mathcal{R} = \mathcal{R}_l$ and (2.46) for $\mathcal{R} = \mathcal{R}_r$. Given that there are two possible configurations for the group and that there are two possibilities for forces, chemo-attraction and chemo-repulsion, then there are four cases we can consider. Specifically for attraction when $\mathcal{R} = \mathcal{R}_l$ and repulsion when $\mathcal{R} = \mathcal{R}_r$. As before the chemotaxis function is essentially given by the difference in concentration over the reacting part of the group, and we again consider a stationary solution (Figure 2-10) and further assume that $c_0 > 0$, then for $\mathcal{R}_r$ we should expect the group to move right-wise and for $\mathcal{R}_l$ left-wise; in other words the group will be attracted for $\mathcal{R}_r$ and $\mathcal{R}_l$ when $c_0 > 0$. Further since the difference in attraction or repulsion is a change of sign of $c_0$, then appealing to intuition we can see that there only two qualitatively unique cases for travelling solutions (consider Figure 2-11) due to symmetry.

Figure 2-11: Travelling solutions of a heterogeneous group for an externally produced chemotactic agent. For travelling solutions there are four possible outcomes that depend on which part of the group is reacting, $\mathcal{R}$, and whether it is attracted or repelled by the concentration gradient of the chemotactic agent (solid black profile). For $\mathcal{R} = \mathcal{R}_l$ the group will move left-wise for repulsion (A) and right-wise (B) for attraction, whereas for $\mathcal{R} = \mathcal{R}_r$ the opposite is true and the group will move left-wise (C) for attraction and right-wise for repulsion (D).
To see this more clearly let us derive the chemotaxis functions for $R_1$ and $R_2$ by substituting $u(x)$ from (2.38) for $u(x)$ in (2.44) we find:

$$c(c; a, c_0) = c_0 \frac{u((\beta + a)/2) - u(\alpha)}{[(\beta + a)/2 - a]} = -c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(x - \beta)} - \lambda_1 e^{\lambda_2(x - \alpha)}\right) \left|_{x=a} \right. + \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(x - \beta)} - \lambda_1 e^{\lambda_2(x - \alpha)}\right) \left|_{x=\alpha} \right.$$

$$= -c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(\beta + a)/2} - \lambda_1 e^{\lambda_2(\beta + a)/2} - \lambda_2 e^{\lambda_1(\alpha - \beta)} + \lambda_1 e^{\lambda_2(\alpha - \alpha)}\right)$$

which for a symmetrically posed group of width $\alpha$:

$$= -c_0 \frac{k_2}{k_1(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(\beta + a)/2} - \lambda_1 e^{\lambda_2(\beta + a)/2} - \lambda_2 e^{\lambda_1(\alpha - \beta)} + \lambda_1 e^{\lambda_2(\alpha - \alpha)}\right)$$

we find the chemotaxis equation for an heterogeneous group when $R = R_1$:

$$c_1(c; a, c_0) = -2c_0k_2 \frac{(\beta + a)}{(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(\beta + a)/2} - \lambda_1 e^{\lambda_2(\beta + a)/2} - \lambda_2 e^{\lambda_1(\alpha - \beta)} + \lambda_1 e^{\lambda_2(\alpha - \alpha)}\right).$$

(2.49)

And substituting $u(x)$ from (2.38) for $u(x)$ in (2.46), we can show by an equivalent derivation when $R = R_2$ that:

$$c_2(c; a, c_0) = -2c_0k_2 \frac{(\beta + a)}{(\lambda_1 - \lambda_2)} \left(\lambda_2 e^{\lambda_1(\beta + a)/2} - \lambda_1 e^{\lambda_2(\beta + a)/2} - \lambda_2 e^{\lambda_1(\alpha - \beta)} + \lambda_1 e^{\lambda_2(\alpha - \alpha)}\right).$$

(2.50)

Now assuming that there are only two equivalent travelling solutions, then is should be the case that (2.49) and (2.50) can be shown to be equivalent for attraction and repulsion respectively. That is it should be true that there are conditions on the model parameters that satisfy $c_1(c; a, c_0) = c_2(c; a, c_0)$. From direct observation we can see this can only be true when $\lambda_{2,1} = -\lambda_{2,1}$, implying $c = -c$ and $c_0 = -c_0$, that is they are equivalent for a change in chemotactic response and direction of motion. Therefore $R$ for attraction is equivalent to $R$ for repulsion and vice versa. For either of these cases we wish to analyse the existence of, and conditions for, travelling solutions to emerge. However we cannot directly determine an analytical derivation given their implicit and non-linear nature, and thus we shall fall back on qualitative techniques.

To determine existence and conditions for travelling solutions we could follow the analysis outlined in 2.2.2 for a homogeneous group. In that section we demonstrated that solutions can be seen to exist if we plot the left and right-hand sides of the chemotaxis equation against each other on the same axis, and that the condition for such travelling solutions is given by taking the derivative at $c = 0$ and solving for the bifurcation parameter $c_0$. It should be clear however, that unlike the homogeneous case, travelling solutions will always exist for a heterogeneous group, regardless of how or where the chemotactic agent is produced for non-zero $c_0$ as a result of the asymmetric distribution of the agent over $R$. The implication of this is that the chemotactic functions are non-zero for $c = 0$, and therefore the
derivative condition suggested in 2.1.2 is meaningless for an heterogeneous group. To see this more clearly let us re-write (2.49) and (2.50) so as to plot the left and right-hand sides against each other on the same axis:

\[
y_1 = c l: \quad y_2 = -\frac{2c_0 k_2}{\alpha k_1 (\lambda_1 - \lambda_2)} \left( \lambda_{2,1} \left( e^{\frac{-a \lambda_{1,2}}{2}} - e^{-a \lambda_{1,2}} \right) + \lambda_{1,2} \left( 1 - e^{\frac{-a \lambda_{2,1}}{2}} \right) \right)
\]  

(2.51)

and let us assume that \( c_0 > 0 \); the reacting part of the group will be attracted to the chemotactic agent. Clearly travelling solutions in (2.51) correspond with the equality \( y_1 = y_2 \), which we can see respectively as graphical intersections between the solid or dashed profiles and the dotted line in Figure 2-12C. The two profiles for each case are for two values of the model parameter \( c_0 \), where we see that in both cases at least one solution will always exist when \( f(c) = 0 \), with two further solutions emerging if \( c_0 \) is of sufficient magnitude.

![Figure 2-12: Attraction for a heterogeneous with the chemotactic agent produced externally will always produce travelling solutions. A-B: if we assume that \( c_0 > 0 \), that is chemo-attraction, then regardless of which sub-population of the group is reacting, \( R \), the difference in concentration over \( R \) will always be non-zero due asymmetry with respect to the chemotactic agent’s profile where we see two cases: one where \( R \) is on the right (A) and \( R \) is on the left (B). C: We illustrate here two pairs of profiles for two values of the parameter \( c_0 \) to demonstrate the effect of increasing its value, where we see that there will always be one solution (intersection of solid black line and small dotted line) and two further solutions emerging when \( c_0 \) increases with one stable and the other unstable. Further we note that there is a case when \( f(c) = 0 \) and \( c \neq 0 \), that we can interpret as limiting cases when the velocity, \( \alpha \), such that the difference in concentrations of either side of \( R \) becomes equal; we will consider this case in the proceeding section.

2.3.5 Generalisation of the Heterogeneous Model

In the preceding section we again found that the condition for travelling solutions is dependent on the prescription of \( c_0 \), but as we can see from Figure 2-12C the derivative condition of 2.1.2 does not hold. Moreover we cannot find a condition on \( c_0 \) that is independent of the variable \( c \), and thus we cannot derive any meaningful results in this respect. However we can make some interesting observations regarding Figure 2-12C, where we see that the right-hand sides of (2.51) are zero for non-zero \( c \) on the left-hand side. We can interpret these two points as limiting cases when the chemotactic mechanism is attraction. To understand this, let us first reconsider the definition of the chemotaxis function given by (2.15), reproduced here for convenience:
Clearly the velocity of the group, \( c \), is given as the difference in the concentration of the chemotactic agent between the left and right boundaries of the group, given respectively as \( \alpha \) and \( \beta \). Given this definition is easy to see that if \( u(\beta) \to u(\alpha) \), then by definition \( c \to 0 \). The condition under which this can happen can be seen with respect to the maximum in the concentration of the agent.

As previously suggested, for a symmetrically posed group in a stationary frame, the maximum coincides with the origin. As the velocity of the frame increases (due to chemo-attraction in this discussion), the maximum moves away from the origin contrary to the direction of motion, and takes on the characteristics of a travelling wave (Figure 2-13); the concentration drops away at the front of the wave and rises behind it. Now if we note that for an attracting group the position of the maximum will move toward, and indeed enter the group of attracting cells, then this fact in tandem with the wave like characteristics implies that \( u(\beta) \to u(\alpha) \) for some critical velocity \( c^* \) (see Figure 2-13).

Figure 2-13: The chemotactic agent is produced by all cells in the group with only a part reacting. A: In a stationary frame the chemotactic agent is symmetrically distributed about the producing group (grey shaded region), \( P \), with the chemotactic mechanism defined such that velocity of the group, \( c \), is given as a gradient, taken as the difference over the reacting group (dashed outline segment), \( R \). B: as the velocity of the frame increases the maximum concentration of the agent moves contrary to the group direction of motion, exhibiting a wave-like characteristic that causes the concentrations on either side of the reacting group to increase on one side and decrease on the other. Thus for sufficient \( c \) we can see that the concentrations will become equal and therefore we expect the velocity of the group to approach a limiting velocity \( c \to c^* \).

Accepting this explanation it should be clear that this will only occur when the chemotactic mechanism is attraction, moreover it occurs only for heterogeneous configurations were the agent is produced externally or internally when the whole group is producing the agent with a sub-population reacting. This latter point speaks to the equivalence between the chemotactic mechanisms for these configurations as previously described in 2.2.6 Motion Due To Chemotaxis.

However we can bring together some of the mechanisms we have studied so far into a generalized model. If we consider again a group that is producing the chemotactic agent and now introduce a border, \( x_B \), within the group, such that it defines what part of the group is reacting, then there are three cases that exist (Figure 2-14). If the border is within the group, for example at the midpoint \( x_B = 0 \), then this is simply the definition of heterogeneity, and we note that it produces equivalent analytical results for both internal and external production of the chemotactic agent. Secondly if the border coincides with the right-hand boundary of the group (\( x = \beta \)), then this is the definition of homogeneity and again we note that,
analytically, it is equivalent for both internal and external production. And finally there is the possibility we have not considered, that is, what happens when the border extends beyond the size of the group, say at $x_h = \beta + 1$.

![Figure 2-14](image)

**Figure 2-14**: Illustration of how the different mechanisms of heterogeneity and homogeneity can be shown by suitable prescription of a varying border defining the division of the group that is reacting. In all panels the red dotted rectangle represents the part of the group that is reacting and the grey rectangle the part that is producing. A: the chemotactically reacting part is smaller than the producing group (border is inside $P$). B: the whole group is producing the chemotactic agent and the whole of reacting ($P = R$). C: The chemotactically reacting part is larger than the producing part ($P > R$).

In this case it is neither heterogeneous nor homogeneous, however it can be shown to be fundamentally linked to the first case as an inversion of the chemotactic mechanism. Putting a slightly different way, we can see that by simply specifying a varying border we can make a fundamental link between the dynamical systems we have considered throughout this chapter. These systems are illustrated in Figure 2-15.

![Figure 2-15](image)

**Figure 2-15**: Illustration of the three fundamental bifurcations resulting from heterogeneity and homogeneity for a varying border. If we make plots in the parametric plane $c_0$ vs. $c$ there are three phase diagrams relating to the three border cases outlined in Figure 2-14. A: When the whole group is producing and sub part reacting we can see that there is always a travelling solution except in the trivial case when $c = 0$. However as $c_0$ is increased two new solutions appear one stable (solid black line) and one unstable (dashed black line) separated by a jump in graph, which is clear example of a fold bifurcation. B: when the border is coincident with the right-hand border we have a homogenous case which we have previously analysed and is clearly prescriptive of a pitchfork bifurcation. C: when the border extends beyond the right-hand boundary, we have a previously unseen case, however it can be seen as an inversion of A, reversing the direction of motion and therefore is also prescriptive of a fold bifurcation. Therefore we can see that all together these mechanisms can be represented one of these three dynamical systems as illustrated in A, B and C. Model parameters for all of the systems were chosen such that $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$ and $a = 50$. 
2.4 Chapter Summary

In the previous sections we derived a model describing a group of hypothetical biological cells, which move in response to the concentration of a locally produced chemotactic agent, and derived conditions using, where possible, a qualitative stability analysis that allow the group to move. In one case we assumed that the cells were homotypic (cells of the same type), which we described as a homogenous group. The implication of the group being homogenous, is that we assume that all cells in the group will respond equivalently to the chemotactic agent and therefore the force, and the resulting velocity \( c \), will be uniform over the group; in simpler terms the group will move coherently given an applied force. We further considered a secondary case where we assumed the group was now composed of two distinct populations of cells, we referred to as heterogeneous, and investigated the various permutations in production of the chemotactic agent, and the population that reacted to it.

To describe the chemotactic agent we employed a two chemical reaction-diffusion system of linear partial differential equations, with one acting as an activator to model the concentration of a hypothetical intracellular \( A \)-mRNA, and the other its reacting extracellular diffusing transcript \( A \). Thus the dynamics of the chemical system can be seen to be dependent on where \( A \)-mRNA is produced. Consequently we investigated two scenarios, one where \( A \)-mRNA is produced by a surrounding population of cells, and the other where it was produced by the cells in the group.

Given these two scenarios we separately defined and derived the corresponding chemotaxis functions, defined as the average in concentration over the group, taken as the difference in concentrations between its left and right boundaries. Defining the chemotaxis functions in this way lead us to a non-linear dynamical system prescribed by an implicit function. The analysis of this system in terms of conditions for motion, were investigated using a qualitative stability analysis and we showed that the systems that resulted followed canonical bifurcation of Pitchfork and Fold.

The results of this chapter are that under various configurations, group of cells can move uniformly under the influence of a chemotactic force, and we showed/derived conditions showing whether the group will move and whether such movement is stable, that is will continue to move indefinitely. Further we found that the production dynamics of the chemical system, whether the chemicals were produced internally or externally, does not display a qualitative or quantitative change in the results of the analysis of the chemotaxis function or its stability properties. More importantly we have showed a rich variety of chemotactic conditions that can give rise to self-regulating motion of a group of cells, both within a tissue and independently on a substrate.
Chapter 3

2D Modeling of a Migrating Group of Cells

3.1 Introduction

In the previous chapter we considered a 1D caricature model based on a group of cells abstracted as a segment of the real line, defining the production domain of a theoretical intracellular mRNA that gives rise to an externally diffusing protein, which we suggested acts to chemotactically attract or repel the group. The advantage of this model is that it afforded us a reduction in complexity, by neglecting biological factors such as intracellular adhesion and cell motility, in favour of analysis of chemical dynamics and the motile dynamical behaviour of the group described by the derivation and analysis of chemotaxis functions. In this chapter we seek to continue this analysis by modelling the group of cells as a two dimensional population of cells, implemented using a modified Cellular Potts Models (CPM). That is, rather than representing the group as an abstract segment, we shall now model it as a discrete, morphologically realistic population of cells. First however we shall consider extending the 1D continuous to a 2D numerical continuous model. In this way we can, in slightly simplified setting, consider the chemical dynamics to compare and contrast with the 1D continuous model before we demonstrate the qualitative results in the CPM.

3.2 2D Continuous Model of Homogenous Migrating Group

In the one-dimensional continuous model we observed chemical dynamics in an advection system, illustrative of Turing wave solutions (or travelling wave solutions) that do not change the shape of their profile in time. The characteristics of these travelling waves are given due to an advective term that is proportional to a chemotactic force, prescribed as the integration of the gradient of the chemotactic chemical over the activating region, or a group as we referred to it. In this section we will consider again this formulation but extend the problem to two dimensions, and consider the group to be represented as a circular domain or disk. In this sense the 2D model allows to take a step closer to the more physiologically realistic representation given by the Cellular Potts Model.
3.2.1 2D Model of a Migrating Group

As suggested above our principal motivation in this section is to analyse the velocity and chemical dynamics of the group with respect to a more relevant model structure, while still overlooking the dynamical nature of group morphology when represented as group of cells. As with the one-dimensional model we again assume that there is a stationary, activating region that is an abstract representation of a group of cells that gives rise to an externally diffusing chemotactic agent $u$. Let us define the activating region or group in the $xy$-plane as a two dimensional disk of radius $r = a/2$ or equivalently the diameter $d = a$, to correspond with the width of the line segment in the one-dimensional model repeated here for convenience:

$$\frac{\partial u}{\partial t} = D\Delta u + c(\Delta u; c_0) \frac{\partial u}{\partial x} - k_1 u + p(x); \quad k_1, k_2, D \in \mathbb{R}^+,$$

where we assume that group only moves on the $x$-axis. The two dimensional model of the chemotactic profile, $u$, is given as an extension of the one-dimensional model to two spatial dimensions, which we note is trivially a change to the diffusive term, given as the first term on the right hand side of the above equation. Where we note that a change of the Laplacian operator, $\Delta$, from one to two dimensions respectively is:

$$\Delta = \left( \frac{\partial^2}{\partial x^2} \right); \quad \Delta = \left( \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \right).$$

(3.1)

In addition the characteristic function, $p(x)$, describing the group is also extended to two dimensions to reflect the circular structure of the group, and thus the two-dimensional representation of the one-dimensional model can be given as:

$$\frac{\partial u}{\partial t} = D \left( \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \right) u + c(\nabla u; c_0) \frac{\partial u}{\partial x} - k_1 u + p(x, y); \quad k_1, k_2, D \in \mathbb{R}^+,$$

(3.2)

and the associated characteristic function:

$$p(x, y) = \begin{cases} k_2 & \sqrt{x^2 + y^2} \leq a \\ 0 & \sqrt{x^2 + y^2} > a. \end{cases}$$

(3.3)

Again we assume that the function is continuous in its first derivatives, $u \in C^1$, to ensure that the two solutions to (3.2), defined on the domains prescribed by (3.3), have equal concentration and curvature on the boundary of the disk. Figure 3-1 illustrates the chemical profiles of $u$ for both one and two-dimensional system for the case when $c = 0$. 


Figure 3-1: Illustrative comparison between profiles of the chemotactic agent in one and two-dimensional systems. A: In the one-dimensional case the group is given to be segment width $a$. B: In a two-dimensional model we show the circular group, with the activating region (black contour circle) centered on the $x,y$-axis giving rise to two-dimensional chemical profile (grey 3D surface).

Given the radially symmetric nature of the two-dimensional model it would seem a logical step to transform (3.2) to different coordinate system such as spherical or cylindrical coordinates. Thus as next step we considered converting to polar a coordinate system.

### 3.2.2 2D Polar Coordinate Representation of 1D Model

We can restate our two dimensional model in a polar coordinate system $x,y \rightarrow r, \theta$, where $r$ is the radial component centered about an arbitrary pole (or origin if you will), with $\theta$ the angular component of rotation. In such a system, it is usual to impose uniqueness given the rotational nature of the coordinates, and this is typically given to be $r > 0$, and $\theta \in [-\pi, \pi)$. The coordinate transformation relies on the trigonometric relationships between the two systems, given by the following:

$$x = r \cos(\theta); \quad y = r \sin(\theta) \quad (3.4)$$

and correspondingly the reverse substitution,

$$r = \sqrt{x^2 + y^2}; \quad \theta = \arctan(y/x). \quad (3.5)$$

The transformation occurs in two parts: firstly we will need to make the above variable substitutions to re-write the derivatives, then secondly repose the characteristic function $p(x,y) \rightarrow p(r, \theta)$. Re-writing the derivates can be a little involved so we shall forgo this lengthy process here (interested readers are directed to Appendix C) and instead we can go directly to it two-dimensional polar representation of the derivative terms:

$$\Delta = \left( \frac{\partial^2}{\partial r^2} + \frac{1}{r} \frac{\partial}{\partial r} + \frac{1}{r^2} \frac{\partial^2}{\partial \theta^2} \right). \quad (3.6)$$

which we can see is entirely dissimilar to that of its Cartesian counterpart, and it follows from the derivation of the above that the transformation of the advective term is:

$$\frac{\partial u}{\partial x} = \left( \cos(\theta) \frac{\partial u}{\partial r} - \sin(\theta) \frac{\partial u}{\partial \theta} \right), \quad (3.7)$$

and lastly the characteristic function:

$$p(r, \theta) = \begin{cases} \frac{k^2}{r} & \text{if } r \leq a \\ 0 & \text{if } r > a, \end{cases} \quad (3.8)$$
and thus we have the two-dimensional polar equivalent of our one-dimensional model

\[
\frac{\partial u}{\partial t} = D \left( \frac{\partial^2 u}{\partial r^2} + \frac{1}{r} \frac{\partial u}{\partial r} + \frac{1}{r^2} \frac{\partial^2 u}{\partial \theta^2} \right) + c \left( \frac{\cos(\theta)}{r} \frac{\partial u}{\partial r} - \frac{\sin(\theta)}{r} \frac{\partial u}{\partial \theta} \right) - k_1 u(r, \theta) + p(r, \theta). \tag{3.9}
\]

### 3.2.3 Solutions to the 2D Polar System

We are now faced with the problem that (3.9) may not admit any analytical solution. Solutions may be possible however when \( c = 0 \), that is for a stationary system. In this situation we note that (3.9) becomes a circularly symmetric problem, independent of \( \theta \) and \( t \), therefore we can re-write (3.9) as:

\[
D \left( \frac{\partial^2 u}{\partial r^2} + \frac{1}{r} \frac{\partial u}{\partial r} \right) - k_1 u(r) = p(r) \tag{3.10}
\]

The general solution to (3.10) is given as the superposition of the particular and complementary solutions, \( u_g = u_c + u_p \). The complementary, or homogenous solution, can be seen to be equivalent to a modified Bessel’s equation:

\[
x^2 \frac{\partial^2 y}{\partial x^2} + x \frac{\partial y}{\partial x} - (x^2 + \alpha^2) y = 0,
\tag{3.11}
\]

which we can see that if we divide throughout by \( x^2 \) and assume that \( \alpha = 0 \) becomes:

\[
\frac{\partial^2 y}{\partial x^2} + \frac{1}{x} \frac{\partial y}{\partial x} + y = 0,
\tag{3.12}
\]

which differs from (3.10) by only a factor \( k_1/D \) in the last term. However we can account for this by making the substitution \( x = \lambda x \) in (3.10) then:

\[
\frac{\partial^2 u}{\lambda^2 \partial r^2} + \frac{1}{\lambda^2 r} \frac{\partial u}{\partial r} - u(\lambda r) = 0,
\tag{3.13}
\]

and now multiplying through by \( \lambda^2 \):

\[
\frac{\partial^2 u}{\partial r^2} + \frac{1}{r} \frac{\partial u}{\partial r} - \lambda^2 u(\lambda r) = 0,
\tag{3.14}
\]

which is clearly equivalent to (3.11) for \( \alpha = 0 \), were we take \( \lambda^2 = k_1/D \). Given this, solutions are known to be in terms of two linearly independent modified Bessel functions of the first \((I_\alpha)\) and second kind \((K_\alpha)\)

\[
u_c(r) = A I_\alpha(r) + B K_\alpha(r)
\tag{3.15}
\]

dependent of the order, \( \alpha \), in this case assumed to be zero, and \( A \) and \( B \) are constant coefficients to be determined from boundary, initial and continuity conditions. The definitions of these modified Bessel function are given to be

\[
I_\alpha(x) = \sum_{m=0}^{\infty} \frac{x^{2m+\alpha}}{2^{2m+\alpha} m! \Gamma(m + \alpha + 1)}
\]

\[
K_\alpha(x) = \frac{\pi}{2} \frac{I_{-\alpha}(x) - I_\alpha(x)}{\sin(\alpha \pi)}
= \frac{\pi}{2} \frac{-(-1)^{\alpha+1} I_{-\alpha}(x) - e^{\alpha \pi i} I_\alpha(x)}{-i \sin(\alpha \pi)}
\tag{3.16}
\]
where \( J_\alpha(x) \) is a Bessel function of the first kind:

\[
J_\alpha(x) = \sum_{m=0}^{\infty} \frac{x^{2m+\alpha}(-1)^m}{2^{2m+\alpha}m!\Gamma(m + \alpha + 1)}.
\]  

(3.17)

Thus using these definitions we can show directly that the solution (3.15) can be written in the rather cumbersome form:

\[
u(r) = \begin{cases} 
\frac{k_2 l_1(ya)}{k_1 l_0(ya) K_1(ya) + l_1(ya) K_0(ya)} & r > \pm a/2 \\
\frac{k_2}{k_1 l_0(ya) K_1(ya) + l_1(ya) K_0(ya)} l_0(yr) & r \leq a/2,
\end{cases}
\]  

(3.18)

where the solution to the particular integral for (3.10) is trivially defined as \( u_p = k_2/k_1 \) and \( \gamma = \sqrt{k_1/D} \).

3.2.4 Comparison between 1D and 2D Profiles

Bringing together the solutions to the 2D polar Bessel function, 2D numerical and 1D analytical problems we can plot the steady states, that is non-travelling solutions without chemotaxis, to illustrate how the profile of \( u \) changes between these different implementations of the model.

In the first instance when we consider Figure 3-2A, we note that the profiles in all cases are illustrative of a diffusive profile, or in a more physical sense the dissipation of heat in a 1D rod with constant heat source at its centre. However there is a significant difference in the maximum value of \( u \) between the 1D and 2D models, where we see an approximate doubling of the maximum in the 1D model as compared to that of 2D.

This difference can be accounted for when we consider the difference in how each model represents the group. In the 1D model the group is represented as line segment and in 2D a disk. The implication of this seemingly innocuous difference brings about a fundamental change in how chemicals spatially diffuse in either system.

![Figure 3-2: Stationary steady-state solutions of the 1D and 2D models illustrating diffusive profiles. In both cases the model parameters where equivalent such that: \( \alpha = 50, k_1 = 0.00025, k_2 = 2k_1, D = 0.5 \). 1D (red profile) is a solution to (2.11) for a steady state (c = 0); 2D Bessel solution is a solution to (3.18) and 2D numeric is a solution given by the description in section 3.2.5.](image)

To understand this difference first let us consider the process of diffusion in the 1D model. If we restrict our view to only consider diffusion with respect to the boundary of the group, that is, the diffusion
in the concentration of \( u(x = a/2) \) (or equivalently \( x = -a/2 \)), then we can see that \( u \) will reduce for \( x \leq a/2 \) and will increase for \( x > a/2 \) until the equilibrium state is reached. In other words, and referring to the previous heat analogy, there is a net thermal flux on the boundary of the heat source as the temperature in the rod tends to thermal equilibrium. Considering this as a spatially discrete process, then it is clear that the flux flows between discrete segments of equal volume/area. However when we consider the 2D model, this is not the case.

**Figure 3-3:** Diffusive Flux is affected by curvature that occurs when we introduce a circular domain that is producing the chemotactic agent. A: In the 1D model the segment is arranged as a rectilinear segment such that when we consider the diffusive flux between segments, it passes between areas of equal are where it is assumed that \( \Delta x = \Delta y \). B: when we introduce curvature due to the circular nature imposed by both the 2D numerical model and the CPM, we can see directly that the diffusive flux passes between areas of unequal size essentially causing a dilution effect that increases the gradient between the areas which can readily be seen by the changes in arc length \( \Delta \theta_1 < \Delta \theta_2 \).

Again if we restrict our view to diffusion on the boundary, a disk in the 2D model, then flux will proceed in an equivalent fashion to that of 1D. However in 2D there is a diluting effect, brought about by the fact that the flux does not flow between discrete segments of equal volume/area (Figure 3-1). That is to say, the flux at the boundary of the disk flow from \( u(r = a) \) to \( u(r = a + \Delta) \), were \( \Delta \) is a unit spatial increase in the radius of the disk, from an area of size \( 2\pi a \) to an area of \( 2\pi a + 2\pi \Delta \) causing a spatial diluting effect not present in the 1D model.

**Figure 3-4:** Analytical results of continually increasing the groups in 1D and 2D models so as to reduce the diluting effect of curvature in the 2D model. The effect of curvature on the boundary of the 2D group causes a spatial diluting effect as the flux from smaller to a larger volume over the boundary proportional to the curvature. The curvature effect can be reduced by increasing the size of the groups and thereby reducing local curvature. Therefore we see that there is a value of \( a \) for which both models have equal concentrations on the boundary. A: The size at which the two models saturate are grossly different, and in terms of the 2D model these sizes become computationally intractable in numerical in simulations and therefore we call upon the Bessel solution derived above. B: As with the 2D case the 1D also saturates to a value at which both concentrations become equal implying the boundary curvature effect due to diffusion is negligible. Model parameters where equivalent to those used in Figure 3-3 such that: \( a = 50, k_1 = 0.00025, k_2 = 2k_1, D = 0.5 \).
From a slightly different point of view, this can be seen as a problem of curvature on the boundary with respect to diffusion, and therefore we can reduce the diluting effect by reducing the curvature. Since the curvature at a point on the boundary is given to be the reciprocal of the radius, that is \( \kappa = \frac{1}{r} \). Then if we continually increase the size of the disk then \( \kappa \) reduces and thus \( \lim_{r \to \infty} \kappa = 0 \), or one would expect the curvature to disappear completely. To illustrate this we can consider a single value of \( u \) on the boundary in both the 1D and 2D models, and continually increase the size of both groups equally, that is the width of the line segment \( a \) in 1D and the radius of the disk \( r = a/2 \) in 2D, then we should find that these values will become equal at some value of \( a \) as the curvature effect becomes negligible. Indeed this is the case and is illustrated in Figure 3-4.

### 3.2.5 Chemotactic Motion of a Circular Group: Numerical Implementation

In this section we shall consider the implementation of the two-dimensional model with reference to chemotaxis as it translates from one dimension to two dimensions. The numerical methods we will apply are based on an explicit Euler scheme, where we implement a forward time centered space scheme (FTCS). The implementation of chemotaxis follows a direct extension from the one-dimensional model, where we will integrate the gradient of the chemotactic chemical, \( u \), over the circumference of the group. That is, we are taking the difference in the chemical concentration of \( u \) bilaterally about the vertical axis (Figure 3-5A) of disk of constant radius \( r_0 \).

\[
c = \frac{c_0}{\pi r^2} \int_{-\pi/2}^{\pi/2} \Delta u(r, \theta) r \cos(\theta) d\theta ; \quad \Delta u = u(r_0, \theta_0) - u(r_0, \theta_0 - \pi) \tag{3.19}
\]

where \( \Delta u \) is the difference in \( u \) for all values of \(-\pi/2 \leq \theta \leq \pi/2\). In addition there is now a dependence on the area of the group, \( \pi r_0^2 \), which in the one-dimensional model was taken to be the length of a line segment, \( a \). As before \( c_0 \) is a constant of proportionality that can be taken as the effective sensitivity of the group to the gradient of \( u \). In all subsequent simulations it will always be the case that \( 2r_0 = a \) when correlating between the models.

![Figure 3-5: Illustration of the implementation of chemotaxis in the two-dimensional model: A: The mechanism of chemotaxis, as in the one-dimensional model, is prescribed as a bilateral difference about the spatial axis transverse to the group's direction of motion, that for a given angle \( \theta_0 \) takes the form \( \Delta u(r_0, \theta_0) \). We assume that the diameter of the group is of fixed size \( d = 2r_0 \) and that the sum of all differences for \(-\pi/2 \leq \theta \leq \pi/2\) represents the total chemotactic force acting on the boundary of the group. B: The illustration of the numerical implementation demonstrates how we discretise the model onto a grid and how we take the difference in concentration, \( \Delta u \), of the chemotactic agent between the gridpoint \((x, y)\) and its vertical reflection at \((x - 2w, y)\).](image-url)
For the numerical implementation of (3.19) we discretize the group/disk onto a grid which approximates the disks with increasing accuracy as we increase the grids resolution (Figure 3-5B). Recalling that the chemotactic force acting on the disk is given by the difference in concentration of the chemotactic agent between the right- and left-hand boundaries, and further that we assume that the direction of motion is confined to the horizontal axis, then the concentration profile of the agent is symmetric with respect to this axis. Therefore the interpretation of (3.19) in a numerical setting is an iteration over the right-hand plane of the grid, and determining the concentrations at points on that half-plane/half-grid that satisfy \( x^2 + y^2 = r \), where \( r \) is given to be the radius of the disk. Since we assumed axial symmetry we can calculate the mirror difference in the opposite half-grid by subtracting the width of the grid, \( 2w \), from the \( x \) coordinate (Figure 3-5B) and thus the difference in concentration on the boundary about the vertical axis is at grid-points \((x^2 + y^2)\) and \(((x - 2w)^2 + y^2)\), for \( 0 \leq x \leq w \) and \(-w \leq y \leq w\), such that \( x^2 + y^2 = r \).

3.2.6 Travelling Solution of a 2D Migrating Group.

In this section we investigate the travelling solutions that arise for a circular group, given the definition of chemotaxis in the previous section, and attempt to correlate these findings with those found in one-dimension. In one-dimension we showed that when we fix all model parameters except \( c_0 \) and make plots in the parametric plane \( c \) vs \( c_0 \), we find that the dynamical behaviour is consistent with a sub-critical pitchfork bifurcation. At small values of the parameter \( c_0 \), less than a critical bifurcation value \( c_0^* \), we find that no travelling solutions exist. However for values \( c_0 > c_0^* \) travelling solutions emerge in pairs, with equal magnitude and opposite sign, with the physical interpretation that the solutions are travelling in either a positive or negative direction on the \( x \)-axis. When we applied a similar analysis to the 2D model we found a qualitatively equivalent behaviour, with the notable exception that there is a discrepancy in the value of \( c_0^* \) between the models, specifically in 1D \( c_0^* \approx 1.8 \) and in 2D \( c_0^* \approx 4.8 \) a difference of \( \approx 63\% \) (Figure 3-6A).

The apparent difference in \( c_0^* \) between the one and two dimensional models can be explained as a result of the curvature in the profiles of \( u \), which was discussed in the previous section. However from a slightly different point of view, we know that the curvature due to diffusion is a function of the radius, \( r \), of the group (or width of the line segment in 1D). An increase in \( r \) necessarily implies an increase in the production/activation domain leading to an increase in the production of the chemotactic agent \( u \). It follows then that the velocity of the group, \( c \), will also increase proportional to this change. Naively one might expect that such a process might continue without limit that is, as \( r \rightarrow \infty \Rightarrow c \rightarrow \infty \). However this is not the case, since if we do not scale diffusion \( (D) \) and the chemotactic response \( (c_0) \), then as \( r \rightarrow \infty \Rightarrow c \rightarrow 0 \).
Figure 3-6: Comparison between 1D analytical and 2D numerical velocity ($c$) characteristics for identical model parameters. A: In the parametric plane $c$ vs $c_\theta$ the behaviour in 1D is descriptive of a sub-critical pitchfork bifurcation were non-zero velocities only occurred above a critical value, $c_\theta^*$, of the bifurcation parameter $c_\theta$. Following a similar analysis in the 2D numerical model we can show a qualitatively equivalent result, with the exception that the value of $c_\theta^*$ is slightly higher: $c_\theta^* \approx 1.8$ in 1D and $c_\theta^* \approx 4.8$ in 2D. The apparent difference is due to local curvature introduced when representing the group as disk, that effectively increases the diffusion effect by what one might call dilution due to the chemical flux between unequal volumes/area. B: We can negate this effect by increasing the sizes of the 1D and 2D groups, which will ultimately have the effect of decreasing local curvature and therefore we should expect both models to saturate to equal velocities for an ideal size of the groups ($a$ in 1D or $2r$ in 2D). Model parameters were such that: $a = 50$ in $\Lambda$ and $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$ for both $\Lambda$ & $B$.

In physical terms this is quite intuitive: The chemotactic force on the group of a given size produces a motile response $c$. If we increase the size of group, for example twice, then we should expect to increase the force proportionally to scale the system. This is nothing more than a consequence of Newton’s second law $F = ma$, from which we can infer that as the size of the group increases, the influence of the advective term reduces. Similarly as diffusion is a constant and independent of the size of the group, it will also become negligible as the size of the group gets very large and so from these two arguments we find:

$$\frac{\partial u}{\partial t} \approx -k_1 u(r, \theta) + p(r, \theta) \Rightarrow u = \frac{p(r, \theta)}{k_1}. \quad (3.20)$$

and therefore, given the definition of chemotaxis we should expect $c \rightarrow 0$ as $r \rightarrow \infty$ (Figure 3-6B).

3.3 Preliminaries Of CPM Models Of Group Migration:

The purpose of this is to give a basic set of terminology for the proceeding material, in that we shall use simple mnemonics to refer to the different CPM configurations. Thus to aid subsequent discussions, let us first consider the visual construction of the models we will be using. This will make discussion simpler at least in terms of naming conventions. Considering Figure 3-7 we have two initial CPM lattices: one illustrating a heterogeneous model of three distinct populations of cells (Figure 3-7A) and the other a homogeneous model of two (Figure 3-7B). The red cells are taken to be a surrounding cellular medium representing a tissue in which the group will travel. In the heterogeneous case the group itself is composed of a four by four square of cells, divided equally into eight green and eight blue cells, while the homogeneous, a single four by four square of green cells. The square nature of the cells are a consequence of the underlying structure of the CPM lattice as a regular square tessellation, which can be taken as an initial condition.
On all occasions when we use the term lattice, it will unambiguously imply all cells of all types constituting a CPM simulation from which there are two illustrated in Figure 3-7. Once again, and in-line with its previous use, the group will represent all cells that are distinct from the surrounding medium of red cells. That is the group may be heterogeneous or homogeneous, but we refer to it is a group nevertheless. A further distinction may be made when we consider chemicals. Since any population can potentially be responsible for production of the chemotactic agent, we may refer to such a population as an activator, since it is the internal concentration of a non-diffusing chemical that gives rise, or activates, the externally diffusing chemical we have referred to as a chemotactic agent, or simply $u$.

Since fundamentally we are dealing with permutations of a population of cells reacting to the agent ($u$), and a population producing it, we shall refer to a particular mechanism with the shorthand form: [Producer][Chemotactic Mechanism][Reactor]. For example in a simulation where the green cells are producing a chemical that attracts the red cells, we shall simply write $GaR$, or when the blue cells are producing a chemical that repels the green cells we shall write $BrG$, and so on, where clearly $a$ and $r$ are the chemotactic mechanisms of attraction and repulsion respectively.

The parameter space of the CPM has within a set that is applicable to all simulations that follow. These are the Boltzmann temperature of the system $T_b = 6.0$ that defines the rate of evolutions of the system and the elasticity or compressibility of cells $\lambda = 0.6$. The ratio of these parameters, $\lambda/T_b$, defines the fluctuation rate of cell boundaries (or cell membranes), where for large $\lambda$ or small $T_b$ the system will tend to freeze and conversely for small $\lambda$ and large $T_b$ the system becomes chaotic and the cellular structure would become unrecognizable. Thus the specification of these parameters is chosen on a qualitative basis due to characteristic morphology one might observe actual biological cells, but also optimally for migratory behavior in the group.

Finally there are parameters associated with differential adhesion, however these parameters are not necessarily fixed for simulations, and this is especially true when we consider the heterogeneous model. We can make an important observation however, in that there are fundamental relationships
between adhesive parameters that define important bonding behavior. We know from the introduction that the CPM uses a symmetric triangular matrix to define the adhesive relationships between cells types, and that the entries are given by the shorthand notation, $J_{m,n}$.

The subscripts identify the indices of matrix entries identifying a particular cell type when $m = n$, or the relationship between two differing types when $m \neq n$, where $J_{m,n} = J_{n,m}$. With this definition if we want two cell types to form a connected tissue we require that $J_{m,n} > 2J_{n,n}$, and for two cell types to remain connected but to not dissociate into each other, we require at least $J_{m,n} > 2J_{n,n}$ and $J_{m,n} > 2J_{m,m}$. However we must note that the rules are predicated on the absence of any other external forces (for example chemotaxis) which we should consider as guidelines.

### 3.4 CPM Homogenous Model of a Migrating Group

As in the 1D case, here we assume the group of cells is composed of a population that are of the same type (homotypic). Being of the same type necessarily implies that individuals in the group will react equivalently to any environmental stimuli, which in this case is the gradient of a locally produced chemotactic agent $u$. Further we again assume that this agent can be produced internally or externally by/to the group. That is either cells in the group or a surrounding population, contains $A$-mRNA that promotes the production of the externally diffusing agent. In the former case referred to as an auto-chemotaxis.

#### 3.4.1 Concentration Profiles for an Internally Produced Chemotactic Agent

The simulation of chemical in the CPM is all but equivalent to that of the one-dimensional models, save two notable differences: Firstly simulations are conducted in a stationary frame of reference, implying we do not need an advection term, and secondly the diffusion term is extended to two dimensions. As in the one-dimensional model we assume that cells give rise to a diffusing chemical $u$, and do so proportional the concentration of an internal non-diffusing chemical denoted by the characteristic function $p(x)$, we previously defined respectively as $A$ and $A$ mRNA. Thus we can rewrite (1.2) as

$$
\frac{\partial u}{\partial t} = D \left( \frac{d^2 u}{dx^2} + \frac{d^2 u}{dy^2} \right) - k_1 u + p(x), \quad k_1, k_2, D \in \mathbb{R}^+. \tag{3.21}
$$

where chemical $u$ decays proporproportional to its own concentration with rate $k_1$ and is produced with rate $k_2$ and $D$ is the constant coefficient of diffusion.

The prescription of $u$ is trivially extended to two dimensions with little modification. However the concentration of $v$ is not so trivial. In the one-dimensional model $p(x)$ was confined to a static line segment, however in the CPM the group is composed of moving cells, implying that the production domain described by $p(x)$ is non-static and is updated on a continual basis as the producing cells move.

Therefore we cannot give a precise mathematical description of $p(x)$, it is given as a function of the CPMs underlying stochastic process. However in general we could describe it in the following way,
were $\sigma_{i,j}(\tau)$ is the lattice site at point $i,j$ on $\mathcal{L}$ with cell type $\tau$, and $\tau_p$ is the type of cell that is producing $p(x)$, as in the one-dimensional model, is held at a constant concentration of $k_2$. Using these equations we ran simulations on the CPM to illustrate the concentration profiles of $u$ and $p(x)$ for a stationary and travelling group of cells (Figure 3-8). Note here that when we say stationary we are referring to a group which is not moving rather than its frame of reference, and equivalently for the travelling group.

\[ p(x) = \begin{cases} k_2, & \forall \sigma_{i,j}(\tau) = \tau_p \\ 0, & \text{otherwise} \end{cases} \] (3.22)

Figure 3-8: Concentration profiles of $u$ both a stationary and travelling group of 25 cells. For a stationary group we can see that production (solid black) is held constant with the cell boundaries (red outlines) and gives rise to diffusing chemical $u$ (B) producing an approximately circularly symmetric distribution (grey shade). For a travelling group (C) the diffusing chemical $u$ (D) now shows a distinct trailing profile that lags behind the group. Compare and contrast these with Figure 2.1. Model where chosen such that $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$, volume constraint $\lambda = 0.6$, temperature $T = 5.0$, and differential adhesion $J_{r,r} = 3.0$ and $J_{w,w} = 9.0$ and $J_{w,r} = J_{r,w} = 7.0$ where the subscript $r$ denotes red cells and $w$ the surrounding substrate.

Considering Figure 3-8 we can see that we have an analogous result to the one-dimensional equivalent, where again we see an approximately symmetric distribution of $u$ for a stationary group. Further for a travelling group $u$’s concentration displays a distinct asymmetry in the form of a travelling wave, with the maximum in concentration displaced contrary to the group’s direction of motion, equivalent to the one-dimensional case. We also note that the group of cells also shares this asymmetry, that is, the group is deformed as individual cells in the group react differently to the varying concentrations of $u$ they are exposed to. Therefore it should be clear that as the chemotactic force on the cells increases, we should expect this type of deformation to increase. However we shall consider this in a subsequent section. In summary the concentration profiles observed in the one-dimensional model are mirrored in the two-dimensional case, with spatial variations brought about by individual cell motility in the group.

3.4.2 Motion Due To Chemotaxis On The CPM.

In the one-dimensional model chemotaxis was implemented as the difference in concentrations over the reacting population of cells, be they the cells comprising the group or an external population. Clearly we are afforded this simplification because we do not consider the populations to have any physiological variances; we do not consider characteristics such as cell density or changes in morphology (shape), for example. However the CPM is a discrete model with an evolving cellular morphology, which implies that the domain of integration for calculating the chemotactic affect over the reacting population is constantly changing, and more importantly it does so stochastically. In addition the discreteness comes
from the fact that the CPM only considers energetic interactions that take place on the boundaries between cells. Putting it a slightly different way, cells are delineated on the CPM surface/lattice as boundary representations, where the spaces contained within boundaries are only considered volumetrically. Therefore to influence the outcome of a CPM simulation, we are required to modify this interaction energy on these boundaries.

Energy in this context can be seen from a mechanical point of view as the concept of work, \( W \) given as the product of a force, \( \mathbf{F} \), and a spatial displacement \( \mathbf{d} \), such that \( W = \mathbf{F} \cdot \mathbf{d} \). In this sense \( W \) is the sum of all the forces acting on a cell boundary, equalling the kinetic energy associated with the cell, with \( \mathbf{F} \cdot \mathbf{d} \) the spatial displacements on cell boundaries due to the underlying stochastic process of the CPM. Thus we can see that the kinetic energy associated with cell boundary fluctuations can be manipulated to produce a motile response, by specification of the vector quantity \( \mathbf{F} \), such that spatial displacements on cell boundaries occurs preferentially in some direction, say \( \mathbf{d} = \mathbf{x} \). Putting this in the context of the CPM, changes in the work done by the system, \( \Delta W \), follows a minimization scheme such that a spatial displacement, \( \mathbf{x} \), of a unit length of a cell’s boundary must act to reduce the energy of the system, otherwise it is rejected. In its basic setting this change in energy is,

\[
\Delta E = E_{\text{adhesion}} + E_{\text{pressure}},
\]

given as the sum of the forces associated with intracellular adhesion, \( E_{\text{adhesion}} \), and the incompressibility of the cells, \( E_{\text{pressure}} \) implemented as a volumetric constraint. Therefore we can influence this change by the addition of a chemotactic force,

\[
\Delta E = E_{\text{adhesion}} + E_{\text{pressure}} + E_{\text{chemo}},
\]

that is proportional to the gradient of \( \mathbf{u} \). That is the work done by the chemotactic force can be defined as \( E_{\text{chemo}} = \mathbf{F} \cdot \mathbf{x} = -\beta \nabla \mathbf{u} \cdot \mathbf{x} \), where \( \beta \) is a constant coefficient that for \( \beta > 0 \) implies attraction and for \( \beta > 0 \) repulsion. Thus boundary displacements acting in the direction of \( \nabla \mathbf{u} \) will maximise energy minimisation, which on average will compel cells to move towards or away from the areas of highest concentration of \( \mathbf{u} \) depending on the specification of \( \beta \).

### 3.4.3 Group Motion for an Internally Produced Chemotactic Agent

The first case we consider is when a compact group of cells is producing the externally diffusing chemotactic agent \( \mathbf{u} \). Recalling that the chemotactic force can either repel or attract cells, and further this force can act on the cells in the group or a surrounding population, we have, at least theoretically, four different scenarios to consider: two for attraction and repulsion of and external population of cells, and two for self-attraction and self-repulsion of the group. However, and informed by the one-dimensional analysis, we know that self-attraction is not feasible since we have shown in 1D that no travelling solutions exist for \( c_0 > 0 \). In addition we have an reacting, external population we have found no results in one-dimension, since we have assumed that chemotactic force is given as an average of the gradient over the group.
However we don’t any such problems in the CPM since the force acts over the boundaries of the cells and hence there is no equivalent averaging taking place, implying the chemotactic force is independent of the size of the group. Therefore the CPM admits new solutions in terms of external population not present in the 1D model. Thus we are left with self-repulsion of the group and attraction or repulsion of the external population. We will consider each of these mechanisms in the following sections.

**Motion For an Isolated Self-Repelling Group (GrG).**

In the simplest setting the one-dimensional analysis demonstrated that a group of cells can self-repel if the value chemotactic forcing parameter (or bifurcation parameter) $c_0$ was of sufficient magnitude. Above this value two travelling solutions would emerge that are equal in magnitude and opposite in sign, describing the a positive or negative direction of motion on the horizontal spatial axis, $x$. In the CPM we are faced with two possibilities, one when an isolated group is self-repelling on a substrate, and the second when there is a surrounding or external population; we shall consider the external case in the subsequent section.

In the first case we constructed a CPM simulation with parameters equivalent to those in the 1D model, and showed that such a group can give results that clearly demonstrate the mechanism of self-repulsion (Figure 3-9). However to achieve this result we had to change the total kinetics rates, $k_1$ and $k_2$ by a factor of 0.1 and increase the chemotactic coefficient $c_0$ by several orders of magnitude.

![Figure 3-9: Simulation of a self-repelling group. A group of nine cells initially arranged as a 3 by 3 square (A) at simulation step zero that diffuses a chemotactic agent into the surrounding environment is given an initial condition such that the chemical it produces is shifted fifteen lattice points left of it true position (B) shown here at step 300. At step 1000 the condition is removed and then the cells become chemotactically sensitive to the chemotactic agent, shown here at 4500 steps after the condition has been removed (C). Subsequently the simulation is allowed to run for a further 4500 steps where we note that the cells have continued to be repelled as predicted in the one-dimensional model (D). To produce this simulation the kinetics coefficients of production and decay were given values equivalent to the one-dimensional model, $k_1 = 0.00025$ and $k_2 = 2k_1$. However to enable a similar outcome the overall kinetics rate was lowered by a factor of 0.7 (or 30%). The coefficient of chemotaxis was also adjusted so that it was four orders of magnitude greater than that in 1D, $c_0 = 1000$.](attachment:image.png)

The explanation of this apparent change is once again due to changes in local curvature on the boundaries of cells in the group. Recalling that chemotaxis is implemented as an energy quantity, that for random boundary fluctuations due to the stochastic process of the CPM, causes such displacement to occur preferentially with the gradient of $u$, be that attraction or repulsion. This clearly implies that the morphology of the group is directly affected by $u$’s gradient, which is of course a rather obvious
statement. However, when we factor in the effect of curvature, we can see that for areas of high or low local curvature the diffusive flux increases or decreases respectively, which in turn so does the local gradient of $u$. Thus in areas of high curvature the local chemotactic effect is amplified, and the net result is deformation of the group, which in extreme can cause total dissociation of the cells.

In this sense the reduction of the kinetics rates serves two purposes: firstly reducing the overall rate of kinetics is equivalent to increasing diffusion, which in turn flattens the gradient of $u$, and therefore reduces the effect of chemotactic amplification associated high curvature. And secondly all local curvature effects become near equal to the total boundary average, implying that deformation is reduced and the group will maintain an approximately circular morphology. Given this explanation it would seem that an increase in $c_0$ is inevitable since if we are flattening the overall gradient of $u$ by reducing kinetics, it merely is logical to assume that we will need to increase the chemotactic response of the cells in the group, by increasing their sensitivity to this change in the gradient.

**Motion for a Group In a Tissue.**

In this section we introduce a homogeneous population of cells, which one can refer to as a tissue, that surrounds our group of cells. We consider that this tissue is homotypic, in that the cells constituting it are all of the same type. However to differentiate them from the group they are distinct in one important way, differential adhesion. Differential adhesion is a fundamental quantity in the CPM that allows us to differentiate between cell types (other than colour that is). Its purpose, as indicated in the introducing chapter, is to allow cells of equivalent types to preferentially adhere to one another. This is clearly important for the cells in our group, since in the absence of any other differentiating factor, there would be no reason for the cells in the group to maintain contact with one another, and therefore they would quickly dissociate when exposed to the chemotactic force. However this not merely a phenomenological aspect of the CPM, in a biological setting this is an analogue of trans-membrane cadherin (calcium-dependent adhesion) bonds that are responsible for essentially gluing biological cells together, consider skin cells or tissue constituting the stomach wall for example.

The addition of an external tissue thus brings with at least two extra forces that will come to bear when consider motile dynamics of the group: viscous forces due to incompressible nature of the tissue, and adhesion forces due to differential adhesion properties between the surrounding tissue and the group. Given this, it is clear that we should expect there to be a significant difference between the motion observed in the previous section this one, at the very least in terms of velocity.

Having a surrounding population of cells also brings with it new possibilities for group motion that are not theoretically possible in the 1D model, which is due to how we derived the chemotactic mechanism as an averaged gradient over the whole group. If we consider active group motion, that is, the chemotactic force is acting on the group and not the surrounding population, we defined this as:

$$F_{active} = \frac{1}{a} \int_{-a/2}^{a/2} \nabla u(x) dx = \frac{u(a/2) - u(-a/2)}{a} = \frac{1}{a} \Delta u.$$  \hspace{1cm} (3.25)
When we consider passive motion, that is, the external population is reacting to the agent, then it would follow that:

\[
F_{\text{passive}} = \frac{1}{-\frac{a}{2} + \infty} \int_{-\infty}^{\infty} \nabla u(x) \, dx + \frac{1}{\infty - \frac{a}{2}} \int_{\infty - \frac{a}{2}}^{\infty} \nabla u(x) \, dx
\]

\[
= u\left( -\frac{a}{2} \right) - u\left( \frac{a}{2} \right)
\]

\[
= -\frac{1}{\infty} \Delta u = 0,
\]

and we can see that the integration leads to zero net force, assuming we could evaluate \(2.2\). When we consider the CPM implementation as described in Section 3.4.2, the integration of the force is now a discrete summation over the cell boundaries, and therefore we have no such dependence on the area of the cells, and therefore the chemotactic force for active and passive movement is given as:

\[
F_{\text{active}} = -F_{\text{passive}} = \Delta u, \tag{3.27}
\]

that states that the passive and active mechanisms rely only on the difference on the concentrations on the boundaries of the group, and that a change from active to passive motion results in the direction of motion being reversed. More specifically, and for a potentially travelling group, if we change the population that is reacting, then to maintain to the same direction of motion we must change chemotactic mechanism from attraction to repulsion or vice versa.

From this it is clear that are at least two scenarios to consider, and using the naming conventions suggested, these are \(GaR\) and \(GrG\). However there is clearly a third scenario which we have not mentioned and this is \(GrR\). From the arguments given above we know that travelling solutions exist when \(GrG\), and indeed this was shown in the analysis of the 1D model. And clearly \(GaR\) must also if we are to believe \(3.27\). However it is not necessarily clear if \(GrR\) will also admit travelling solutions. To demonstrate that no solutions exist when \(GrR\), consider the following scenario, illustrated in Figure 3-10:

Consider the simple case of a stationary group, in a tissue, that is diffusing a chemotactic agent into the surrounding environment. In this setting, regardless of who is reacting, it should be clear the net result of the chemotactic force acting on the tissue or the group is clearly zero. In other words the group is positioned symmetrically within the agent, and in the absence of symmetry breaking event, such as a spatial perturbation, the chemotactic force acts equally in all directions (Figure 3-10:B).

Assume now that we impose some initial condition, such that we force the group to move through the tissue, for example mechanically drag it, and subsequently let it go. At the instant of release, and depending on whether we dragged it to the left (Figure 3-10:A) or to the right (Figure 3-10:C), the concentration profile of the agent would exhibit wave-like characteristic. We now ask the question will the group continue to move? In more formal terms, will the perturbation from the stationary equilibrium lead to an instability in the form a travelling solution.
In simple terms the answer will be yes if the resulting chemotactic force acts in the direction of motion imposed by the initial condition, in this case either positively or negatively on the $x$-axis; will it become unstable? In the active case, that is $GrG$, we know from all previous definitions that chemorepulsion is given (in the simplest sense) as the negative of the difference in concentration of the boundary of the group, that is $GrG = -\Delta u$. From this it is clear to see that this results in a travelling solution. Further it follows from (3.27) that $GaR$ will also admit a travelling solution. However if we consider $GrR$, recalling this is a passive chemo-repulsion mechanism, then it equivalent to $-F_{\text{active}} = \Delta u$, and it is clear that the group will return to its stationary state, and thus the mechanism of $GrR$ is not feasible.

To corroborate these findings we ran two CPM simulations, one $GrG$ and the other $GaR$. In both simulations the parameters of the model were identical, and further they were identical to those used in the 1D model except for a relative decrease in the overall kinetics as explained in section 0. Specifically we set $k_1 = 0.00025$, $k_2 = 2k_1$, $D = 0.5$ and $a = 50$. The size of the surrounding population was set at 55 cells wide by 31 cells high, and the group as square of 3 by 3 cells. As indicated in the previous scenario, in the absence of any symmetry breaking event, the cells will not move appreciably from their starting position, and so we provide this by offsetting the group spatially from the chemotactic agent it produces, more simply a shift in the profile of $u$ relative to the group. This represents an initial condition of sorts that introduces asymmetry, and thereby instigates motion in the group (see Figure 3-9B). We allow the initial condition to run for 1200 simulation steps, at which point we remove “shift” and the simulation is allowed to continue naturally, that is in the absence of any artificial conditions. The results for $GrG$ and $GaR$ are illustrated in Figure 3-11.
Figure 3-11: Passive (GaR) and active (GrG) motion for a homogenous group in a tissue. The group, represented by green cells, is placed in a cell medium (or tissue) represented by red cells, give rise to an externally diffusing, chemotactic agent which acts to either repel the cells in the group, GrG (B), or attract the cell of the surrounding tissue, GaR (A). In both mechanisms the cells are given an initial condition at $T = 0$ (as given in Figure 3-9B) and motion forced to travel form left to right (as in Figure 3-9B). All model parameters are identical with the exception of chemotactic mechanism changing from attraction to repulsion and the cell type that reacts from red to green. In both cases it can be seen that indeed that both mechanisms result in uniform direct motion, indicated by both groups reaching the same spatial end point. However there is a clear discrepancy in the time it took each mechanism to reach this end point. For GaR it took ~2900 steps and for GrG it took ~8000 steps, this is a time difference of ~70%. In a purely mathematical sense, these two mechanisms should be identical as shown in (3.27), but this is simply not the case. We shall delay consideration of this problem until later, in this context of this result, it is only important to note that both mechanisms achieve the same result, and more importantly using an identical parameter set.

Model parameters were chosen such that $k_1 = 0.000025$, $k_2 = 2k_1$, $D = 0.5$, volume constraint $\lambda = 0.6$, temperature $T = 5.0$, and differential adhesion $J_{r,r} = 3.0$, $J_{g,g} = 2.0$, $J_{r,r} = J_{r,g} = 7.0$ and $J_{w,w} = 0.0$ and $J_{w,g} = J_{w,r} = 9.0$ where the subscript $r$ denotes red cells and $w$ the surrounding substrate.

From Figure 3-11 it is clear that GaR and GrG both produce the desired result of a directed, uniformly travelling group of cells. However there is a clear discrepancy between the chemotactic mechanisms in the amount of time group it takes the group to reach the spatial end point, here arbitrarily set at $x = 300$ grid units on the lattice. Accounting for this discrepancy is clearly important as (3.27) demonstrates that there should be no such difference, moreover the mechanisms should be equivalent for an identical set of parameters, but this clearly is not the case.

Our initial investigation centered on the boundary representation of the cells on the lattice, and the effect of integrating the forces over such boundaries. However the problem arises, and indeed a solution becomes more elusive, when one considers the group and surrounding tissue to be an incompressible medium, such as fluid. And of course in this sense, we negate any effect(s) that compressibility might have on the cellular structure. Of course what we alluding to, is that the time discrepancy we see between GaR and GrG, is entirely due to the compressible nature of the CPM.

In practice incompressibility in the CPM is function of the volume of each cell, which is imposed as a quadratically increasing energy penalty as the volume of each cell deviates from its pre-assigned target volume. To understand how this works in practice, one needs to consider that the CPM evolves simulations as attempts of a cell’s boundary to expand or contract due to randomly imposed fluctuations that will succeed or fail depending on how the system energy is affected. In short the fluctuations manifest as changes in the volumes of the cells as unit lengths of each cells boundary is gained or lost according simple energy based rules. Thus it is entirely possible for cells to maintain a marginal increase or decrease in their volume over time, if that increase or decrease is energetically beneficial to the energy state of the system; energy minimization.
Figure 3-12: The chemotactic force on the group can affect its velocity due to compression or expansion of cell volumes that increase or decrease the size of the group. A: When the mechanism is GrG there is a net increase in the size of the group which in turn enlarges the production domain. C: when the mechanism is GaR there is a net reduction in the size of the group leading a reduction in production domain of u. B-D: We can get clearer picture of how the concentrations of u are affected when we view the concentration profiles from a side elevation, where we see significant relative drop in the concentration of u between GrG (B) and GaR (D). E: If we set symmetric initial conditions then the group is symmetrically posed within the profile u and will remain stationary. This allows us to increase the chemotactic force c₀ to observe the effect on the group’s volume α. Over the range of c₀ shown we observe an decrease of ~17% in the groups volume.

If we consider again the two chemotactic mechanisms of GrG and GaR in light of this, then for GrG it should be clear that the force acts to expand the volume of the cells in the group, while for GaR the force acts compress the volume. More specifically, when the mechanism is GrG then force acts predominantly from inside to outside of the group, as the maximum concentration of u will always be within its outer boundary. Inversely when the mechanism is GaR the force acts from outside to inside, for exactly the same argument: u’s maximum is with the group’s outer boundary.

The net effect of compression and expansion on the velocity of the group then, or the time it takes the group to reach the end-point, is revealed when we recall that the group represents an activating region for u. If the group maintains an increase or decrease in volume, then so does the activating region of u, and consequently there will be, respectively, an increase or decrease in the chemotactic forces proportional this change. Therefore the discrepancy in time, is function of compression/expansion and the chemotactic force. These points are illustrated in Figure 3-12.

3.4.4 Group Motion for an Externally Produced Chemotactic Agent

In this section we are again concerned with active and passive mechanisms of motion for the group. As before the active mechanism is where the cells in the group are responsible for their own motion, that is, they are chemotactically reacting to the chemotactic agent u, the passive mechanism is where the group moves as a result of chemotactic forces acting on an external population, referred to as a tissue. We will continue this analysis, but with the difference that the agent is now produced by the surrounding tissue.
As with the previous section we are informed by our 1D analysis that Ra\(\text{G}\) is a feasible mechanism for motion of the group, and by arguments of symmetry given by (3.27) this implies that Ra\(\text{R}\) must also admit travelling solutions. If this point is not necessarily clear, then it is suggested to once again consider the scenario illustrated in Figure 3-10, but with the chemical externally produced. In addition there is a mechanism of Ra\(\text{G}\), however this mechanism can again be shown to be not feasible, by arguments of the previous section.

![Figure 3-13: Passive (Ra\(\text{R}\)) and active (Ra\(\text{G}\)) motion for a homogeneous group for an externally produced chemotactic agent.](image)

Initially the cells in the simulation are allowed to relax before the simulation begins proper at \(T = 0\). To initiate motion the group is initially attracted to a distant attracting point (white encircled cross) with constant force to allow the agent to saturate to a travelling profile until \(T = 2000\), after which the group is allowed to move independently. In both cases the parameter set is identical to those for an internally produced as, with changes only to who produces to the agent, the chemotactic mechanism, and who is reacting. In all other respects these simulations are identical to used in Figure 3-11. For passive (A) and active (B) motion we see once again that the group moves independently and uniformly from left to right marked by three time periods representing the beginning middle and end of each simulation. Again we notice that there is a distinct difference in time elapsed between the two mechanisms, however as suggested this can be explained by compression/expansion effects due to the chemotactic forces on the group. Model parameters are as in Figure 3-11.

In general then it would seem that for a change from internal to external production, we are merely inverting the problem, and this is clear when we consider, for example, that for internal we have Gr\(\text{G}\) and Ga\(\text{R}\) and for external we have Ra\(\text{R}\) and Ra\(\text{G}\). Clearly there is only an inversion of the type of cell that is reacting, and the type of cell that is producing, but not the chemotactic mechanism. From this point of view the story of a heterogeneous model is complete, and it remains to demonstrate the final mechanisms of Ra\(\text{R}\) and Ra\(\text{G}\) (Figure 3-13).

Once again we constructed two simulations, one for Ra\(\text{R}\) and one for Ra\(\text{G}\), using parameters identical to those used in the homogenous group with an internally produced chemotactic agent. Practically this literally entailed copying the simulation files and simply changing the population that produces the agent and the population that reacts to it. In both simulations we found almost identical results to those where the chemical is produced internally by the group, with a single exception that in the external case the simulations are apparently slightly faster than those of the internal case by ~30%.
Figure 3.14: Compression and expansion of the cells in the group due to the chemotactic force can increase or decrease production of the chemotactic agent and therefore affect velocity of the group. As with the case of internal production there is a net change in the volume of cells in the group due to the chemotactic force, that shows when the mechanism is active (A), there is a net increase in volume leading to an increase in chemotactic gradient (C) which consequently increase the velocity of the group. Inversely when the mechanism is passive (B) there is a net decrease in the volume of the group leading to decrease in the chemotactic gradient (C), consequently decreasing the velocity of the group.

However we account for this once again by arguments of compression and expansion due to the chemotactic force, and this is corroborated when we see that the relative differences between active and passive mechanisms, for both internal and external cases, shows an approximate four-fold difference in the time elapsed. Therefore if it were possible to remove the compression effects we would expect to see four identical results. For completeness the dynamics of the chemicals are illustrated to demonstrate the effects compression and expansion has on the group Figure 3.14, where we see clearly the significant drop in concentration for $RrR$ relative to $RaG$.

3.5 Heterogeneous Model of a Migrating Group

Possibly the most interesting feature of the heterogeneous models is that there is no permutation for which we cannot find a travelling solution, that is, no matter who is producing the chemotactic agent, or who is reacting to it the group can move coherently and uniformly. This is direct contrast the homogenous model where we showed that not all mechanisms are feasible, for example the two symmetric cases $RrG$ and $GrR$. However the heterogeneous model brings with it problems associated with differential adhesion, which is due in no small part to what we define a successful simulation to be.

Since the group is bisected into two distinct cell populations, we know from previous discussion that the primary differentiating factor of cell type in the CPM is differential adhesion, which allows us, by simple energy rules, to determine what one might call affinity between cells of differing types. However the group, regardless of heterogeneity, is still a group and specification of this “affinity” by differential adhesion needs to account for the sub-populations affinity to the group and affinity to their own populations. In other words, a successful simulation is one where the group (and by necessity is sub-populations) moves as a coherent whole, while preventing the sub-populations from dissociating into surrounding tissue, or indeed another sup-population.
In all the simulations that follow we began with a parameter set identical to the 1D analytical model, with the single exception that the chemotactic forcing parameter was found to optimal for $c_0 = 350$. This disparity can be explained by several factors include frictional forces due to contacts with a surrounding tissue, and of course effects of compression and expansion. Model parameters specific to the CPM will be identical to all previous simulations, with the exception of cell adhesion as suggested previously.

### 3.5.1 Group Migration for an Internally Produced Chemotactic Agent

In this section we consider that a sub-population of the group is responsible for the production of the chemotactic agent that can react with either another population in the group, itself, or the surrounding tissue. Unlike the homogeneous models, we do not require any initial conditions for the group to begin moving, such as an initial shift in the chemotactic agent relative the position of the group or a distant attracting point. In those models it was required to break the symmetry that occurs naturally. When the group is heterogeneous however, asymmetry is defined explicitly due to the composition of the group. Put another way, we can see that in all configurations we will have one population producing the agent, another reacting to it, and a third population that essentially breaks the symmetry, either by differential adhesion, opposing chemotactic forces or the approximately incompressible nature of the cellular medium.

From a physical point of view the compressible nature of the cellular medium, as we have shown, causes significant changes in volume of the group from increased pressure due to the chemotactic forces being applied. In those cases however, the forces were acting almost symmetrically on the whole group, which implies that as the force increases, the pressure increases, and without any means of releasing that pressure the volume decreases. In this case however, the chemotactic force is applied asymmetrically. In this sense when the chemotactic force acts to compress the reacting cells there is an equal and opposite expansion effect in another population, analogous to squeezing one half of a balloon.

Since we know that these mechanisms will always produce travelling solutions, all that remains is to illustrate such solutions which can be seen in Figure 3-15.
Figure 3-15: Simulations illustrating travelling solutions of the homogeneous group with an internally produced chemotactic agent. Here we demonstrate the four possible mechanisms of travelling solutions for an internally produced chemotactic agent, in each pair (left and right panels) illustrate both the cells at the beginning middle and end of the simulation, and the concentration field of the agent. (A-B): Blue cell producing the chemotactic agent that attracts the green cells. (C-D): Green cells producing the agent that repels the blue cells. (E-F): Blue cells producing the agent that repels the red cells. (G-H): Green cells producing the agent that attracts the red cells.
3.5.2 Group Migration for an Externally Produced Chemotactic Agent

When the chemical is produced by the external population then we have an inverted problem, however since we have already shown that solutions exist we will only illustrate solutions as we did in the previous section.

![Simulation illustrations](image)

**Figure 3-16**: Simulations illustrating travelling solutions of the heterogeneous group with an externally produced chemotactic agent. (A-B): Cells in the surrounding medium (red cells) are producing the chemotactic agent which acts to attract the blue cells in the group. (C-D): Again the red cells are producing the agent, but this time the green cells in the group are repelled.

3.6 Chapter Summary

The primary motivation of this chapter was to demonstrate, albeit on a qualitative basis, that we could corroborate the analysis of the previous chapter where we considered a 1D continuous model. To do this we first introduced a continuous numerical model to first illustrate the chemical dynamics and chemotaxis functions when the group is represented by a more realistic geometry that is a disk.

Using this representation we showed clearly that the dynamics are equivalent in both models, with the exception that when we extend our model to this new geometry, there is an inevitable change in the diffusive characteristics of the chemotactic agent. This change manifested as a discrepancy in the bifurcation point at which the travelling solutions emerge (Figure 3-6A). However the difference in the bifurcation, and indeed the resultant velocity profiles, was demonstrated to be a difference in the rates of diffusion we referred to as a diluting effect, due to the group being represented as a disk giving a volumetric difference due to curvature on the disk boundary. However we accounted for this showing that if the size of the disk (and line segment in 1D) are steadily enlarged then the effect of this curvature would become negligible, and both 1D and 2D models would become equal (Figure 3-6B).

Due to computational complexity in the 2D model, increasing the size of the disk quickly become computationally intractable, and so we turn to a polar representation that admitted solutions in the form of modified Bessel functions. Using this solution we could demonstrate how the concentration values changed on the boundaries between 1 and 2D models, in a steady state solution, which clearly showed that
as both the sizes of the groups were increased in both models, the concentrations on the boundaries of
the groups saturated to a common value, indicating that the effect of curvature had become negligible
(Figure 3-4), and therefore the velocities will become equal.

Notwithstanding these differences the biggest difference between the 1D and 2D model was due
to the introduction of an external surrounding population of cells. In the 1D model we had derived an
expression for the chemotactic function that was dependent on the size of the group, and this became
insoluble when considered the external population to be reacting to the chemotactic agent (3.26). However in the CPM we had no such barriers and so we could introduce two new simulations that were
not possible in the 1D models, which gave four mechanisms of group motion, given by two pairs
symmetric problems: $GaR/GrG$ and $RrR/RaG$. In all cases we demonstrated again that for identical
parameter sets, all of these models produce travelling solutions for the group.

However while we could demonstrate that such solutions exist, there was a significant difference in
each symmetric pair when the external population was reacting or the group itself. This manifested as a
near four-fold increase in velocity of the group relative to when the external population was reacting,
(compare Figure 3-11 and Figure 3-13). However this was shown to be a function of compression and
expansion between the mechanisms. When the group is reacting to an external chemical there is a net
reduction in volume of the group, directly affecting the production domain of the agent, leading to drop in
velocity. Conversely when the group is reacting to an internal chemical there is a net increase in the
volume of the group which increases the production domain of the agent and of course increasing the
velocity.

When we introduced heterogeneity into group, we found that there were no permutations we
could not find a travelling solution for. And in a simple sense there is very little to say about these
simulations, however they do have far reaching implications in other studies, such in primitive streak
progression in gastrulation in the chick embryo, however in the context of this chapter we merely illustrate
the results verbatim, as dynamically these results are quite trivial.

Finally we have demonstrated that there are a variety of chemotactic mechanisms that can lead to
group motion, and in a more morphologically realistic setting (CPM), demonstrated all that was predicted
in the 1D analysis and realized new mechanisms previously thought not possible. In the proceeding
chapter we will see through an experimental study, how the mechanisms we have demonstrated here and
in the previous chapter, can explain early developmental processes in the embryogenesis of chick egg.
Chapter 4

COORDINATION OF CELL DIFFERENTIATION AND MIGRATION IN MATHEMATICAL MODELS OF EMBRYONIC AXIS EXTENSION

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4.1 Abstract

Vertebrate embryos display a predominant head-to-tail body axis whose formation is associated with the progressive development of post-cranial structures from a pool of caudal undifferentiated cells. This involves the maintenance of active FGF signaling in this caudal region as a consequence of the restricted production of the secreted factor FGF8. FGF8 is transcribed specifically in the caudal precursor region and is down-regulated as cells differentiate and the embryo extends caudally. We are interested in understanding the progressive down-regulation of FGF8 and its coordination with the caudal movement of cells which is also known to be FGF-signaling dependent. Our study is performed using mathematical modeling and computer simulations. We use an individual-based hybrid model as well as a caricature continuous model for the simulation of experimental observations (ours and those known from the literature) in order to examine possible mechanisms that drive differentiation and cell movement during the axis elongation. Using these models we have identified a possible gene regulatory network involving self-repression of a caudal morphogen coupled to directional domain movement that may account for progressive down-regulation of FGF8 and conservation of the FGF8 domain of expression. Furthermore, we have shown that chemotaxis driven by molecules, such as FGF8 secreted in the stem zone, could underlie the migration of the caudal precursor zone and, therefore, embryonic axis extension. These mechanisms may also be at play in other developmental processes displaying a similar mode of axis extension coupled to cell differentiation.

4.2 Introduction

During embryonic development, generation of cell diversity needs to be coordinated with tissue growth in order to achieve the right size, cell number and shape of the different organs. Depending on the developmental context this is implemented differently. Several developmental systems with predominant growth along one axis share a similar strategy: cells at one end of the domain remain undifferentiated and give rise progressively in time and space to cells that have a more restricted fate and can differentiate further. This occurs for example during growth of plant root meristemes, caudal extension of short germ band insects and worms, extension of the vertebrate limb bud, growth of bones, and caudal extension of the vertebrate body axis [31, 32, 33, 34, 35, 7]. In this paper we focus on the latter process, namely we are interested in understanding how the migration and differentiation of cells associated with the caudal extension are controlled at the molecular and cellular level.

Vertebrate embryos display very important differences along their rostro-caudal (head-to-tail) axis from very early stages of development which are manifested, for example, by the orientation and movement of the primitive streak along the rostro-caudal axis. This is a transient structure, composed of cells that form a groove in the epiblast, through which cells ingress to form the mesoderm and the endoderm. The primitive streak displays a rostral tip (named Hensen’s node), which has an important pattern organizing role on the cells that develop in its vicinity and influences the primitive streak.
dynamics. Primitive streak development goes through an initial phase of rostral elongation followed by caudal regression.

Formation and rostral elongation of the primitive streak is associated with cell movements that may have a lateral intercalation component \[36\] or be of chemotactic nature \[4, 9\]. Regression of the primitive streak is associated with the movement of a group of cells surrounding and including Hensen’s node, that behaves as a precursor region for postcranial mesoderm and neural tube. Although some stem-like cells giving rise to several lineages may reside in this caudal precursor region, different populations have been discovered to give rise preferentially to distinct lineages. The mesodermal layer of Hensen’s node gives rise to the notochord while the rostral primitive streak gives rise to somites. The ectodermal layer of Hensen’s node gives rise to the floorplate of the neural tube while the ectoderm adjacent to the primitive streak gives rise mainly to lateral (non-floorplate) neural tube \[37\] and some somitic tissue \[38, 39, 40, 41\]. Cells in this region proliferate and their daughter cells can either continue to move caudally and remain in the caudal precursor region as the streak regresses or can be left behind and consequently exit this region (Figure 4-1).

![Figure 4-1: Progressive down-regulation of FGF8 at the caudal precursor zone.](image)

Schematic drawing showing expression of FGF8 (purple) in embryos of 10 (A) and 14 (B) somites respectively. Transcription of FGF8 (red) only occurs at the primitive streak and adjacent epiblast but FGF8 mRNA extends into the presomatic mesoderm and adjacent neural tube due to maintenance of the transcript as the embryo extends. Cells which are left behind the moving caudal neural precursor zone (blue dot) do not regress caudally and stop transcribing FGF8.

In general, it is thought that cells either remain in the caudal precursor region or transit to a more differentiated state depending on the degree of activation of signaling pathways which in turn depends on their exposure to specific morphogens produced by particular cell populations. A precise molecular marker for this precursor population has not been described, but in the epiblast layer, according to fate maps, it may correspond to cells that transcribe FGF8 as detected with the FGF8 intronic probe \[6\]. We
will refer to this population as the caudal neural precursor region (CNPR) (which includes the caudal lateral epiblast [42], and the epiblast layer of the node and gives rise primarily to neural tube although it also contains mesoderm precursors). FGF8 is not just a marker of the caudal precursor region but it is also a crucial player in the regulation of cell maturation within its domain of influence. Cells with active FGF signaling pathway remain undifferentiated, both in the neural plate and in the mesoderm, while those with low or no activation of FGF signaling can progress to a more differentiated state (if the right signals are present) [5, 43]. It is therefore important to understand how this signaling pathway is regulated and in particular how the production of FGF8 at both the protein and mRNA levels is controlled.

Some aspects of the regulation of FGF8 expression are known. FGF8 mRNA is characterized by high stability so that cells that have stopped transcription of the gene can maintain its expression for a considerable time interval resulting in a graded distribution of the RNA in the extending axis [6]. Figure 4-1 illustrates that although FGF8 transcription takes place in the CNPR, the area where FGF8 mRNA is present extends further rostrally. FGF8 levels can be down-regulated by retinoic acid (RA) that is secreted from somites and this could in theory be sufficient for the progressive down-regulation of FGF8 [5]. However, in the absence of RA, FGF8 mRNA is still progressively down-regulated [5] although its region of expression is expanded.

Although mechanisms responsible for the control of FGF8 transcription remain unknown, it is clear that they must be coupled to caudal extension of the embryonic axis, a crucial process that takes place as FGF8 is down-regulated. Caudal extension involves movements in all three embryonic layers that rely on different cellular behaviors that are region and embryo dependent. Many efforts have been made to understand the mechanisms that regulate convergence and extension of the mesodermal layer in fish and frogs where region-specific cellular behavior such as directed migration towards the midline (due to cellular intercalation) have been described [44]. More recently, a random cell motility gradient has been observed in chick presomitic mesoderm that contributes to axial elongation [45]. In addition, other phenomena such as stem-cell like mode of growth [46] reviewed in [42] and active movement of cells towards the caudal end [47] have been identified for neural plate and notocord elongation respectively. Extension of the embryo constitutes, therefore, a multi-factorial process where all these aspects of cell behavior are coordinated [44].

In this paper we will focus on two main features of vertebrate embryonic axis extension, namely progressive generation of cells not producing FGF8 and migration of the caudal precursor zone. We will use mathematical methods to analyze these processes.

Concentration dynamics of FGF8 and RA during caudal extension in chick embryo have been modeled previously in [48]. It was shown there that the dynamics of the concentration profiles of FGF8 and RA could, in theory, be explained by specific interactions (mutual inhibition) between FGF8 and RA which can be described by the system of nonlinear partial-differential equations having a propagating front solution.
The ability of local self-enhancement and long-ranging inhibition of morphogen gradients to give rise to a propagating front-like behavior has also been addressed in [49], where it was suggested that stationary patterns (Turing) form due to the growth of the medium (tissue grow due to cell proliferation).

The main feature of our approach is that we take into account the movement of the FGF8 production domain and consider its effect on the dynamics of the FGF8 concentration profile, as well as the effect of FGF8 concentration profile on the differentiation and movement of cells. We perform our study using two distinct models. First, we develop and consider different modifications of a continuous one-dimensional model to check hypotheses concerning dynamics of morphogens and mechanisms of cell motion. Then we verify the obtained results by use of a multi-cell simulation method (the Glazier-Graner Hogeweg model, the GGHM, aka the Cellular Potts model or CPM) originally developed by Glazier and Graner [20, 30] and recently used to simulate and analyze the migration of cells in various biological tissues [50] including the formation of cell flows at the early stages of the chick embryo gastrulation [4].

Furthermore, with new experimental observations, we analyze modeling outcomes and further explore the mechanisms that underlie progressive down-regulation of FGF8 and its role in the caudal precursor zone migration. We focus on the events that occur in the epiblast region that will give rise to the spinal cord (the CNPR) as this is a tissue where the regulation of FGF8 transcription occurs but similar interactions may be relevant for mesoderm maturation. Based on our modeling and experimental results we suggest that the movement of the caudal precursor region is essential for the observed dynamics of the concentration patterns of involved morphogens, and that the interplay between these morphogens and the cells producing them is responsible for the progressive generation of differentiated cells as well as for the migration of the CNPR. We also show that the integration of cell proliferation, differentiation and movement allows the CNPR to maintain a constant size and preconditions the constant speed of its migration so that the moving stem zone regulates regression of the primitive streak. Table S1 outlines the summary of our models.
Table 1: Summary of models used. Using continuous one-dimensional and individual-based two-dimensional models we have considered migration of the domain of transcription (DoT) under a few distinct sets of assumptions concerning proliferation, differentiation and movement of cells forming the DoT. Summary of the mechanisms with references to the figures and supplementary movies demonstrating simulation outcomes has been provided. Again $\mathcal{A}$ is the externally diffusing chemotactic agent/transcript of an intracellular gene $\mathcal{A}$-mRNA and $T_A$ is a constant representing the threshold of $\mathcal{A}$ at which differentiation occurs.

The mechanisms of embryonic axis extension we propose here may also be applicable to different systems where production of a morphogen by a domain of moving cells is responsible for progressive differentiation.

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<th>Schematic</th>
<th>1D or 2D, reference to figure, movie</th>
<th>Differentiation</th>
<th>Domain of movement</th>
<th>Directionality</th>
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<td>DoT</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>No</td>
</tr>
<tr>
<td><img src="image2" alt="Schematic" /></td>
<td>1D, Figure 4-3, Movies S2, S3</td>
<td>A-self repression: $T_A$ threshold</td>
<td>DoT</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>No</td>
</tr>
<tr>
<td><img src="image3" alt="Schematic" /></td>
<td>1D, Figure 4-5</td>
<td>A-self repression FGF8 activation by $\mathcal{A}$</td>
<td>DoT</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>No</td>
</tr>
<tr>
<td><img src="image4" alt="Schematic" /></td>
<td>1D, Figure 4-5</td>
<td>A-self repression FGF8 repressed by $\mathcal{A}$</td>
<td>DoT</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>No</td>
</tr>
<tr>
<td><img src="image5" alt="Schematic" /></td>
<td>2D, Figure 4-6, Movie S4</td>
<td>A-self repression: $T_A$ threshold</td>
<td>Only DoT cells can move</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>Yes</td>
</tr>
<tr>
<td><img src="image6" alt="Schematic" /></td>
<td>2D, Figure 4-7, Movie S5</td>
<td>A-self repression: $T_A$ threshold</td>
<td>No movement when $\mathcal{A}$ goes below threshold</td>
<td>Imposed directionality</td>
<td>Constant speed, $c$</td>
<td>Yes</td>
</tr>
<tr>
<td><img src="image7" alt="Schematic" /></td>
<td>1D, Figure 4-8, Movie S6</td>
<td>Not considered</td>
<td>DoT</td>
<td>Initial bias and then towards low $\mathcal{A}$: chemorepulsion</td>
<td>Dependent on $\mathcal{A}$ concentration gradient: chemorepulsion</td>
<td>No</td>
</tr>
<tr>
<td><img src="image8" alt="Schematic" /></td>
<td>2D, Figure 4-9, Movies S8-S10</td>
<td>Not considered</td>
<td>Only DoT cells are considered and can move</td>
<td>Initial bias and then towards low $\mathcal{A}$: chemorepulsion</td>
<td>Dependent on $\mathcal{A}$ concentration gradient: chemorepulsion</td>
<td>No</td>
</tr>
<tr>
<td><img src="image9" alt="Schematic" /></td>
<td>2D, Figure 4-10, Movie S11</td>
<td>A-self repression: $T_A$ threshold</td>
<td>Only DoT cells can move</td>
<td>Initial bias and then towards low $\mathcal{A}$: chemorepulsion</td>
<td>Dependent on $\mathcal{A}$ concentration gradient: chemorepulsion</td>
<td>Yes</td>
</tr>
</tbody>
</table>
4.3 Results

4.3.1 Concentration profiles in the continuous one-dimensional model

We posed the general theoretical problem of what simple regulatory network could account for the restricted transcription of a gene within a domain of constant size considering that cells that transcribe the gene proliferate and move as the main axis of the embryo extends. To address this problem we first developed a continuous one-dimensional model (installation, templates for basic simulations and source codes are available from the web site: http://pcwww.liv.ac.uk/~mf0u4027/biochemsim.html). The simplest version of the model includes two variables: one for dynamics of hypothetical A-mRNA, transcribed exclusively in a fixed-sized domain that moves (say to the right) with constant speed \(\ell\), and one for the secreted protein it encodes (protein \(A\)). The basic model therefore considers concentrations of the following two species:

1. mRNA (non-diffusible) which is maintained (produced) at a constant level exclusively in a domain of constant size moving with a constant speed. Further on we will refer to the domain of mRNA transcription as to the DoT.
2. Protein A (diffusible) whose production rate is proportional to the level of A-mRNA.

Figure 4-2 shows the stationary concentration profiles of both species with the assumption that the decay rates are given by linear functional terms (see the description of 1D model in the Materials and Methods Section). The transition process from the initial conditions (when both concentrations are zero everywhere except for the DoT where the concentration of A-mRNA is one) to the stationary solution is shown in Movie S1. Also, since the DoT is moving, the concentration profiles of A-mRNA and protein A do not form symmetric pattern with respect to each other: A-mRNA decays gradually behind the DoT and the maximum in protein \(A\) profile lags behind (shifted to the left in the graph) the midpoint of the DoT. This shift becomes more pronounced with the increase of the DoT speed, \(\ell\), and depends on the kinetic rates of A-mRNA and protein \(A\) (see equation (4.6), Material and Methods). Due to this shift the maximum in concentration of the protein can lie outside the DoT (see Figure 4-2, also Figure 4-3B). The condition for this is given by inequality (4.7) in the Materials and Methods Section.
4.3.2 Self-regulation of the size of the DoT via negative feedback

So far in our model the size of the DoT (which reflects the number of cells transcribing A-mRNA) has been fixed. Now we would like to take into account that the cells forming the DoT proliferate and differentiate (i.e., can stop transcribing A-mRNA under the appropriate conditions). When proliferation is taken into account the DoT size should gradually increase over time unless a regulative mechanism ensures this does not occur. The shape of the concentration profile of protein A in Figure 4-2 gives an idea of a possible and simple mechanism for regulating the size of the DoT that would not involve any component external to the system. If we assume that cells stop transcription of A-mRNA when the level of protein A rises above some threshold, $T_A$, (see the diagram on Figure 4-3A), this would define the position of the left side border of the DoT (as a coordinate of the point where the level of protein A gets above $T_A$) while the position of the right side border is predefined and given as a coordinate of a point moving to the right with speed $c$ (see Figure 4-3B). Now the size of the DoT is defined by the negative feedback loop where protein A inhibits the transcription of A-mRNA.
In this version of the model the size of the DoT is defined by the value of the threshold \( T_A \): when the concentration of the protein gets above \( T_A \), transcription of the \( A \)-mRNA stops and this eventually defines the DoT size. This mechanism of the DoT size regulation works if the value of parameter \( T_A \) is below the maximum possible value of protein \( A \) concentration (which is \( k_3/k_2 \), see equations (4.2) and (4.4) in the Materials and Methods section). Obviously, the size of the DoT is small for low values of \( T_A \) and increases with \( T_A \). Simulations and analysis of the model show that the size of the DoT is generally an increasing function of the DoT’s speed (see Figure 4-3C). Simulations also indicate that this size is not necessarily a constant and can oscillate over time (compare Movies S2 and S3 showing formation of a
DoT of stationary and oscillating sizes). Oscillations are observed when the kinetics rate of $A$-mRNA ($k_4$) or protein $A$ (note that for simplicity we presume that $k_3 = 2k_2$ in all simulations, i.e. the rates of protein $A$ production and decay are varied in a proportional manner) are too small with the transition (bifurcation) value depending on the speed of migration of the active transcription domain (Figure 4-3D). Comparison of the numerical and analytical results indicates that the domain in the model parameter space, where oscillations are observed numerically, strongly correlates when no stationary solution exists according to analytics (compare dashed blue and solid red lines in Figure 4-3D and see inequality (4.10) in the Materials and Methods Section).

4.3.3 Size regulation of the FGF8 domain of transcription

So far, we have kept our model general and have not named the molecules that would be represented as morphogen $A$. Going back to the CNPR, we are interested in understanding the mechanisms that regulate the size of the FGF8 transcription domain and therefore the CNPR. The simplest possibility would be that FGF8 corresponds to morphogen $A$. In this case, based on the results presented in Figure 4-3, we could suggest that high FGF8 levels stop transcription of its own gene and thus regulates the size of its domain of transcription.

Previous reports suggest that FGF8 may be able to promote the stability of its own mRNA transcript [6], but no experimental evidence for its influence on the rate of its own transcription has been found. In order to examine the dependence of FGF8 transcription on FGFR signaling, we treated chick embryos with the FGFR antagonist PD173074 for 4h [51]. This treatment did not change significantly the

Figure 4-4: FGF8 transcription is not altered by FGF signaling. A–B: FGF8 transcription at the caudal precursor zone in control (A) and FGFR antagonist treated (B) chick embryos. No changes in FGF8 transcription are observed following a blockade of FGF signaling. C: Schematic drawing showing the origin of the explants shown in D–E. D–E: FGF8 transcription in caudal precursor zone chick explants following culture in the presence of control (D) and FGF4 containing media (E).
domain of FGF8 transcription (compare panels A and B in Figure 4-4) (n = 6). In addition, the treatment of cultures of the caudal precursor zone with FGF4 (which activates FGF receptors more efficiently than FGF8 [52]) for 9h did not alter FGF8 transcription (n = 3, Figure 4-4 panels D and E). These results in the chick embryo are consistent with the phenotype of FGFR1 mutant mouse embryos where the expression of FGF8 in the caudal precursor region is not altered [53]. Therefore we conclude that FGF8 is not self-repressing.

![Figure 4-5](image)

**Figure 4-5:** Possible mechanisms of the involvement of FGF8 in the caudal gene regulatory network. A-B: as before the concentration of protein A is proportional to its transcript A-mRNA given my equations (4.1) and (4.2) (see the Materials and Methods Section for details). However we now impose a feedback control on transcription by regulating the size of the A-mRNA DoT due to a threshold of protein A’s concentration (T_A) modeled in equation (4.8), where the size of the DoT is now a function of model parameters and not constant and is given by equation (4.9). The rate of FGF8 transcription is proportional to the level of protein A (see equations (4.11) and (4.12)). Note that the FGF8 DoT extends behind the A-mRNA DoT. C-D: the transcription of FGF8 and A-mRNA are launched independently in (roughly) the same group of cells, while both down-regulated by the same signal provided by protein A (again see equations (4.1) (4.2) and (4.11) (4.12) coupled with condition (4.8)). Note that the concentration profiles for FGF8 DoT and A-mRNA DoT in this case basically coincide. Values of parameters (in equations (4.1) and (4.2)) are the same as in Figure 4-2. For extra parameters (equations (4.11) and (4.12)): D_A = 0.5, k_{31} = k_{41} = 0.0003k_{32} = k_{42} = 2k_{31}

In terms of our model this means that FGF8 dynamics could be regulated by the caudal self-repressing morphogen A. We have explored several possible relationships between this self-repressing morphogen (protein A) and FGF8. If FGF8 transcription was activated by protein A then the profile of FGF8 transcription (and therefore the extent of the CNPR) would lag behind the domain of the A-mRNA transcription (Figure 4-5A–B). Alternatively, if the transcription of A-mRNA and FGF8 is initiated in a similar caudal domain and protein A is repressing simultaneously the transcription of both, then the DoT of FGF8 would coincide with (or at least will not significantly differ from) the DoT of A-mRNA (Figure 4-5 C–D). Both possibilities are feasible in principle but for simplicity, in the following sections, we will
consider the latter option, where the domains and concentration profiles of morphogen A and FGF8 are equivalent.

4.3.4 Maintenance of the migrating DoT size in GGHM

We have explored some features of the migrating DoT by means of a continuous 1D model. However this model does not provide the most appropriate framework to model cells that are proliferating and moving and therefore we have extended our study by developing and using an individual-cell based model represented by Glazier-Graner-Hogeweg Model (GGHM) [28]. This modeling approach allows us to test more carefully the phenomena emerging from the individual cell behaviors.

The epiblast caudal precursor region in the chick embryo consists of a unicellular layer of cells and therefore we can use the 2D version of the GGHM to capture events taking place over the CNPR (here we are not dealing with the influence of external signals coming, for example, from the mesoderm). Installation of the program and templates for reproduction of our simulations are available from http://pcwww.liv.ac.uk/~mf0u4027/biocellsim.html.

The version of the GGHM, which corresponds to the 1D model used above (see Figure 4-3), includes two cell types only (Figure 4-6A–B): cells transcribing A-mRNA (red) and cells which do not transcribe it (green). The dynamics of A-mRNA and protein A are defined the same way as in the 1D model except for: (a) equations (4.1) and (4.2) (see Materials and Methods Section) are written for the laboratory frame of reference (c = 0) and (b) diffusion term in equation (4.2) is extended into 2D. To be in line with the differentiation mechanism suggested for 1D model we assume that the red cells proliferate and convert into green cells when the level of protein A reaches the threshold value, $T_A$. We also attribute motility properties to cells, namely, we presume that red cells move actively while green cells do not and can only follow red cells due to adhesive contacts.

We start the simulation with a group of red cells (the DoT) moving in a particular direction (to the right in Figure 4-6B) under the influence of a preset force. This force is given by the extra term $E_f = -\beta_f (x \cdot l)$ (where $x$ - shift of red cell’s interface and $l$ – the unit vector pointing to the right) in the definition of the energy which counts for the work done by the horizontal force applied to moving cells. This permits us to leave the study of the mechanisms of cell motion for later (see below). While moving and proliferating, cells in the DoT transcribe A-mRNA which in turn allows the production of protein A (see Figure 4-6C and D). In places where the level of protein A reaches its threshold level, $T_A$, red cells differentiate into green. Simulations show that, while moving and proliferating, red cells (forming the DoT) leave a trail of differentiated green daughter cells (Figure 4-6B and Movie S4) very similar to what happens in the embryo where the CNPR gives rise to more mature tissue progressively. The size of the DoT (number of red cells) is regulated by the kinetics of both A-RNA and protein A: increasing either kinetics constant $k_1$ or $k_2$ (assuming for the latter that $k_3 = 2k_2$) decreases the size (area) of the DoT (see Figure 4-6E).
4.3.5 Promotion of cell migration by a caudal morphogen

The simulations presented in Figures 4-2, Figure 4-3, Figure 4-5 and Figure 4-6 were performed under the assumption that the ability of cells to move correlates strongly with their ability to transcribe the $A$-mRNA gene so that the high level of protein $A$ switches off both abilities of a cell. However other mechanisms that relate motility of cells to morphogen concentration may fit better to experimental results and known signaling molecules produced in the caudal precursor zone, in particular FGF8. Our 2D (GGHM) model can be used to check some of these mechanisms.

In our model we were dealing with a hypothetical protein $A$ rather than FGF8, but as we have previously explained, if protein $A$ down-regulates the transcription of both $A$-mRNA and FGF8, the concentration profile of protein $A$ is equivalent to that of FGF8. The signaling pathways that regulate the movement of cells in the caudal precursor zone are not well established, although it is known that FGF signaling controls the ability of spinal cord precursor cells to move [54]. Down-regulation of FGFR signaling in one cell promotes its exit from the CNPR which suggests that FGF signaling keeps cells moving and allows them to accompany the regressing Hensen’s node [55].

![Figure 4-6: The DoT migration in the GGHM. A: Schematic diagram of the used model (identical to the diagram in Figure 4-3). B: Three consecutive images from the simulation of primitive streak regression. Initially there is a group of 25 red cells (the DoT) forming a square tissue. The level of $A$-mRNA is high and constant in all red cells. Red cells move (to the right), proliferate and differentiate, i.e. red cell transforms into the green cell when the level of protein $A$ at any point inside the red cell gets above the threshold value $T_A = 0.8$. Green cells do not move nor produce $A$-mRNA, for simplicity we have also assumed that they do not proliferate. Cell differentiation is regulated by the level of morphogen $A$ (as in Figure 4-3). Parameters: $k_1 = 0.001$, $k_2 = 0.003$, $\beta = 4.5$. C, D: Concentrations of $A$-mRNA (C) and protein $A$ (D) are shown in shades of red. The border line between red and green zones is along an isoline in the concentration field of protein $A$ corresponding to the threshold value $T_A = 0.8$. E: Increase in the rate of $A$ kinetics, $k_2$, (assuming that $k_3 = 2k_2$) reduces the size of the DoT (or number of cells forming the DoT) exponentially.]
To incorporate this feature into our model we decouple the ability of cells to transcribe $A$-mRNA from their ability to move and we introduce an intermediate cell type: cells that do not transcribe $A$-mRNA but can move (blue cells in Figure 4-7). Therefore differentiation of red cells into green cells takes place in two steps. Step 1: we assume that transcription of $A$-mRNA is down-regulated by protein $A$, i.e. production is stopped when the level of protein $A$ achieves some threshold level $T_A$ (analogous to 1D model represented in Figure 4-3A and Figure 4-5C). Step 2: all cells transcribing $A$-mRNA move and those that do not transcribe $A$-mRNA keep moving until the level of protein $A$ gets below another threshold $T_M$ ($T_M < T_A$). The shape of tissue formed by cells forming the DoT and their descendants (all daughter cells) is given on Figure 4-7B (see also Movie S5). Comparing Figure 4-6B and Figure 4-7B we can see that both modifications of the model allow regulation (stabilization) of the DoT size.

### 4.3.6 Chemotactic mechanism for the DoT migration

Up to now we were presuming that the DoT is moving and that the speed of its motion is given by the preset parameter $c$. Examination of Figure 4-3 reveals a possible mechanism of this motion. Assume that the cells forming the DoT are chemotactic to a morphogen they produce. For example, we can assume that the speed of the DoT migration is proportional to the gradient of $A$, i.e. $\nabla A = c_0 \frac{\partial A}{\partial x}$. The gradient can be taken at some specific point, for example at the right border of the DoT, or we could use an average gradient over the entire DoT, i.e. the difference between concentrations of the protein $A$ on two borders of the DoT divided by the size of the DoT. Computer simulations show that both these assumptions can cause the DoT migration with constant speed and therefore the motion of the CNPR can have a chemotactic nature. Simulations with the first assumption, i.e. the speed of the DoT is defined by the gradient of $A$ on one particular side, show that starting from a wide range of initial conditions (and also for a wide range of values of $c_0$), we obtain (after some transition period) a DoT migrating moving with constant speed (see Movie S6). Simulations with the second assumption, i.e. that the speed is defined by the average gradient of morphogen $A$ over the DoT, also show the desired behavior, but we need to apply special initial conditions: for example, we force the DoT to move for some initial time and then switch this force off and chemotaxis on (see Movie S7).
Figure 4-7: The DoT migration in the GGHM with 3 cell types. A: Schematic diagram of the used model. B: Snapshots from the simulation: red cells move and produce A-mRNA, blue cells move but don’t produce A-mRNA and green cells do not move and do not produce A-mRNA. Red cells transform into blue when the level of protein A rises above $TA = 0.75$, blue cells to green when the level of protein A drops below $TM = 0.6$. Only red cells proliferate. Parameters: $k_1 = 1023$, $k_2 = 5 \cdot 10^{-4}$, $\beta = 3.8$. C, D: concentration profiles of A-mRNA (C) and protein A (D) after 20500 time steps of simulation.

Thus the migration of the DoT can be due, in theory, to chemo-repulsion of its constituent cells by protein A. This mechanism of migration is very similar to that of so called “ballistic” motion of a point which is a source of its own chemo-repellent [56]. In our case this “ballistic” effect is even more profound: the chemorepellent is produced not only inside the DoT but also behind it (where the concentration of A-mRNA is nonzero) and this adds to the difference between concentrations of the chemorepellent at the front and back sides of the DoT. Simulations as well as analytical studies of the model show that the migration (with constant speed) of a self-repelled DoT is only possible when the parameter $c_0$, defining the strength of chemotaxis, is above a certain threshold (see Figure 4-8A, where this threshold is roughly 0.6). The concentration profiles of A-mRNA and protein A, as well as the size of the DoT, depend on the parameter $c_0$ similar to their dependences on the parameter $c$ in the non-chemotactic version of the model (equations (4.8)–(4.10)) with fixed speed of the DoT migration (compare Figure 4-3C with Figure 4-8B).
Figure 4-8: The DoT migration due to chemotaxis in 1D model. The speed of the DoT migration is defined by the formula $c = c_0(A_l - A_r)$ where $A_l$ and $A_r$ are concentrations of protein $A$ on the left and right borders of the DOT. A: The version of the model where the DoT size is fixed (chemotaxis without mRNA self-repressive production control). The domain in the parameter plane “$c_0$ versus $c_0$” where the travelling DoT should definitely exist according to the analysis of the model is on the right side of the solid red line (this line represents the border of the domain defined by inequality (4.16) in the Materials and Methods section. Transition points (between existence and nonexistence) of migrating DoT’s in simulations are given by blue markers, dashed blue line connecting these markers gives the numerically obtained border. B: The version of the model where the DoT size is controlled by the protein $A$ (chemotaxis with mRNA self-repressive production control). The size of the DoT depends on the parameter $c_0$. The difference $A_l - A_r$ depends on $c_0$ and saturates when $c_0 \rightarrow \infty$ giving a linear asymptotic (red) for the dependence of the DoT size on $c_0$.

The role of chemotaxis in the migration of caudal precursor zone has not yet been addressed experimentally, but it is known that FGF8 can act as a chemo-repellent upon mesenchyme cells during gastrulation in the chick embryo [9], and that down-regulation of FGF signalling does not allow the caudal movement of cells following node regression [54]. Although, in our model, we consider the self-repressing morphogen $A$ as the chemo-repellent for cells forming the DoT, the same result would be obtained if we consider that the actual repellent is FGF8 whose dynamics coincides with that of protein $A$ (as discussed above).

4.3.7 Chemo-repulsion in GGHM

As we have noted in the previous section, the migration of the DoT can have a chemotactic nature and, in addition, there is strong evidence that FGF8 can act as a chemorepellent in several contexts [57]. Using GGHM we can analyze this problem to a much greater extent than was possible in the framework of the 1D model.

Let us first consider a simplified problem by ignoring cell proliferation and differentiation. Assume that the DoT is represented by a group of (non-proliferating) cells which produce some chemotactic agent (protein $A$ or FGF8, in our case), such that the cells are repelled by this agent. Is it possible that this group of cells will migrate (move along a straight line) due to this repulsion and thus reproduce migration of the DoT? Our simulations show that the group of cells repelled by a chemical they produce can exhibit three types of behavior (Figure 4-9). Cells can stay as a compact group and move randomly or meander with little net relocation (Figure 4-9B, Movies S8) or exhibit oriented motion, as in the case of the CNPR (Figure 4-9C, Movie S9). Movement of cells can deform the shape of the DoT (Figure 4-9D, Movie S10) or even break it so that they form a few smaller groups of cells each moving independently. The type of
observed behaviour is defined by model parameters and can be altered by varying the number of cells, their adhesiveness (defined by the adhesion matrix $\mathcal{J}$), kinetics rates of protein $A$ (production $k_3$ and decay $k_2$), $A$-mRNA (decay $k_1$) and chemotactic forcing, $\beta$, (see Materials and Methods Section). Generally, the model’s parameter space can be represented as a collection of domains corresponding to each type of observed dynamic behavior. Figure 4.9E shows the location of these domains on a parametric plane corresponding to two key parameters (responsible for the type of dynamics exhibited by self-repelling group of cells), namely chemotactic force as defined by parameter $\beta$ and protein $A$ decay rate, $k_2$ (see Materials and Methods Section). When the chemotactic forcing is weak ($\beta$ is less than some threshold value, and this threshold depends on $k_2$) the group of cells meanders and shows no net migration. The meandering behaviour is intrinsic to GGHM (corresponds to thermal fluctuations when $T > 0$) and it is a counterpart of the resting DoT in continuous 1D model: as we saw previously the DoT in 1D model does not migrate (or no travelling solutions exist) when $c_0$ is below than some threshold value (see inequality (4.16)) and this threshold depends on the DoT size (see Figure 4-8A). A meandering group of cells starts to move along a straight line when we increase the chemotactic forcing (by increasing the parameter $\beta$) or the protein $A$ decay rate, $k_2$. This type of behaviour is also in a line with our observations on 1D model where the DoT starts to migrate when chemotactic forcing $c_0$ is above some threshold value (Figure 8A). On the other hand further increase in either of these parameters results in deformation of moving tissue so that rounded tissue transforms into an umbrella-like shape. There is no 1D counterpart for this kind of behaviour.
Thus, the migration of the A-mRNA DoT can be explained by the chemotactic response of its constituent cells to protein A (i.e. the FGF8 DoT migrates due to repulsion by FGF8). Now we can put proliferation and differentiation of cells back into this model and adjust model parameters so that we observe oriented motion of tightly packed red cells leaving the trail of differentiated daughter cells (Figure 4-9F, see also Movie S11). The result from this simulation mimics the regression of the CNPR indicating that the interactions we have considered are sufficient to account for the observed maintenance of a compact group of cells that proliferate, migrate and differentiate during vertebrate embryonic axis extension.

4.3.8 Experimental study of regulative properties of the FGF8 DoT

In order to challenge the ability of the model to reproduce experimental results, we have performed an experiment where the FGF8 DoT (which in our model is equivalent to the A-mRNA DoT) was split into two and the changes in the expression of FGF8 were analyzed after 20h culture (Figure 4-10A, B). In the rostral moiety, FGF8 was maintained caudally suggesting that FGF8 does not require signals from the caudal-most region of the embryo for its maintenance. In addition, this experiment also
shows that the capacity of FGF8 to progressively down-regulate is also intrinsic to the caudal moiety (Figure 4-10B).

**Figure 4-10: Regulative properties of the FGF8 DoT.** FGF8 expression can be maintained in the absence of caudal-most signals and can be progressively down-regulated in the absence of rostral signals. A: Schematic diagram showing the experimental separation of the rostral and caudal parts of the FGF8 DoT (cutting experiment). B: FGF8 expression following the experimental separation of the caudal precursor zone into two. White arrows show how FGF8 is maintained at both the rostral and caudal moieties and black arrow shows the progressive down-regulation of FGF8 in the caudal moiety. C: Simulation of cutting experiment: the chemotactically moving DoT is cut into two pieces (see images at T=0 and T = 650). We allow the piece corresponding to the caudal moiety to move while the movement in the rostral moiety is arrested by the cut. The concentration of A-mRNA (shown by shades) which is associated with the location of the DoT reproduces the corresponding pattern for FGF8 shown in B. Parameters: $k_1 = 5 \cdot 10^{-4}$, $k_2 = 2.5 \cdot 10^{-4}$, $\beta = 1950$.

Using our complete model, we have performed simulations where we split the DoT into two (Movie S12) and follow the behavior of the two moieties. As shown in Figure 4-10C we find the maintenance of a caudal population of red cells (those producing A-mRNA or the equivalent FGF8-mRNA) in the rostral moiety and the progressive generation of a green population (that have stopped producing A-RNA) in the caudal moiety, very similar to what is observed in experiments (Figure 4-10A, B).

This result suggest that our model captures essential features of the biological network regulating FGF8 expression and encourage the search for a morphogen $A$ with both the ability to self-repress transcription of its encoding RNA and of FGF8. In addition, it posses the possibility that chemotaxis may play a role in caudal elongation of the embryo.
4.4 Discussion

The aim of this work is to explore possible mechanisms of progressive differentiation and regression of the caudal neural precursor region (CNPR, as defined by the region where cells transcribe FGF8 mRNA) in the chick embryo. We have focused on essential features of regression of this region such as progressive differentiation and the conservation of its size and speed of migration. These features were incorporated into two distinct modeling approaches which we used to evaluate a set of hypotheses concerning the mechanisms of differentiation and motion of cells in the CNPR. Our results are summarized in the Table S1.

The regulation of FGF8 dynamics during the regression of the caudal precursor zone was previously addressed using a mathematical model [48] where it was suggested that a mechanism involving FGF8 self-activation could, in theory, account for the progressive down-regulation of FGF8 provided a high FGF8 degradation rate was considered. According to this model the dynamics of FGF8 can be described as a propagating concentration wave, which is one of the patterns that form in nonlinear reaction-diffusion systems [58].

In the present work we have also addressed the regulation of FGF8 dynamics but we consider both intracellular (mRNA) and diffusing (protein) species of FGF8 and most importantly we take into account the movement of the domain of FGF8 transcription.

Cells in the precursor region proliferate and differentiate in such a way that the size of the region (identified by FGF8 transcription) does not change significantly over time. The size of a growing tissue usually involves control of proliferation such that when the tissue reaches a certain size, cells stop dividing. This may involve a mechanism which is able to measure population size as has been described in bacteria quorum sensing in Dictyostelium population, the Drosophila imaginal disc [59] or for the mesoderm community effect [60]. It is generally hypothesized that the concentration of signaling factors change as the size of the tissue increases until they reach a threshold value that dictates an arrest in cell proliferation. In our scenario, maintenance of a population with constant size is not due to an arrest of proliferation but to the balance of proliferation versus differentiation that is spatially controlled such that only cells at the rostral end of the domain differentiate (i.e. stop transcribing FGF8). This involves a mechanism where the strength of the signal regulating cell differentiation correlates with the size of the cell population, i.e. the signal is provided by a morphogen whose overall production is related to the size of the zone. In terms of our model this could be morphogen A produced from A-mRNA which in turn is produced exclusively in the precursor region. This is reflected by our model assumption that cell differentiation takes place when the level of morphogen A rises above some threshold, $T_A$. This assumption allows the control of the DoT size, although (depending on parameter values) the size can be stationary or oscillating (see Figure 4-3). An interesting problem is whether the DoT size is stationary or oscillating in experimental conditions.

Another important problem is what morphogen is actually under self-repression control and can be involved in the regulation of the DoT size. One possibility would be that FGF8 is actually able to repress transcription of FGF8 mRNA, however this is not supported by our experimental evidence as
manipulations of the level of FGFR activation in experimental conditions do not seem to affect the size of the area where FGF8 mRNA is expressed. This brings us to an alternative assumption that, for example, another morphogen is responsible for the regulation of FGF8 transcription. Two possibilities have been considered: morphogen $A$ activates FGF8 transcription or it represses FGF8 transcription (see Figure 5). Both are able to maintain a domain of FGF8 transcription of constant size; however the latter network would account more easily for the maintenance of FGF8 expression in the rostral fragment following the splitting of its domain of expression.

Several secreted proteins are present in the caudal zone that could correspond to $A$-mRNA such as WNTs (WNT3A, WNT8C) and BMPs (BMP7, BMP4). They could participate in the mechanisms presented in Figure 4-5. Independently of the particular mechanism that regulates production of FGF8 in our models, the relevant feature of the regulatory networks that allows the maintenance of a constant size of the domain transcribing $A$-mRNA is the presence of a negative feed-back loop involving protein $A$.

It is known that retinoid acid signalling from the somites is involved in down-regulation of FGF8: in the absence of RA the domain of FGF8 is expanded. However, down-regulation of FGF8 still occurs in RA-deficient embryos and our experiments of embryo sectioning show that progressive down-regulation can occur in the absence of rostral signals. In our model we did not take into account the influence of the rostro/caudal gradient of RA in shaping the FGF8 pattern. Future work will be required to incorporate into the models more elements concerning the gene regulatory network involved in FGF8 regulation such as the influence of RA, which is itself influenced by FGF signaling and Wnt8C, which is regulated by RA and FGF8 [7, 61].

Our models assume the existence of concentration thresholds of morphogen $A$ that determine whether $A$-RNA (or FGF8) is transcribed or not. Several molecular mechanisms underlying such an all-or-nothing response of cells could be relevant in this context, such as nonlinear saturating autocatalytic feedback of a gene product [2] or mutual inhibition [49]. It has been suggested that mutual inhibition of FGF8 and RA gradients may be involved in setting a bi-stability switch of FGFR versus Retinoic acid receptor activation. However so far, no experimental evidence indicates that such a switch could be involved in controlling whether FGF8 is transcribed or not [62].

Coordination of differentiation and axis extension can be found during growth of plant meristemes and in vertebrate limb bud development. In these cases, however, the mechanism involved must be different to caudal extension as differentiation coupled to axis extension relies on an external cell population that secretes a morphogen that regulates proliferation and maintains neighboring cells in an undifferentiated state. In the case of the root meristeme this is the quiescent center, in the case of the apical shoot meristeme it is the organizing center [33, 34] and in the case of the limb it is the apical ectodermal ridge that secretes FGFs [35].

The other feature that we have explored using our models is the mechanism of domain migration. Several cellular behaviors have been shown to contribute to regression of the primitive streak-node and extension of the embryo. Convergence (at the midline) and extension seem to be at play in mesoderm.
Besides, stem-cell like mode of growth and caudal movement of cells has also been observed in the neural tube and axial mesoderm [54, 55]. At the caudal neural plate, FGF signaling is required for cells to accompany the regressing primitive streak and precocious down-regulation of the pathway results in cells exiting the node-streak region. The version of the GGHM with differentiation (incorporating the influence of FGF8 on FGF8 transcription and cell motility) shows that such a mechanism is able to maintain a cohesive group of cells moving at constant speed (Figure 4-6). Further extension of the model with the assumption that the reason why cells move caudally is related to FGF8 concentration (FGF8 acts as a chemo-repellent) allows us to simulate the correct behaviour of cells that can move coherently in one direction provided there is an initial cause for the migration. A stationary group of cells producing a chemotactic agent maintains a symmetric condition with respect to the agent’s concentration profile and will not move unless other events (such as noise) are involved. Indeed, Hensen’s node (which we considered here as a part of stem zone) changes direction of its motion when the progression of primitive streak is replaced by its regression. We don’t know what is responsible for reversing the motion of Hensen’s node but most likely it is due to some external signals, while the repulsion by morphogen A in our model is rather an internal process as the production of this morphogen is closely associated with the processes in the caudal precursor region itself.

In summary, we have used mathematical models to explore possible mechanisms for the progressive differentiation of the caudal stem zone coordinated with the embryonic rostro-caudal extension. We have found that the self-repression of a caudal morphogen could be involved in driving progressive differentiation of the caudal stem zone and that chemo-repulsion here may be part of the mechanism responsible for the axis extension. Further experimental evidence is required to assess the role of FGF in regulating motility of ectodermal cells and to find out the signalling pathways that may be at the core of these mechanisms.
4.5 Materials and Methods

In this section we describe the mathematical models as well as the experimental techniques used for obtaining the results presented in this work. For our study we have developed two models: continuous (1D) and cell-based (2D, Glazier-Graner Hogeweg model also known as Cellular Potts model). Dynamics of morphogens was modelled in the same way in both models while the migration of the DoT - using different techniques. In the 2D model we have considered a tissue consisting of a single layered group of cells. Each cell can produce and/or degrade genes and proteins and, in addition, move in response to the forces (adhesive, chemotactic) acting upon it. Also, the 2D model incorporates the ability of cells to grow and proliferate.

4.5.1 One-dimensional continuous model

1D simulations were performed in a medium of fixed size in a frame of reference moving with the DoT. To describe the dynamics of morphogens we have used reaction-diffusion equations with an added advection term to take into account the DoT migration.

Basic Model.

The basic model is represented by two equations: one – for the dynamics of the concentration of a non-diffusible agent which we call A -mRNA and the second – for the concentration dynamics of corresponding protein A. The concentration of A -mRNA (denoted as \( u_1 \)) is equal to 1 (i.e. constant) inside the DoT of fixed size, a, while outside is given by the equation:

\[
\frac{\partial u_1}{\partial t} = c \frac{\partial u_1}{\partial x} - k_1 u_1
\]

Parameter \( k_1 \) defines the rate of A-mRNA decay while parameter \( c \) defines the speed of DoT migration or the speed of the frame of reference. The concentration of protein A (denoted as \( u_2 \)) is defined by the equation:

\[
\frac{\partial u_2}{\partial t} = D_2 \frac{\partial^2 u_2}{\partial x^2} + c \frac{\partial u_2}{\partial x} - k_2 u_2 + k_3 u_1
\]

where parameter \( D_2 \) defines its diffusion constant while \( k_2 \) and \( k_3 \) are the rates of protein decay and production. Production of the protein A is assumed to be proportional to the concentration of A-mRNA while its decay is proportional to its own concentration. The stationary solution of the system (4.2) can be found analytically. One can check directly that the solution:

\[
\begin{align*}
    u_1 &= \begin{cases}
        e^{-x} & \text{for } x \leq 0 \\
        1 & \text{for } 0 \leq x \leq a \\
        0 & \text{for } x > a
    \end{cases}
\end{align*}
\]
satisfies (4.1) while the solution:

\[ u_2 = \begin{cases} 
\frac{k_3}{D_2(\lambda_1 - \lambda_2)} \left( \frac{\lambda_2 - \lambda_1}{(\alpha - \lambda_2)(\alpha - \lambda_1)} \right) e^{\alpha x} + \left( \frac{1}{\alpha - \lambda_1} + \frac{1 - e^{-\lambda_1 x}}{\lambda_1} \right) e^{-\lambda_1 x} & , \quad x \leq 0 \\
\frac{k_3}{D_2(\lambda_1 - \lambda_2)} \left( \frac{e^{\lambda_2 x} - 1}{\alpha - \lambda_2} + \frac{1 - e^{\lambda_1 (x-a)}}{\lambda_1} \right) & , \quad 0 \leq x \leq a \\
\frac{k_3}{D_2(\lambda_1 - \lambda_2)} \left( \frac{1}{\alpha - \lambda_2} + \frac{1 - e^{\lambda_2 a}}{\lambda_2} \right) e^{\lambda_2 x} & , \quad x > a
\end{cases} \]

where

\[ \lambda_{1,2} = -\frac{c}{2D_2} \pm \sqrt{c^2 + 4k_2D_2} ; \quad \frac{k_1}{c} \] (4.5)
satisfies (4.2). Figure 2 shows typical concentration profiles given by ((4.3)–(4.4)).

Since the DoT is moving, the maximum of the concentration of protein A lags behind the middle of the DoT, i.e. \( x_{\text{max}} < 0.5a \) where \( x_{\text{max}} \) is the location of the maximum. For a slowly moving DoT the maximum is located inside the DoT (\( 0 < x_{\text{max}} < 0.5a \)) with its coordinate defined by the condition that the derivative of the \( u_2 \)-solution inside the DoT, \( 0 \leq x \leq a \) (see (4.4)), is zero. This gives:

\[ x_{\text{max}} = \frac{D_2}{\sqrt{c^2 + 4k_2D_2}} \left[ \lambda_1 a + \ln \left( 1 + \frac{\lambda_2}{\alpha - \lambda_2} \right) \right] \] (4.6)

his coordinate is \( a/2 \) when \( c = 0 \) and decreases with the increase of \( c \). When the DoT’s speed is too high the maximum lags behind the DoT, i.e. \( x_{\text{max}} < 0 \). The condition for this case can be given, for example, by the following inequality:

\[ \lambda_1 a = \ln \left( 1 - \frac{\lambda_1}{\alpha} \right) \] (4.7)

when \( x_{\text{max}} \) defined by (6) becomes negative. This condition is also confirmed by consideration of the maximum for \( u_2 \) solution behind the DoT (\( x \leq 0 \) in (4.4)).

**Model for the regulation of the DoT size.**

In order to consider the proliferation and differentiation of cells in the DoT we extend the basic model by the assumption that the location of the left side (or back side in respect to the direction of motion) of the DoT is controlled by the signal provided by protein \( A \). That is, the maintenance of \( A \)-mRNA, whose concentration is constant inside the DoT, is switched off (cells forming the DoT differentiate) when the concentration of protein \( A \) achieves the threshold value \( T_A \). In terms of the model ((4.1)- (4.2)) and its stationary solution ((4.3)–(4.4)) this gives:

\[ u_2(0) = \frac{k_3}{D_2(\lambda_1 - \lambda_2)} \left( \frac{1}{\alpha - \lambda_2} + \frac{1 - e^{-\lambda_1 a}}{\lambda_1} \right) = T_A \] (4.8)

and, therefore, the size of the DoT, \( a \), is not a preset parameter but a function of other model parameters, including \( T_A \):

\[ a = -\frac{1}{\lambda_1} \ln \left[ 1 - \lambda_1 \left( \frac{T_A D_2(\lambda_1 - \lambda_2)}{k_3} - \frac{1}{\alpha - \lambda_2} \right) \right] \] (4.9)
Furthermore, the threshold value, \( T_A \), is generally achieved in two points (on the either side of the maximum whose location is given by (4.6). As the concentration of \( u_2 \) should not get above \( T_A \) in the DoT the differentiation should take place before the maximum is achieved, i.e. condition (4.7) is to be held. Combining equation (4.9) with the inequality (4.7) we define the condition when the stationary solution to the stated problem exists:

\[
\frac{\alpha}{\alpha - \lambda_2} \geq 1 - \lambda_1 \left( \frac{T_A D_2 (\lambda_1 - \lambda_2)}{k_3} - \frac{1}{\alpha - \lambda_2} \right) \tag{4.10}
\]

An important case to consider is when the concentration of the protein A is low and doesn’t reach the threshold value, \( T_A \), anywhere in the medium. In the simulations we presume for this case that the size of the DoT is increasing over time (due to proliferation of cells) and the coordinate of the DoT's left border is gradually decreasing (the degree of “graduality” represents the proliferation rate). Simulations show that the size of the DoT is fixed and stable under the condition given by the equation (4.10). Furthermore, simulations show that if this condition does not hold the size of the DoT oscillates over time (see Figure 4-3 and Movies S2 and S3).

**For Variable Models.**

Modelled protein A down-regulates its own transcription while experimental results shown in Figure 4-4 indicates that FGF8 is not involved in the control of its own transcription. Thus protein A does not correspond to FGF8 and we need to analyze possible relationships between these two morphogens. We have examined two possibilities:

Transcription of FGF8 is proportional to the concentration of protein A (see Figure 4-5 A, B). This is expressed in the following equation for the concentration (\( u_3 \)) of FGF8 mRNA:

\[
\frac{\partial u_3}{\partial t} = c \frac{\partial u_3}{\partial x} + k_{31} u_2 + k_{32} u_3. \tag{4.11}
\]

Transcription of FGF8 mRNA and A-mRNA take place in the (nearly) same group of cells: they have been switched on independently from each other but both switched off by the signal provided by protein A. In this scenario the concentration of FGF8 mRNA is calculated the same way as the concentration of A-mRNA in the basic model (see above, equation (4.1)).

In both cases the concentration of FGF8 protein (\( u_4 \)) is given by the equation:

\[
\frac{\partial u_4}{\partial t} = D u_4 \frac{\partial^2 u_4}{\partial x^2} + c \frac{\partial u_4}{\partial x} + k_{41} u_2 + k_{42} u_4. \tag{4.12}
\]

i.e. similarly to the concentration of protein A, \( u_2 \), (see equation (4.2)) it is a diffusible agent and its production is proportional to the level of its corresponding gene (FGF8-mRNA, equation (4.11)) and decays proportional to its own concentration.

**Modeling chemotaxis.**

In this version of the model, parameter \( c \), defining the DoT migration speed, is calculated with the assumption that the migration is taking place due to chemotaxis, i.e. the speed is proportional to the
gradient of the chemotactic agent \[63, 19\]. We have assumed that protein \( A \) acts as a chemo-repellent on cells forming the DoT and the speed of migration is defined either by its gradient in some specific point, say on the front (right-side, \( x = a \)) of the DoT:

\[
c = -c_0 \frac{\partial u_2}{\partial x} \bigg|_{x=a},
\]  

(4.13)

(see Movie S6) or by the average gradient over the DoT,

\[
c = -c_0 \frac{u_2(a) - u_2(0)}{a},
\]  

(4.14)

(see Movie S7).

The analysis of conditions when the DoT can migrate due to self-repulsion is relatively simple when we consider the chemotactic movement of a DoT of fixed size, \( a \), i.e. consider solution ((4.3)–(4.4)) remove condition (4.8) and add condition (4.14) which gives:

\[
c = \frac{k_3 c_0}{D_2 (\lambda_1 - \lambda_2)} \left( \frac{a}{(\alpha - \lambda_2)\lambda_2} \left(1 - e^{\lambda_2 a}\right) + \frac{1}{\lambda_1} \left(1 - e^{-\lambda_1 a}\right) \right)
\]  

(4.15)

Where the right hand side is also function of \( c \) (\( \lambda_1, \lambda_2 \) and \( a \) are functions of \( c \), see the definitions given by (4.5)). When \( c = 0 \) the right hand side of (4.15) is zero, i.e. one stationary solution (with \( c = 0 \)) exists for all sets of parameters. One can show that the right hand side of (4.15) is positive and tends to zero when \( c \) tends to infinity. Travelling solutions correspond to the points where \( c \neq 0 \) and plots of functions

\[
y_1 = c \quad \text{and} \quad y_2 = \frac{k_3 c_0}{D_2 (\lambda_1 - \lambda_2)} \left( \frac{a}{(\alpha - \lambda_2)\lambda_2} \left(1 - e^{\lambda_2 a}\right) + \frac{1}{\lambda_1} \left(1 - e^{-\lambda_1 a}\right) \right)
\]  

(4.16)

intersect. At least one such point exists, if the derivative of the function \( y_2 \) (the derivative of the RHS of (4.15)) is more than 1 at \( c = 0 \). This condition can be expressed by the formula:

\[
\frac{k_3 c_0}{2ak_1 \sqrt{k_2 D_2}} \left(1 + \frac{k_1}{k_2} - \left(1 + \frac{k_1}{k_2} + \frac{k_1}{\sqrt{k_2 D_2}}\right)e^{-a \sqrt{k_2 D_2}}\right) \geq 1
\]  

(4.17)

Therefore for sets of model parameters satisfying (4.16) (see Figure 8A) we can expect the existence of a moving DoT, moving with constant speed. Whether more than one such solution exists and whether such solutions exist when condition (4.16) is violated should be rigorously analyzed in a more detailed study. We have plotted the function given by the RHS of (4.15) versus variable \( c \) for various sets of model parameters. It looks that this function always has only one maximum. Therefore we expect that inequality (4.16) gives the condition for the Pitch-Fork bifurcation, i.e. we have only one solution (corresponding to \( c = 0 \)) when model parameters do not satisfy (4.16) and two extra solutions appear (corresponding to the DoTs moving in opposite directions) when (4.16) is satisfied. But indeed this conclusion should be verified by proper analysis.
Details of simulations and verification of parameters.

For simulations we used the explicit Euler’s method with central differencing scheme for diffusion and alternating up- and down-wind schemes for advection. Typical initial conditions: all concentrations are equal to zero with the only exception: \( u_1 = 1 \) in the DoT which has a predefined location and size. Default values of the parameters: diffusion coefficients \( D_2 = D_4 = 0.5 \); kinetics rates \( k_1 = k_{31} = k_{41} = 0.0003, k_2 = 0.00025, k_3 = 2k_2, k_{32} = k_{42} = 2k_{31}; \) speed \( c = 0.015 \), chemotaxis \( c_0 = 2 \).

Default values for the time and space steps \( \Delta t = 1 \) and \( \Delta x = 1 \) for which we found the simulations to be fairly accurate: two-fold reduction of space step together with four-fold reduction of time step \( (\Delta t = 0.25 \) and \( \Delta x = 0.5) \) was altering measured quantities (such as maximums in concentration profiles, the DoT size when differentiation is on, and the DoT speed when it is drove by chemotaxis) by less than 3%. Also the simulations were performed in sufficiently large domain to reduce the influence of medium boundaries (doubling the size of the medium has changed measured quantities by less than 1%).

To scale the model parameters we estimate the DoT size to be \( 1 \) mm and its speed to be \( 0.1 \) mm/hour. Comparing this with the simulations shown on Figure 4-3 where \( c = 0.015 \) (space units over time units) and the DoT size is 40 (space units) we conclude that the space unit corresponds to about 20 \( \mu \)m and the time unit – to 10 seconds. This means that \( D = 0.5 \) corresponds to \( 2 \cdot 10^{-1} \text{sm}^2/\text{sec} \) and kinetics coefficient \( k = 0.0003 \) – to \( 3 \cdot 10^{-2} \) sec.

A few words about justification of the parameter values used in our simulations. Firstly, the analysis of the model represented by equations (4.1) and (4.2) with extra conditions (4.8) and/or (4.14) indicates that qualitatively the solution is the same for any set of parameters represented by positive numbers. Furthermore, we can take three arbitrary numbers to represent the values of three parameters appropriate for scaling dimensions associated with time, distance and concentration. In our case we decided that the concentration of A-mRNA inside the DoT is 1, the DoT is represented by about 40 grid points (or its size is 40 space units if the grid size is 1) and the speed of the DoT is something between 0.1 and 0.01. The choices for the DoT size and speed are dictated by the accuracy issue. We have checked that 40 grids for the DoT gave considerably more accurate solutions than say 10 grids and, on the other hand, approximately the same accuracy as 100 grids. Similarly, if we assume that the time step is 1 then speed \( c \) should be less than 0.1 (say 1/40) to provide enough accuracy in numerical calculation of concentration profiles. Diffusion \( D = 0.5 \) is convenient when it comes to the numerical scheme (the highest possible value when the explicit Euler scheme with time and space steps \( \Delta t = \Delta x = 1 \) is still stable) and still in a range of diffusion coefficients known for proteins. Kinetic rate \( k_1 \) has been chosen in a way that the space scale for the mRNA degradation is comparable with the size of the DoT: this is done to fit with the observations concerning the sizes of \( FGFB \)-mRNA transcription and expression (Figure 4-1 and Figure 4-4). Other kinetic constants have been chosen to be of the same order as \( k_1 \). And finally, concerning \( k_2 \) and \( k_3 \): the ratio of these two constants is only important for the choice of the threshold value \( T_A \): the ratio
2 has been chosen only to bring concentrations of $A$-mRNA and protein $A$ to the same scale (Figure 4-2, Figure 4-3, Figure 4-5).

**Glazier-Graner-Hogeweg Model.**

This is a computational individual-based model originally developed by Graner and Glazier [20, 30]. In this model we consider the DoT as a group (25 by default) of cells, each, in turn, is represented by a number of grid points (50 grid points per cell in our simulations) on a regular (square-shaped 2D in our case) lattice (see also Methods Section in [4]). Movement of a cell (or change in its shape) means that the cell looses or gains some grid points on the lattice. In terms of the underlying tissue this implies that the grid points are associated with different cells at different times. To calculate whether a particular grid point will be associated with a different cell at next time step a variation principle is used to minimize a quantity representing “the energy” of the system.

Contrary to the original implementation of the GGHM which was based on Monte Carlo algorithm involving the random choice of the pixel followed by the random choice of its neighbor and following calculations of probability of change [20] we have implemented a synchronous model: at each time step we calculate the probability to change the state for all grid points. For each grid point, we randomly select a neighbor (one out of the eight nearest) and calculate how the energy of the system will change after changing the state of the grid point to that of its neighbor. If this change results in an energy decrease we allow the change to occur; if the energy is increased we calculate the probability of that change, $p$, using the Boltzmann function: $p = \exp(-\Delta E/T)$ where the parameter $T$ can be referred to as the “temperature” of the system.

The energy is defined in a way that its change accounts for the work done by different forces acting upon moving or deforming cells. The definition of energy used in our implementation of the model takes into account three forces, the adhesive forces between cells, the force associated with the incompressibility of cells (pressure) and forces developed by chemotactically moving cells:

$$E = E_{\text{adhesive}} + E_{\text{pressure}} + E_{\text{chemotaxis}}$$  \hspace{1cm} (4.18)

The following definitions of the terms on the right hand side of equation (4.17) are commonly used in various modifications of GGHM [4, 50]:

An adhesive energy associated with cell-to-cell contacts is defined by the adhesion matrix $J_{k,l}$ ($J_{k,l} = J_{l,k}$) which refers to an interface between neighbouring grid points which belong to different cells (numbers $k$ and $l$ represent cell types of these cells). The energy, $J_{k,l}$, characterizes the strength of a particular cell’s adhesive contacts (stronger contacts correspond to smaller energies). To consider adhesive contacts between cells and the surroundings we treat the latter as a special cell of its own type. In our simulations we, as a rule, consider 3 cell types: the surrounding was considered as a cell of its own type – cell type 1; cell type 2 – cells which form the DoT; cell type 3 – cells which form the DoT trail or the differentiated daughter cells. The default adhesion matrix for adhesive bonds between each pair of different cell types is:
To control the size of a cell (say $k^{th}$ cell), $A_k(t)$, a target area (in case of our 2D cells number of grid points forming the cell can be considered as its area), $T_k$, is introduced. The $k^{th}$ cell is given an energy $E_{vol,k} = a(A_k(t) - T_k)^2$, where $a$ is a positive constant ($a = 0.6$ in all our simulations). Constant $a$ represents the cell’s resistance to compression and we can call it the “incompressibility” coefficient (it had different names such as “Lagrange multiplier specifying the strength of the area constraint” in [20], or “volume elasticity” in [64]) referring to another approach when the motion of cells in a tissue is seen as a flow in incompressible viscous liquid described by Navier-Stokes equation [65, 66] and where this term would correspond to the gradient of pressure. The “incompressibility” energy reaches its minimum (zero) when the cell’s actual and target areas are equal. To take into account the growth of cells the target area, $T_k$, is considered to be an increasing function of time. To model cell proliferation we split big cells (which contain 100 or more grid points) into two small cells. The split is performed along a straight line (having a random orientation) crossing the cell’s centre of mass.

To implement the chemotactic effect of the agent, whose concentration is denoted as ‘$u$’, to a moving (or deforming) cell we introduce the change in chemotactic energy $\Delta E_{ch,k} = \beta_k (x = \text{grad}(u))$, where $\beta_k$ is a constant describing the chemotactic response of cells of type $k$ to the chemotactic agent $u$ and $x$ is a vector representing the local displacement of the given cell’s boundary. This energy change refers to the work done by chemotactic force $\Delta F_{ch,k} = F_{ch,k} \cdot x$ and therefore corresponds to the chemotactic force, $F_{ch,k} = \beta_k \text{grad}(u)$, exerted by chemotactically responding cell. The identical definition of the chemotactic force was introduced and used earlier in the hydrodynamic model of Dictyostelium development [67]. The most common implementation of chemotaxis in our model: $\beta_k = 0$ if $k \neq 2$ and $\beta_k = \beta$ ($\beta \neq 0$) for $k = 2$ or for “red” cells forming the DoT. This means that there is an energy gain or loss in the system related to the relocation of the red cell’s boundary which depends on the local gradient of chemotactic agent. We note that a positive value for the parameter $\beta$ corresponds to the process of chemo-repulsion while a negative value for $\beta$- to chemo-attraction.

Detailed description of the GGHM model and its modifications and applications to various problems in developmental biology are given in [50]. One of the greatest advantages of GGHM is that it allows modeling the dynamics of biological tissue while being focused on behavior of individual cells. The simplicity of the model allows modeling of tissue which contains up to 105 cells on a single PC. Parallel implementation of the software [64] allows an increase in this number up to 107–108 which is close to the actual number of cells in many real tissues. Furthermore, the GGHM allows relatively simple modifications to address various problems associated with mechanics and deformations of cells. For example the GGHM allows consideration of cells of different shapes. Cells in the version of the GGHM which we use here are predominantly round-shaped. To model, for example, elongated cells the GGHM can be extended by the introduction of the anisotropy in adhesive properties of cells [68] or by the
introduction of cellular subunits which compose cells of desired shape and stiffness [69]. The GGHM has also been extended to address three-dimensional problems [70] and its simulation code is available publically (the CompuCell3D package at http://www.compucell3d.org).

**Details of simulations.**

We have implemented a synchronous model: at each time step we calculate the probability to change the state for all grid points. For each grid point, we randomly select a neighbour (one out of the eight nearest) and calculate how the energy of the system will change after changing the state of the grid point to that of its neighbour. If this change results in an energy decrease we allow the change to occur; if the energy is increased we calculate the probability of that change, \( p \), using the Boltzmann function:

\[
p = \exp(-\Delta E/T)
\]

where the parameter \( T \) can be referred to as the “temperature” of the system (we set \( T = 6 \) in all our simulations).

All simulations start with a group of 25 cells (forming an artificial square-shaped tissue) representing the DoT. In the simulations where we do not consider cell proliferation, we assume that all cells have a constant target volume (\( T_k = 50, \forall k \)) which does not depend on the cell age. In the simulations where we take into account cell growth and proliferation, we assume that the initial target volumes of cells are randomly distributed among the cells in the range (30–70) and then the target volume of each cell is increased by one unit every 10 time steps with probability 1/3. When the actual volume of a cell reaches 100, the cell divides along a line crossing through the cell’s centre of mass with a random direction of the cleavage plane. After division the target volumes of both daughter cells are reset to 50 and they start to increase again over time. This implies that the average time required for a cell to double in size and proliferate is equal to 1500 time steps. One time step scales as 70 seconds (as it derived in the next section) and therefore the effective proliferation rate in the model is one division per 30 hours. In experimental conditions the proliferation rate is much higher (one division per 6 hours) but on the other hand, in experimental conditions, many cells leave the stem zone (and epiblast) and transform into mesenchyme cells. Since in our model we don’t consider formation of mesenchyme cells (this would require three-dimensional version of the model) we have to reduce the proliferation rate of cells in the epiblast (to compensate the mesenchyme formation). Furthermore we did not consider the proliferation of differentiated (green) daughter cells as this would not influence the phenomena which we are interested in but add unnecessary details into simulations and graphical outputs used in the figures.

**Verification of parameter values in GGHM.**

Parameters used in the GGHM can be split into two sets. One set is used for the definition of energy in the system and is associated with adhesiveness (entries \( J_{k,l} \) in the adhesion matrix), incompressibility (parameter \( a \) defining incompressibility) and chemotactic responses of cells (parameter \( b \) defining chemotactic response) as well as temperature \( T \) in Boltzmann function. The second set of parameters is used to define the dynamics of morphogen concentrations (kinetics and diffusion of morphogens). The First set of parameters forms a core of the GGHM and verification of the parameter
values used for this set can be found in the literature ([30, 50] including more references in [50]). Here we can briefly note that the most important point concerning the entries \( J_{k,l} \) in the adhesion matrix is their ratios: \( J_{k,l} = J_{l,k} > 2J_{k,k} \) for cells to stay together and form a tissue. Also \( J_{k,l} = J_{l,k} > 2J_{l,l} \) for cells of types \( k \) and \( l \) to sort out or to stay sorted out. The values of the entries \( J_{k,l} \) are scaled with the values of parameters \( a \) and \( \beta \) in order to scale all three considered forces (associated with adhesion, pressure and chemotaxis) relative to each other. The value of parameter \( T \) defining the rate of the evolution in the system is also scaled with the values of \( J_{k,l} \), \( a \) and \( \beta \). The ratio \( a/T \) defines the amplitude of the cell shape fluctuations (or cell membrane fluctuations). These fluctuations freeze at high values of \( a \) as well as at low values of the Boltzmann temperature \( T \). If we will keep all parameters of the model constant and vary only the temperature we will see that the rate of dynamics in the model will be low at low temperatures, then the processes accelerate with the increase of the temperature and eventually they slow down again when the temperature becomes too high. We have measured the speed of migrating group of cells as a function of the Boltzmann temperature (keeping all other model parameters at their “default” values) and found that the highest speed is observed at \( T = 6 \) (see Figure S1). It was noted in [71] that the Boltzmann temperature, \( T \), defines the intrinsic cell motility in GGHM. Therefore \( T = 6 \) (which we have chosen for our simulations) corresponds to the highest possible intrinsic cell motility for the given set of other model parameter values.

It was shown on many occasions that the GGHM is robust: small variations in the values of model parameters do not alter qualitatively the outcome of simulations. Besides, it was shown that the GGHM parameters can be rescaled so that the outcome of simulations is absolutely the same. For example, the simulation of the primitive streak progression was performed in [4] in tissues containing 625 and 15000 cells without any notable difference in the outcome.

The concentration fields of morphogens were calculated in a way similar to that for the 1D model. The level of \( A \)-mRNA was set equal to 1 in all (red) cells forming the DoT and was decaying in differentiated (green) cells according to the equation \( \dot{u}_1 = -k_1 u_1 \) similar to what was in the 1D (compare with the equation 1 where \( c = 0 \)). The equation for protein \( A \) includes diffusion, production and decay and is given by the equation \( \dot{u}_2 \) (see above) where \( c = 0 \). There are no advection terms in the 2D model as the events are considered in the laboratory frame of reference. As the GGHM is considerably slower (as compared with our 1D model) we have increased the speed of computations by ensuring slightly faster processes (faster moving DoT and faster kinetics for chemicals). For \( k_1 = 0.0001 \), \( k_2 = 0.0003 \) and \( \beta = 4.5 \) (see Figure 4-6) the speed of the DoT is roughly 60 space steps per 1000 time steps (should correspond to 0.1 mm/hour) and the DoT size is roughly 32 grid points (should correspond to 1 mm). This means that 1 space unit roughly corresponds to 30 \( \mu \)m, 1 time step to 70 seconds causing for dimensional diffusion and kinetic coefficients to be slightly (2 to 3 times) less than for the set of parameters used in the 1D model.
Experimental Methods.

Stage Hamburger and Hamilton (HH) 9–10 chick embryos were obtained from fertilized eggs (Granja Santa Isabel, Cordoba, Spain) and dissected in L15 culture medium (Invitrogen).

Embryos were cultured in 4 well dishes on top of collagen beds and with 0.2 ml of culture medium (OptiMEM (Invitrogen), fetal calf serum, glutamax and gentamicine) containing 0.1% DMSO (control) or PD173074 (10 μM in 0.1% DMSO, Sigma). Caudal explants (including 3 embryonic layers) were cultured in collagen as described in [72] in the presence of BSA (control) or hFGF4 (330 ng/ml, Sigma). For splitting the caudal domain into two, embryos were prepared following the EC culture method [73], a cut was performed caudal to the node with a microsurgical knife and embryos where cultured for another 20h. Embryos and explants were fixed in 4% PFA and processed for in situ hybridization with probes to detect either nascent [6] or total FGF8 following standard methods.

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Author Contributions.

Conceived and designed the experiments: RDC BV. Performed the experiments: NCH RDC BV. Analyzed the data: NCH RDC BV. Contributed reagents/materials/analysis tools: RDC BV. Wrote the paper: NCH RDC BV.
Chapter 5

DISCUSSION AND CONCLUSIONS

In this thesis we have set out to understand (and explain) mechanisms of self-regulated motion of a group of cells due to reaction of the group to the gradient of locally produced chemical, referred to as chemotaxis. Such problems are abound in nature with examples appearing in both plant and animal kingdoms, illustrating the ubiquitous nature of chemotaxis as a mechanism for migration and reorganisation in biological cellular systems. However while vast amounts of data has been collected over protracted period of time, it is only relatively recently that computational power and techniques have advanced to the state where these data can be analysed and correlated. Further, advances in microscopy and genetic sequencing to show how the expression of genes controls the development of biological organisms at the cellular level, that bridge the gap between the microscopic and macroscopic, that is, gene expression and cell behaviour. It is in this context, molecular and cellular dynamics, that this thesis attempts to explain self-regulation of migratory behaviour (cellular dynamics), due to local gene expression (molecular dynamics).

For the rest of this chapter we shall briefly summarize the preceding work, and consider how what steps we can take next to further the work, and end with and short conclusion.

5.1 Summary Of Thesis

In the most basic setting, we consider a small group of cell that is chemotactically reacting to a locally produced chemical. This chemical in a biological context, is the product of a gene expression within some population of cells, translating into an externally diffusing protein. The diffusive nature gives rise to a spatial distribution, that in a stationary setting can visualised as the normalised probability curve (the bell curve). If our group of cells are exposed to such a curve then depending on how they reacted to the gradient of this protein, they could either move towards (chemo-attraction) or away (chemo-repulsion) from the point of highest concentration. This raises two fundamental questions in the thesis: who is producing the chemical and who is reacting to it? And under what conditions will the group move, or will not as the case may be?

In chapter 2 we began with a reduced one-dimensional analytical model, essentially a caricature representation of the group of cells, as a segment of the real line. This allowed us to neglect the physiological aspects of the cells in the group in order analyse the chemical and chemotactic dynamics in a simplified setting. Using this simplified model we demonstrated a variety of mechanisms for group migration, for different assumptions about where the chemical was produced and what cells reacting to it. The results of this chapter where that there are essentially two dynamical systems that can result, that depend on the composition of the group being homogeneous or heterogeneous. From these dynamical systems we were able to derive analytical expressions/conditions that result in travelling solutions, that
were shown to be dependent on chemotactic forcing parameter $c_0$. Thus we were able to derive the fundamental characteristics of group motion.

In chapter 3 we extended our model to two-dimensions, with a numerical analytical model where we represented the group as a circle (or disk), and found there are fundamental differences between one and two-dimensional representations due to how the geometry of the group affects the diffusive characteristics of the chemotactic agent. Regardless of this difference we demonstrated a strong qualitative and quantitative agreement between these models. With this in hand we then showed, using a modified Cellular Potts Model, that when we introduce a more realistic morphology, and indeed more realistic physiology, we can demonstrated all of the results of the analytical/numerical model, essentially corroborating the robustness of the self-regulatory mechanism of chemotaxis.

In the final chapter 4 we performed an extensive experimental study that brings together the analytical, numerical and computational techniques illustrated in the previous chapters, applied to a study of early growth in the central nervous system in the chick egg. In this work [11] we demonstrated how coordination of proliferation, differentiation and regulation of morphogens bringing about self-regulated migratory behaviour observed in axial extension of the central nervous system, during neurulation. While there are still questions relating to identity of specific morphogen responsible processes, the underlying mechanisms of self-regulated chemotaxis is elegant and relies on few assumptions, specifically those outlined in Chapters 2 & 3.

5.2 Discussion

In general the mechanisms underlying biological migratory behavior or spatial reorientation of an object, can be a function of several different mechanisms that are dependent on numerous factors such as: environment, motivation, physiology and stimulus, to name but a few. Clearly the specification of these factors for a given object is governed by its size and/or complexity. For example if we suggest the object in question is a hungry lion, we could prescribe a set of values for the above factors with ease. However if we suggest that object is the organizing centre Hensen’s node in gastrulation in the chick embryo, few could, with certainty, prescribe values for all of the above factors. Hence we are forced to take a reductionist view and consider the problem in terms of one or two varying factors, while holding all the others constant if you will.

In this context of this thesis we only consider stimulus, that is, what signal compels the object to migrate, and physiology, how the object responds to the signal. Hopefully by now it is clear that by stimulus we mean the presence of diffusing chemical, and by physiology, we mean chemotactic movement.

The choice of chemotaxis was not taken arbitrarily, indeed the literature is abound with it as a fundamental mechanism in many systems of growth, development and survival throughout the plant and animal kingdoms. Mathematical models of chemotaxis tend to focus on an arbitrary point source [56], while mathematical biological models focus on a select set of model organisms such a bacterium [63] and ameobae [65, 67, 19], due their relative simplicity and having their behavior macroscopically visible to the
naked eye. In our thesis however, our experimental work is in the very early stages of embryogenesis, where there are literally innumerable regulatory networks of chemicals involved, not to mention the physiological aspect of motility itself in such a compact cellular medium [9], that are not so easily discernible by macroscopic observation. However there is evidence in the literature that suggests there are certain morphogens [5, 7] that are coupled with the ability of cell population to move. Specifically we consider extension of the primitive streak as coordinated/orchestrated by Hensen’s node, a small compact group composed of approximately 20,000 cells that appears to maintain a consistent size while migrating uniformly during primary body axis extension. Given this it is reasonable to assume that the primary mechanism involved in migration of the node is chemotaxis, however there are question relating to other local and global forces due to passive and active cellular flows [9] that could also have a dramatic impact on migration, however we largely neglect these in this thesis. In addition it not clear exactly which chemical/morphogen is responsible directly or indirectly for node migration [11], however our models were designed such that, were it to be found, it would not change the outcomes of our results.

When we considered the composition of the group in our models, that is the model representation of Hensen’s, we assumed it was either homogeneous or heterogeneous, in that was either a homotypic or heterotypic population of cells. In a sense we can consider the homogeneity as a base case in our analysis, that is, the simplest assumption we can make is that the node is composed of single type of cell. In this case we showed using various mechanisms of chemo-attraction and chemo-repulsion, with itself and a surrounding population, that there are symmetries arising between the group and an external population that lead to pairs of travelling solutions, however we also showed that there are symmetric cases that do not, the results of which we correlated between analytical (1D model) and numerical methods (CPM).

When we consider heterogeneity, we found that there was no configuration for which we could not find traveling solutions, and in the sense the results of these simulations were rather uninteresting. However when initially investigating cellular models we reproduced results [4] where heterogeneity plays a crucial role in the dynamics of progression of the primitive streak; a preceding event to regression we studied in Chapter 4.

During the early stages of development of the chick embryo a small condensation of cells arises in the posterior marginal zone of the embryo within a region known as Koller’s sickle (Figure 5-1). This small condensation is the first emergence of Hensen’s node and it progresses over the surface of the embryo, and as it does so Koller’s sickle follows and forms a streak behind the node. The dynamics of this process are assumed to be a function of both passive and active cell motility, where passive is due to cell flows that occur in the other cell population (such as Koller’s sickle) and active chemotactic force acting on the node. In this setting the heterogeneity of the simulations seems to be a determining factor for successful streak progression, and therefore its role in embryogenesis could be an important mechanism underlying primitive streak regression.
Figure 5-1: Primitive streak progression during the early stages of embryogenesis may rely on heterogeneity between cell populations in the embryo in determining successful streak progression. A: Initial conditions for simulations with Hensen’s node (yellow cells) located with Koller’s sickly (blue cells) are opaca (green cells) and area pullicada (red cells) and black and white cells above where chemical dynamics can be observed. B: Hensen’s node diffuses a morphogen that acts to attract Koller’s sickly cells. C: Hensen’s node difuses morphogen that repels area pellucida cells. D: Area pellucida difuses chemical that attracts the cells in Hensen’s node. E: Keller’s sickle diffuses morphogen that repel cells in Hensen’s node.

In addition to these simulations we also undertook a minor study with Prof Weijer lab in Dundee to produce simulations that could corroborate analysis that he had found in investigating a morphogen suspected (FGF8) of playing a role in chemotaxis during streak progression (Figure 5-2). The configuration of the experiment was such that, just prior to the progression of the streak, a bead saturated with the morphogen was implanted into the embryo lateral to the streaks normal progression, to determine its effect on the streak. The results showed clearly that the expected behavior of the morphogen on the streak was contrary to its previously known effects, that is, it was previously assumed that it acts as a chemo-repellent. However it clearly demonstrates that cells in the streak where chemo-attracted to the bead, illustrating that if it didn’t have a differential effect then we would not have observed any reaction.

Figure 5-2: Bead simulations illustrating differential FGF8 signalling of motility. Experiments were conducted to determine the effect of FGF8 on cell motility in cell populations on the surface of the embryo where a bead saturated in FGF8 was placed adjacent to the primitive streak. The streak was attracted to the bead (A,B,C) indicating that FGF8 seems to act as an attractor which is counter to FGF8s known effect as an repellent, and secondly that FGF8 must have a differential effect on cell populations otherwise the bead would have a negligible effect. Simulations were conducted to understand the mechanisms involved where different scenarios of attraction and repulsion where investigated. D: Initial conditions. E: Yellow attracts blue and grey repel yellow. F: Yellow repels red and grey repel yellow. G: Blue and Grey repels yellow.
The discussion wouldn’t be complete without reference to competing models, or at least models by their definition lend themselves to problems of the kind we have described in Chapter 4. In this sense there are two models that are generally accepted as being involved in the emergence of spatially complex structures and patterns in biological organisms. The first of these is based upon the work of Alan Turing [16] in the so-called reaction-diffusion systems, where it has been shown that in a two-component (and possible more) activator-inhibitor system complex stationary patterns can emerge between interacting morphogen gradients as a result of growth within tissue [2, 74, 49, 35], that stimulates instabilities as the concentrations of the morphogens rise.

The reaction begins when the concentration of a morphogen $A$ begins to increase at some point, and as it does so it catalyses the production of a secondary morphogen $B$, which in turn acts to down-regulate $A$. If $B$ diffuses at a faster rate than $A$ then locally the concentration of $B$ will be less than that of $A$ and thus its ability to down-regulate $A$ locally will be small. However at a longer range the concentration of $B$ will exceed that of $A$ and thus will act to totally down-regulate down $A$; put another way we will have short range activation and long range inhibition. At furthermore range $B$ degrades in the absence of $A$ and in this way new instabilities may arise by the same mechanism outlined above. In such a context it easy to see how different spatial patterns can arise, such as spots or stripes, dependent of diffusion and kinetics rates of the morphogens.

The second model assumes that there are propagating waves [48] that develop mutual interactions, whereby both morphogens catalyse the decay and production of each other while still enhancing their own production by autocatalysis. The interaction of these morphogens can give rise to spatially stable travelling wave-fronts (not to be confused with Turing waves) in the concentrations of one or more morphogens, such that they travel along a spatial axis.

In both of these models it is assumed that the morphogens involved illicit programs of differentiation in the cell populations that are exposed to them. In a stationary setting of the first case, it has been shown in many development contexts such as the patterning on the bodies of zebra fish, correspond to expressions of morphogen concentrations defining skin pigmentation. In the travelling setting of the second case wave-fronts can describe a similar program of cellular differentiation due to thresholding as the wave-fronts travel over a development field (populations of cells exposed to the waves), similar to that described in positional information [15], albeit in a non-stationary setting.

The latter mechanism of travelling wave-fronts would seem to be more relevant to the models we propose in this thesis, since it is not necessarily clear how stationary morphogen concentrations can lead to migratory behavior. Bringing things back to the setting of primitive streak regression of Chapter 4 travelling wave-fronts could in theory be responsible for migratory behavior of the node, in that the wave-front could act as a chemo-attraction or chemo–repellent depending on the directionality of the waves propagation. In this sense the nodes migration is an active response the chemicals gradient of the wave, that in simple terms literally pushes (or drags) the node along in front of the wave, analogous to a surfer.
on an ocean wave. This is in direction contrast to our models where we assume that it is the motion of the node that brings about the wave, initiating a travelling solution.

5.3 Conclusion

In this thesis we derived a novel caricature one-dimensional model for the analysis of chemical and chemotactic dynamics of an abstract representation of a group of cells (Hensen’s node). We demonstrated that heterogeneity and homogeneity play a key role in what type of motile dynamics occurs, in that the composition of the group, and the resultant chemotactic mechanism, produces two types of non-linear dynamical systems (pitchfork and fold bifurcations). We showed directly that solutions to these systems depend on exact conditions that determine whether the segment will move, that is travelling solutions emerge, under the assumption that such solutions manifest as travelling waves and whether these waves remained stable. That is, once travelling (wave) solution emerged, the profile of the wave would remain constant in time.

To corroborate this analysis we investigated a cell-centered computational model (Cellular Potts Model) that takes into account a physiology and morphology of cells in more realistic manner. Using this model we encountered several phenomenological features intrinsic to two-dimensional modeling not present in our one-dimensional model, however we were able to account for these features and demonstrated conditions that could mitigate these differences. Further we were able to demonstrate all that had been demonstrated in our one-dimensional analysis, and found that other mechanisms or travelling solutions can occur. However while it evident that the CPM based modeling corroborates our 1D analysis, it is also true that there remains a question mark over the CPMs, as there has been no formal analysis of its parameter space (by ourselves or others) and therefore, at best, it can only be considered a qualitative model.

Bringing together the analysis of both one and two-dimensional modeling, we applied what we had learnt to an experimental setting of migration of Hensen’s node and regression of the primitive streak during embryogenesis in the chick embryo. Through unique experiments, performed by Dr. R.D. del-Corral, we correlated our numerical and analytical results with experimental observations and demonstrated, that there is indeed good agreement in terms of morphogen dynamics and nodal migratory behavior. These experiments also concluded that a specific morphogen that has been identified with regulating migratory behavior of cells during gastrulation (FGF8), a morphogen we assumed could be responsible for nodal/streak regression, did not correlate with experimental evidence. However our models were made sufficiently general that if such a morphogen were to be found, we could incorporate it into our models with equivalent results. Notwithstanding this small contradiction, our results clearly demonstrated an elegant biophysical model that requires few assumptions, and produced robust reproducible results.
Appendix A

**DERIVATION OF SOLUTION FOR A HOMOGENOUS GROUP WITH AN INTERNALLY PRODUCED CHEMOTACTIC AGENT.**

We start by considering the production of a chemical over the spatial domain as a reaction-diffusion system of equations as defined in the manuscript as (2.3) and (2.4), repeated here for convenience:

\[
D \frac{d^2 u(\xi)}{d\xi^2} + c \frac{du(\xi)}{d\xi} - k_1 u(\xi) + p(\xi) = 0, \\
\]

where \( \xi = x - ct \) when we converted to a co-moving frame of reference. If we consider the system for a group with an internally produced chemical as given in (A.2), where the partitions of the spatial axis are given by the arbitrary constants \( \alpha \) and \( \beta \):

\[
\begin{align*}
0 &= D\Delta u + cu_x - k_1 u, & x < \alpha \\
0 &= D\Delta u + cu_x - k_1 u + k_2, & \alpha < x < \beta \\
0 &= D\Delta u + cu_x - k_1 u, & x > \beta
\end{align*}
\]  

then solutions are given by solving each equation independently and then employing boundary conditions to solve for the unknown coefficients. Let us refer to each equation by the subscript \( i \in \{1, 2, 3\} \), where each index corresponds to the intervals defined left to right on the \( x \)-axis thus giving three equations:

\[
\begin{align*}
\dot{u}_1 &= D\Delta u + cu_x - k_1 u, & x < \alpha \\
\dot{u}_2 &= D\Delta u + cu_x - k_1 u + k_2, & \alpha < x < \beta \\
\dot{u}_3 &= D\Delta u + cu_x - k_1 u, & x > \beta
\end{align*}
\]

with corresponding boundary conditions:

\[
\begin{align*}
u_1(-\infty) &= u_3(\infty) = 0, \\
u_1(\alpha) &= u_2(\alpha), \\
u_2(\beta) &= u_3(\beta), \\
\frac{du_1}{dx}|_{x=\alpha} &= \frac{du_2}{dx}|_{x=\alpha}, \\
\frac{du_2}{dx}|_{x=\beta} &= \frac{du_3}{dx}|_{x=\beta}
\end{align*}
\]

The equations are clearly second order linear partial differential equations, where \( u_{1,2} \) are homogenous and \( u_3 \) is inhomogeneous. The derivation of the homogenous solution is equivalent for all 3 equations, which differ only in their unknown coefficients. Therefore we can consider the homogenous solution for all 3 equations,

\[
D \frac{d^2 u_i}{dx^2} + c \frac{du_i}{dx} - k_1 u_i = 0
\]  

(A.10)
were from the form of (A.1) we can see the solutions will be of the form

\[ u = c_1 U_1(x) + c_2 U_2(x) \]  

(A.11)

were \(c_1\) and \(c_2\) are unknown coefficients and \(U_1\) and \(U_2\) are linearly independent solutions of (A.10). Let us assume the solution is of the form \(u_j = e^{\lambda_j x}\) and substitute this expression into (A.10) giving:

\[ D\lambda^2 e^{\lambda x} + c\lambda e^{\lambda x} - k_1 e^{\lambda x} = 0, \]

\[ \Rightarrow D\lambda^2 + c\lambda - k_1 = 0, \]  

(A.12)

the solutions of the characteristic equation (A.12) give the corresponding eigenvalues:

\[ \Rightarrow \lambda_{1,2} = \frac{-c \pm \sqrt{c^2 + 4k_1D}}{2D}, \]  

(A.13)

and therefore we can substitute (A.12) into (A.11) giving us the complementary solution

\[ u = c_1 e^{\lambda_1 x} + c_2 e^{\lambda_2 x} \]  

(A.14)

Finally we can determine the particular solutions of \(u_3\) by noting that the forcing term is a constant and therefore we can assume that the solution is of the form:

\[ u_3 = c_3 k_2 \]  

(A.15)

were again \(c_3\) is an unknown coefficient. Substituting (A.11) into (A.10) we find:

\[ k_1 c_3 k_2 = k_2, \]

\[ \Rightarrow c_3 = \frac{1}{k_1} \]  

(A.16)

Therefore we can define the general solution of (A.1) as the system:

\[ u_i = \begin{cases} 
  u_1 = A e^{\lambda_1 x} + B e^{\lambda_2 x}, & x < \alpha \\
  u_2 = C e^{\lambda_1 x} + D e^{\lambda_2 x} + \frac{k_2}{k_1}, & \alpha < x < \beta \\
  u_3 = E e^{\lambda_1 x} + F e^{\lambda_2 x}, & x > \beta 
\end{cases} \]  

(A.17)

To solve for the remaining coefficients we can employ boundary conditions ((A.6)- (A.9)) with which we can define the following system of equations:

\[ A e^{\lambda_1} = C e^{\lambda_1} + D e^{\lambda_2} + \frac{k_2}{k_1} \]  

(A.18)

\[ E e^{\lambda_1} = C e^{\lambda_1} + D e^{\lambda_2} + \frac{k_2}{k_1} \]  

(A.19)

\[ \lambda_1 A e^{\lambda_1} = \lambda_2 C e^{\lambda_1} + \lambda_2 D e^{\lambda_2} \]  

(A.20)

\[ \lambda_1 E e^{\lambda_1} = \lambda_2 C e^{\lambda_1} + \lambda_2 D e^{\lambda_2} \]  

(A.21)

which can be solved algebraically. Solve (A.18) for \(A\) and (A.19) for \(E\):

\[ A = C + D e^{(\lambda_2 - \lambda_1)} + \frac{k_2}{k_1} e^{-\lambda_1} \]  

(A.22)

\[ E = C e^{(\lambda_1 - \lambda_2)} + D + \frac{k_2}{k_1} e^{-\lambda_2} \]  

(A.23)
Substitute (A.22) into (A.20):
\[ \lambda_1 \left( C + D e^{a(\beta_2 - \beta_1)} + \frac{k_2}{k_1} e^{-a \lambda_1} \right) e^{a \lambda_1} = \lambda_1 C e^{a \lambda_1} + \lambda_2 D e^{a \lambda_2} \]  
(A.24)

Solve (A.24) for D:
\[ \lambda_1 \left( C + D e^{a(\beta_2 - \beta_1)} + \frac{k_2}{k_1} e^{-a \lambda_1} \right) e^{a \lambda_1} = \lambda_1 C e^{a \lambda_1} + \lambda_2 D e^{a \lambda_2} \]
\[ C + D e^{a(\beta_2 - \beta_1)} + \frac{k_2}{k_1} e^{-a \lambda_1} = \frac{\lambda_2}{\lambda_1} D e^{a(\beta_2 - \beta_1)} \]
\[ D e^{a(\beta_2 - \beta_1)} = \frac{\lambda_2}{\lambda_1} D e^{a(\beta_2 - \beta_1)} - \frac{k_2}{k_1} e^{-a \lambda_1} \]
\[ D \left( \lambda_1 e^{a \lambda_1} - \lambda_2 e^{a \lambda_2} \right) = - \frac{k_2}{k_1} \lambda_1 \]
\[ D = \frac{\lambda_2 k_2}{k_1 (\lambda_1 e^{a \lambda_1} - \lambda_2 e^{a \lambda_2})} \]
(A.25)

Substitute (A.23) into (A.21):
\[ \lambda_2 \left( C e^{\beta(\alpha_1 - \alpha_2)} + D + \frac{k_2}{k_1} e^{-\beta \lambda_2} \right) e^{\beta \lambda_2} = \lambda_2 C e^{\beta \lambda_2} + \lambda_2 D e^{\beta \lambda_2} \]
\[ C e^{e^{\beta \lambda_2}} = \frac{\lambda_2}{\lambda_2} C e^{\beta(\alpha_1 - \alpha_2)} + D \]
\[ C e^{e^{\beta \lambda_2}} = \frac{\lambda_2}{\lambda_2} C e^{\beta(\alpha_1 - \alpha_2)} = \frac{k_2}{k_1} e^{-\beta \lambda_2} \]
\[ C \left( \lambda_2 e^{\beta \lambda_2} - \lambda_2 e^{\beta \lambda_2} \right) = - \frac{\lambda_2 k_2}{k_1} e^{\beta \lambda_2} \]
\[ C = \frac{-\frac{\lambda_2 k_2}{k_1} e^{\beta \lambda_2}}{k_1 (\lambda_1 e^{\beta \lambda_1} - \lambda_2 e^{\beta \lambda_2})} \]
(A.26)

Now find A by substituting (A.25) and (A.26) into (A.18):
\[ A e^{a \lambda_1} = \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_1} - \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1} \]
\[ A e^{a \lambda_1} = \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1} \]
\[ A e^{a \lambda_1} = \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1} \]
\[ A e^{a \lambda_1} = \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1} \]
\[ A e^{a \lambda_1} = \frac{k_2}{k_1 (\lambda_1 - \lambda_2)} e^{a \lambda_2} + \frac{k_2}{k_1} \]
\[ A = \frac{k_2}{k_1} \left( e^{-a \lambda_1} - e^{-a \lambda_1} \right) \]
(A.27)
Now find $E$ by substituting (A.25) and (A.26) into (A.19):

$$E \beta \lambda_2 = \frac{k_2}{k_1} \lambda_2 \left( e^{\beta \lambda_1} \frac{e^{\beta \lambda_1}}{\lambda_1} - \frac{k_2}{k_1} \lambda_2 \right) + \frac{k_2}{k_1} \lambda_2$$

$$E \beta \lambda_2 = \frac{k_2}{k_1} \lambda_2 e^{\beta \lambda_1} k_1(\lambda_1 - \lambda_2) e^{\alpha x_1} k_1 - k_1(\lambda_1 - \lambda_2) e^{\beta \lambda_1} k_1 \lambda_1 e^{\beta \lambda_2} + k_2(\lambda_1 - \lambda_2) e^{\beta \lambda_1} k_1(\lambda_1 - \lambda_2) e^{\alpha x_1} k_1$$

$$E \beta \lambda_2 = \frac{k_2}{k_1} \lambda_2 - k_2 \lambda_1 e^{\beta \lambda_1} - e^{\beta \lambda_2} + k_2(\lambda_1 - \lambda_2)$$

$$E = -\frac{k_2}{k_1} \lambda_2 e^{\beta \lambda_1} - e^{\beta \lambda_2} + k_2(\lambda_1 - \lambda_2)$$

$$E = \frac{k_2}{k_1} \lambda_2 - e^{\beta \lambda_2} - e^{\alpha x_1} k_1(\lambda_1 - \lambda_2)$$

Thus with expressions for $A, C, D$ and $E$ we can now substitute these values into (A.17) and after a little simplification we finally have:

$$u_1 = \begin{cases} 
\frac{k_2 \lambda_2}{k_1 (\lambda_1 - \lambda_2)} \left( e^{\beta (x-\beta)} - e^{\alpha (x-\alpha)} \right), & x < \alpha \\
\frac{k_2}{k_1 (\lambda_1 - \lambda_2)} \left( \lambda_2 e^{\beta (x-\beta)} - \lambda_2 e^{\alpha (x-\alpha)} \right), & \alpha < x < \beta \\
\frac{k_2 \lambda_2}{k_1 (\lambda_1 - \lambda_2)} \left( e^{\beta (x-\beta)} - e^{\alpha (x-\alpha)} \right), & x > \beta 
\end{cases}$$

$$u_1 = \begin{cases} 
\frac{k_2 \lambda_2}{k_1 (\lambda_1 - \lambda_2)} \left( e^{\beta (x-\beta)} - e^{\alpha (x-\alpha)} \right), & x < \alpha \\
\frac{k_2}{k_1 (\lambda_1 - \lambda_2)} \left( \lambda_2 e^{\beta (x-\beta)} - \lambda_2 e^{\alpha (x-\alpha)} \right), & \alpha < x < \beta \\
\frac{k_2 \lambda_2}{k_1 (\lambda_1 - \lambda_2)} \left( e^{\beta (x-\beta)} - e^{\alpha (x-\alpha)} \right), & x > \beta 
\end{cases}$$
Appendix B

Derivation of Cubic Approximation to the Chemotaxis Function

In this document we wish to find a third order approximation to the chemotaxis function of equation (2.14) derived in the main body of the thesis in section 2.2.2 Motion Due To Chemotaxis, reproduced here for convenience:

\[ c(a; c_0) = \frac{k_2c_0}{ak_1(\lambda_1 - \lambda_2)} \left( \lambda_1 (1 - e^{ak_2}) + \lambda_2 (1 - e^{-ak_1}) \right). \] (B.1)

We start by stating the requirement of taking the 3rd order derivative and deriving this expression then substituting this expression in to the Taylor approximation for \( c = 0 \). Thus we first need to find the complete expression for:

\[ \frac{d^3c(a; c_0)}{dc^3} = \frac{k_2c_0}{ak_1} \left( \frac{d^3}{dc^3} \left( \frac{\lambda_1 - \lambda_1 e^{ak_2}}{\lambda_1 - \lambda_2} \right) + \frac{d^3}{dc^3} \left( \frac{\lambda_2 - \lambda_2 e^{-ak_1}}{\lambda_1 - \lambda_2} \right) \right) \] (B.2)

First however we can make the observation that the third order derivative expressions in (B2) are equal:

\[ \frac{d^3}{dc^3} \left( \frac{\lambda_1 - \lambda_1 e^{ak_2}}{\lambda_1 - \lambda_2} \right) = \frac{d^3}{dc^3} \left( \frac{\lambda_2 - \lambda_2 e^{-ak_1}}{\lambda_1 - \lambda_2} \right) \] (B.3)

For simplicity we shall first expand these expressions in terms of the derivative operator, and then individually find the corresponding evaluations of these derivatives. We choose to take this approach for clarity as the expression will become very cumbersome. However we cannot arrive at our goal without evaluating the intermediate steps, so let us consider a symbolic derivation of the derivatives in (B3) and later derive each derivative expression separately. Thus first let us make the following substitutions:

\[ f(c) = \lambda_1 - \lambda_1 e^{ak_2} \text{ and } g(c) = \lambda_1 - \lambda_2 \]

\[ \Rightarrow \frac{d^3}{dc^3} \left( \frac{\lambda_1 - \lambda_1 e^{ak_2}}{\lambda_1 - \lambda_2} \right) = \frac{d^3}{dc^3} \left( \frac{f(c)}{g(c)} \right) \] (B.4)

Now let us determine the expression for (B4) starting with:

\[ \frac{d}{dc} \left( \frac{f(c)}{g(c)} \right) = \frac{df(c)}{dc} g^{-1}(c) + f(c) \frac{dg^{-1}(c)}{dc} \] (B.5)

and

\[ \frac{d^2}{dc^2} \left( \frac{f(c)}{g(c)} \right) = \frac{df(c)}{dc} g^{-1}(c) - f(c) g^{-2}(c) \frac{dg(c)}{dc} \] (B.6)
\[
\frac{d^3 f(c)}{dc^3} = \frac{d}{dc} \left( \frac{d^3 f(c)}{dc^3} g^{-1}(c) - \frac{df(c)}{dc} g^{-2}(c) \frac{dg(c)}{dc} \right. \\
\left. - \frac{df(c)}{dc} g^{-2}(c) \frac{dg(c)}{dc} + 2f(c)g^{-3}(c) \left( \frac{dg(c)}{dc} \right)^2 - f(c)g^{-2}(c) \frac{d^2 g(c)}{dc^2} \right) \\
= \left( \frac{d^3 f(c)}{dc^3} g^{-1}(c) - \frac{d^2 f(c)}{dc^2} g^{-2}(c) \frac{dg(c)}{dc} \right.
\]
\[
- \frac{df(c)}{dc} g^{-2}(c) \frac{dg(c)}{dc} + 2 \frac{df(c)}{dc} g^{-3}(c) \left( \frac{dg(c)}{dc} \right)^2 - \frac{df(c)}{dc} g^{-2}(c) \frac{d^2 g(c)}{dc^2} \\
+ 2 \frac{df(c)}{dc} g^{-3}(c) \left( \frac{dg(c)}{dc} \right)^2 - 6f(c)g^{-4}(c) \left( \frac{dg(c)}{dc} \right)^3 + 4f(c)g^{-3}(c) \frac{dg(c)}{dc} \frac{d^2 g(c)}{dc^2} \\
- \frac{df(c)}{dc} g^{-2}(c) \frac{d^2 g(c)}{dc^2} + 2f(c)g^{-3}(c) \frac{d^2 g(c)}{dc^2} \frac{dg(c)}{dc} - f(c)g^{-2}(c) \frac{d^3 g(c)}{dc^3} \\
= \left( \frac{d^3 f(c)}{dc^3} g^{-1}(c) - 3 \frac{d^2 f(c)}{dc^2} g^{-2}(c) \frac{dg(c)}{dc} \right.
\]
\[
+ 6 \frac{df(c)}{dc} g^{-3}(c) \left( \frac{dg(c)}{dc} \right)^2 + 6f(c)g^{-4}(c) \frac{dg(c)}{dc} \frac{d^2 g(c)}{dc^2} - 6f(c)g^{-3}(c) \left( \frac{dg(c)}{dc} \right)^3 \\
- 3 \frac{df(c)}{dc} g^{-2}(c) \frac{d^2 g(c)}{dc^2} - f(c)g^{-2}(c) \left( \frac{dg(c)}{dc} \right)^2 \right) 
\]
\[
(B.7)
\]

From the expression in (B.7) we can see there are 1st, 2nd and 3rd order derivatives in both \( f(c) \) and \( g(c) \), so now we shall evaluate all of these derivatives and substitute them back into (B.7). Further since we are deriving the cubic for a Taylor approximation, we shall evaluate each derivative at \( c = 0 \) before substitution. First let us find the derivatives for \( f(c) \) for \( \lambda_1 \):
\[
\frac{df(c)}{dc} = \frac{d(\lambda_1 - \lambda_1 e^{a_1 v})}{dc} = \frac{d\lambda_1}{dc} (1 - e^{a_1 v}) + \lambda_1 \frac{d(1 - e^{a_1 v})}{dc} \\
= \frac{d\lambda_1}{dc} (1 - e^{a_1 v}) - \lambda_1 a \frac{d^2 \lambda_2}{dc} e^{a_1 v} 
\]
\[
(B.8)
\]
and the second derivative of \( f(c) \):
\[
\frac{d^2 f(c)}{dc^2} = \frac{d^2 (\lambda_1 - \lambda_1 e^{a_1 v})}{dc^2} = \frac{d}{dc} \left( \frac{d\lambda_1}{dc} - \lambda_1 a \frac{d^2 \lambda_2}{dc} e^{a_1 v} \right) \\
= \left( \frac{d^2 \lambda_1}{dc^2} - \frac{d^2 \lambda_1}{dc^2} e^{a_1 v} - a \frac{d^2 \lambda_2}{dc} e^{a_1 v} \\
- \lambda_1 a \frac{d^2 \lambda_2}{dc^2} e^{a_1 v} - \lambda_1 a^2 \left( \frac{d\lambda_2}{dc} \right)^2 e^{a_1 v} \right) \\
= \left( \frac{d^2 \lambda_1}{dc^2} - \frac{d^2 \lambda_1}{dc^2} e^{a_1 v} - 2a \frac{d\lambda_1}{dc} \frac{d\lambda_2}{dc} e^{a_1 v} - \lambda_1 a \frac{d^2 \lambda_2}{dc^2} e^{a_1 v} - \lambda_1 a^2 \left( \frac{d\lambda_2}{dc} \right)^2 e^{a_1 v} \right) 
\]
\[
(B.9)
\]
And the third derivative of $f(c)$:

$$
\frac{d^3 f(c)}{dc^3} = \frac{d^3(\lambda_1 - \lambda_1 e^{a\lambda_2})}{dc^3} = \frac{d}{dc} \left( \frac{d^2 \lambda_1}{dc^2} - \frac{d^2 \lambda_2}{dc^2} e^{a\lambda_2} - 2a \frac{d \lambda_1}{dc} \frac{d \lambda_2}{dc} e^{a\lambda_2} \right)
$$

$$
-\lambda_1 a \frac{d^2 \lambda_2}{dc^2} e^{a\lambda_2} = \lambda_1 a^2 \left( \frac{d \lambda_2}{dc} \right)^2 e^{a\lambda_2}
$$

$$
= \frac{d^3 \lambda_1}{dc^3} - \frac{d^3 \lambda_2}{dc^3} e^{a\lambda_2} - a \frac{d^2 \lambda_1}{dc^2} \frac{d \lambda_2}{dc} e^{a\lambda_2}
$$

$$
-2a \frac{d \lambda_1}{dc} \frac{d \lambda_2}{dc} e^{a\lambda_2} - 2a \frac{d \lambda_1}{dc} \frac{d^2 \lambda_2}{dc^2} e^{a\lambda_2} - 2 \lambda_1 a^2 \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^2 e^{a\lambda_2}
$$

$$
- \lambda_1 a \frac{d^3 \lambda_2}{dc^3} e^{a\lambda_2} - \lambda_1 a \frac{d^2 \lambda_2}{dc^2} \frac{d \lambda_2}{dc} e^{a\lambda_2} - \lambda_1 a \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^2 e^{a\lambda_2}
$$

$$
- \lambda_1 a \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^2 e^{a\lambda_2} - 2 \lambda_1 a^2 \frac{d^2 \lambda_2}{dc^2} \frac{d \lambda_2}{dc} e^{a\lambda_2} - \lambda_1 a^3 \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^3 e^{a\lambda_2}
$$

$$
= \frac{d^3 \lambda_1}{dc^3} - \frac{d^3 \lambda_1}{dc^3} e^{a\lambda_2} - 3a \frac{d^2 \lambda_1}{dc^2} \frac{d \lambda_2}{dc} e^{a\lambda_2}
$$

$$
- 3a \frac{d \lambda_1}{dc} \frac{d^2 \lambda_2}{dc^2} e^{a\lambda_2} - 3a \frac{d \lambda_1}{dc} \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^2 e^{a\lambda_2}
$$

$$
- 3a^2 \frac{d \lambda_1}{dc} \frac{d \lambda_2}{dc} \frac{d \lambda_2}{dc} e^{a\lambda_2} - \lambda_1 a \frac{d^3 \lambda_2}{dc^3} e^{a\lambda_2} - \lambda_1 a \frac{d^2 \lambda_2}{dc^2} \frac{d \lambda_2}{dc} e^{a\lambda_2} - \lambda_1 a^3 \frac{d \lambda_2}{dc} \left( \frac{d \lambda_2}{dc} \right)^3 e^{a\lambda_2}
$$

Now let’s consider the derivatives for $f(c)$ for $\lambda_2$:

$$
\frac{df}{dc} = \frac{d}{dc} (\lambda_2 - \lambda_2 e^{-a\lambda_1}) = \frac{d \lambda_2}{dc} \left( 1 - e^{-a\lambda_1} \right) + \lambda_2 a \frac{d \lambda_1}{dc} e^{-a\lambda_1}
$$

$$
= \frac{d \lambda_2}{dc} \left( 1 - e^{-a\lambda_1} \right) + \lambda_2 a \frac{d \lambda_1}{dc} e^{-a\lambda_1}
$$

And now the second derivative:

$$
\frac{d^2 f}{dc^2} = \frac{d^2 (\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^2} = \frac{d}{dc} \left( \frac{d \lambda_2}{dc} - \frac{d \lambda_2}{dc} e^{-a\lambda_1} + \lambda_2 a \frac{d \lambda_1}{dc} e^{-a\lambda_1} \right)
$$

$$
= \left( \frac{d^2 \lambda_2}{dc^2} - \frac{d^2 \lambda_2}{dc^2} e^{-a\lambda_1} + 2 \frac{d \lambda_2}{dc} \frac{d \lambda_1}{dc} e^{-a\lambda_1} + \lambda_2 a \frac{d^2 \lambda_1}{dc^2} e^{-a\lambda_1} - \lambda_2 a \frac{d \lambda_1}{dc} \left( \frac{d \lambda_2}{dc} \right)^2 e^{-a\lambda_1} \right)
$$

(B.10)
And the third derivative:

\[
\frac{d^3 f(c)}{dc^3} = \frac{d^3(\lambda_2 - \lambda_2 e^{-a_3})}{dc^3} = \frac{d^2\lambda_2}{dc^2} - \frac{d^2\lambda_2}{dc^2} e^{-a_3} + 2a \frac{d\lambda_2}{dc} \frac{d\lambda_1}{dc} e^{-a_3} + \lambda_2 a \frac{d^2\lambda_1}{dc^2} e^{-a_3} - \lambda_2 a^2 \left(\frac{d\lambda_1}{dc}\right)^2 e^{-a_3} + \lambda_2 a^3 e^{-a_3} \left(\frac{d\lambda_1}{dc}\right)^3 e^{-a_3} + \lambda_2 a^2 \frac{d\lambda_1}{dc} e^{-a_3} + \lambda_2 \frac{d\lambda_1}{dc} e^{-a_3} \left(\frac{d\lambda_1}{dc}\right)^3 e^{-a_3}
\]

We can find the derivatives of \( g(c) \) by evaluating the derivatives of \( \lambda_{1,2} \) up to third order for \( c = 0 \):

\[
\lambda_{1,2} = \left. \frac{d\lambda_{1,2}}{dc} \right|_{c=0} = \frac{\pm \sqrt{c^2 + 4k_1d}}{2d} \quad \text{(B.14)}
\]

\[
\frac{d\lambda_{1,2}}{dc} = \left. \frac{d^2\lambda_{1,2}}{dc^2} \right|_{c=0} = \frac{\pm 2k_1}{2d\sqrt{c^2 + 4k_1d}} = \frac{\pm 1}{4d\sqrt{k_1d}} \quad \text{(B.15)}
\]

And so we can substitute (B14-B17) into the corresponding derivatives of \( \frac{d^{(n)}g(c)}{dc^n} \) for \( n = 0,1,2,3 \):

\[
g(c)|_{c=0} = \left( -\frac{k_1}{d} \right) = 2 \frac{k_1}{d} \quad \text{(B.18)}
\]

\[
\left. \frac{d^2g(c)}{dc^2} \right|_{c=0} = 0 \quad \text{(B.19)}
\]

\[
\left. \frac{d^3g(c)}{dc^3} \right|_{c=0} = 0 \quad \text{(B.20)}
\]

\[
\left. \frac{d^4g(c)}{dc^4} \right|_{c=0} = 0 \quad \text{(B.21)}
\]
We have found all of the expressions for the derivatives and now we shall back substitute all of the values in reverse order to find the approximation at \( c = 0 \).

\[
\left. \frac{d^3(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^3} \right|_{c=0} = \left( -3a \frac{1}{4d\sqrt{k_1 d}} \left( -\frac{1}{2d} \right) e^{-a\sqrt{k_1 d}} + 3a \left( \frac{1}{2d} \right) \frac{1}{4d\sqrt{k_1 d}} e^{-a\sqrt{k_1 d}} \right) - 3a^2 \left( -\frac{1}{2d} \right)^2 e^{-a\sqrt{k_1 d}} \\
+ 3 \frac{k_1}{d} a^2 \left( \frac{1}{4d\sqrt{k_1 d}} \right) - \frac{1}{2d} e^{-a\lambda_1} - \frac{k_1}{d} a^3 \left( -\frac{1}{2d} \right)^3 e^{-a\sqrt{k_1 d}}
\]

\[
= \left( 3a - \frac{1}{8d^2\sqrt{k_1 d}} - 3a \frac{1}{8d^2\sqrt{k_1 d}} + 3a^2 \frac{1}{8d^3} - 3 \frac{k_1}{d} a^2 \frac{1}{8d^2\sqrt{k_1 d}} + \frac{k_1}{d} a^3 \frac{1}{8d^3} \right) e^{-a\sqrt{k_1 d}}
\]

\[
= \left( 3a^2 \frac{1}{8d^3} - 3a^2 \frac{1}{8d^3} + \frac{k_1}{d} a^3 \frac{1}{8d^3} \right) e^{-a\sqrt{k_1 d}}
\]

\[
= \left( \frac{k_1}{d} a^3 \frac{1}{8d^3} \right) e^{-a\sqrt{k_1 d}}
\]

(B.22)

And therefore we have:

\[
\left. \frac{d^3(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^3} \right|_{c=0} = a^3\sqrt{k_1 d} \frac{e^{-a\sqrt{k_1 d}}}{8d^4}
\]  

(B.23)

Now we can make the observation that while signs have changed the \( \lambda \)s they have also changed position then so will the signs of the resultant derivatives therefore:

\[
\left. \frac{d^3(\lambda_1 - \lambda_1 e^{a\lambda_1})}{dc^3} \right|_{c=0} = \left. \frac{d^3(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^3} \right|_{c=0} = a^3\sqrt{k_1 d} \frac{e^{-a\sqrt{k_1 d}}}{8d^4}
\]

(B.24)

Now find:

\[
\left. \frac{d^2(\lambda_1 - \lambda_1 e^{a\lambda_1})}{dc^2} \right|_{c=0} = \left. \frac{d^2(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^2} \right|_{c=0} = \frac{d^2\lambda_2}{dc^2} - \frac{d^2\lambda_2}{dc^2} e^{-a\lambda_1} + 2a \frac{d\lambda_2}{dc} \frac{d\lambda_1}{dc} e^{-a\lambda_1} + \lambda_2 a \frac{d^2\lambda_1}{dc^2} e^{-a\lambda_1} - \lambda_2 a^2 \left( \frac{d\lambda_1}{dc} \right)^2 e^{-a\lambda_1}
\]

\[
= -\frac{1}{4d\sqrt{k_1 d}} + \frac{1}{4d\sqrt{k_1 d}} e^{-a\lambda_1} + 2a \frac{1}{4d^2} e^{-a\lambda_1}
\]

\[
- \frac{k_1}{d} a \left( \frac{1}{4d\sqrt{k_1 d}} \right) e^{-a\lambda_1} + \frac{k_1}{d} a^2 \left( \frac{1}{4d^2} \right) e^{-a\lambda_1}
\]

\[
= -\frac{1}{4d\sqrt{k_1 d}} + \frac{a\lambda_1}{4d\sqrt{k_1 d}} + \frac{ae^{-a\lambda_1}}{2d^2} - \frac{ae^{-a\lambda_1}}{4d^2} + \frac{a^2\sqrt{k_1 d} e^{-a\lambda_1}}{4d^2\sqrt{k_1 d}}
\]

\[
= -a\lambda_1 + 2a\sqrt{k_1 d} e^{-a\lambda_1} - a\sqrt{k_1 d} e^{-a\lambda_1} + \frac{1}{2d} \frac{k_1}{d} a e^{-a\lambda_1}
\]

\[
= \frac{-d + de^{-a\lambda_1} + a\sqrt{k_1 d} e^{-a\lambda_1} - a\sqrt{k_1 d} e^{-a\lambda_1} + a^2\sqrt{k_1 d} e^{-a\lambda_1}}{4d^2\sqrt{k_1 d}}
\]

(B.25)
And following a similar argument it can be shown that:

\[
\frac{d^2(\lambda_1 - \lambda_2 e^{a\lambda_1})}{dc^2} \bigg|_{c=0} = -\frac{d^2(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc^2} \bigg|_{c=0} = -\frac{-d + de^{-a\lambda_1} + a\sqrt{k_1} e^{-a\lambda_1} + a^2 k_1 e^{-a\lambda_1}}{4d^2}\sqrt{k_1 d} \quad (B.26)
\]

Now find:

\[
\frac{d(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc} \bigg|_{c=0} = \frac{1}{2d} + \frac{1}{2d} e^{-a\sqrt{\frac{k_1}{d}}} + \frac{\sqrt{k_1}}{d} \frac{1}{2d} e^{-a\sqrt{\frac{k_1}{d}}}
\]

\[
= -\frac{1}{2d} + \frac{e^{-a\sqrt{\frac{k_1}{d}}}}{2d} + \frac{a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{2d^2}
\]

\[
= -\frac{-d + de^{-a\sqrt{\frac{k_1}{d}}} + a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{2d^2} \quad (B.27)
\]

And following a similar argument to (B.5.3.1) we it can be shown that:

\[
\frac{d(\lambda_1 - \lambda_2 e^{a\lambda_1})}{dc} \bigg|_{c=0} = \frac{d(\lambda_2 - \lambda_2 e^{-a\lambda_1})}{dc} \bigg|_{c=0} = -\frac{-d + de^{-a\sqrt{\frac{k_1}{d}}} + a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{2d^2} \quad (B.28)
\]

Now we have the derivatives evaluated at \(c = 0\) (B17-B28) for substitution into (B7):

\[
\frac{d^3}{dc^3}(f(x)) \bigg|_{c=0} = \frac{a^3 \sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{8d^4} \left(\frac{2\sqrt{k_1}}{\sqrt{d}}\right)^{-1}
\]

\[
= -3 \left(-\frac{-d + de^{-a\sqrt{\frac{k_1}{d}}} + a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{2d^2}\right)^{-2} \frac{1}{2d \sqrt{k_1 d}}
\]

\[
= \frac{a^3 \sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{8d^4} \frac{1}{2\sqrt{k_1} d}
\]

\[
= -3 \left(-\frac{-d + de^{-a\sqrt{\frac{k_1}{d}}} + a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{2d^2}\right)^{-1} \frac{1}{4k_1 2d \sqrt{k_1 d}}
\]

\[
= \frac{a^3 e^{-a\sqrt{\frac{k_1}{d}}}}{16d^3} - \frac{3}{16} \left(-\frac{-d + de^{-a\sqrt{\frac{k_1}{d}}} + a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{d^2 k_1 \sqrt{k_1 d}}\right)
\]

\[
= \frac{a^3 e^{-a\sqrt{\frac{k_1}{d}}}}{16d^3} + \frac{3d - 3de^{-a\sqrt{\frac{k_1}{d}}} - 3a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{16d^2 k_1 \sqrt{k_1 d}}
\]

\[
= \frac{1}{16d^2 k_1 \sqrt{k_1 d}} \left(\frac{k_1 \sqrt{k_1} d a^3 e^{-a\sqrt{\frac{k_1}{d}}} + 3d - 3de^{-a\sqrt{\frac{k_1}{d}}} - 3a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{d}\right)
\]

\[
= \frac{1}{16d^2 k_1 \sqrt{k_1 d}} \left(\frac{k_1 \sqrt{k_1} d a^3 e^{-a\sqrt{\frac{k_1}{d}}} + 3d - 3de^{-a\sqrt{\frac{k_1}{d}}} - 3a\sqrt{k_1} d e^{-a\sqrt{\frac{k_1}{d}}}}{d}\right)
\]

\[
= \frac{3d}{16d^2 k_1 \sqrt{k_1 d}} \left(e^{-a\sqrt{\frac{k_1}{d}}} \left(\frac{k_1(a^3 - 3ad)}{d^2} - 1\right) + 1\right)
\]
\[
\begin{align*}
= & \frac{3}{16d k_1 \sqrt{k_1 d}} \left( e^{-\frac{\kappa_1}{\sqrt{d}}} \left( \frac{k_1 (k_1 a^3 - 3ad)}{d} - 1 \right) + 1 \right) \\
& \quad \text{(B.29)}
\end{align*}
\]

Now considering (B2), (B3) we can now substitute (B29) into a third order Taylor approximation, first however we note that for an odd function we have:

\[
-h(c) = h(-c) \implies \left. \frac{d^{2n} h(c)}{dc^{2n}} \right|_{c=0} = 0; \quad \forall n \in \mathbb{N}
\]

so when we take the cubic Taylor approximation:

\[
h(c) \approx \sum_{n=0}^{3} \frac{h^{(n)}(0)}{n!} (c - 0)^n
\]

\[
= \frac{h^{(0)}(0)}{0!} (c - 0)^0 + \frac{h^{(1)}(0)}{1!} (c - 0)^1 + \frac{h^{(2)}(0)}{2!} (c - 0)^2 + \frac{h^{(3)}(0)}{3!} (c - 0)^3
\]

we can show that \( h(0) = h^{(2)}(0) = 0 \) then,

\[
\frac{1}{1!} \left. \frac{d h(c)}{d c} \right|_{c=0} c + \frac{1}{3!} \left. \frac{d^3 h(c)}{d c^3} \right|_{c=0} c^3.
\]

We can find linear term from (B5):

\[
\frac{d}{d c} \left( f(c) \right) = \left( \frac{-d + de^{-\frac{\kappa_1}{\sqrt{a}}} + a \sqrt{k_1 d} e^{-\frac{\kappa_1}{\sqrt{a}}}}{2 \sqrt{d}} \right) \left( \frac{2 \sqrt{k_1}}{d} \right)^{-1}
\]

\[
= \left( \frac{-d^2 + 2 d^2 e^{-\frac{\kappa_1}{\sqrt{a}}} + 2d \sqrt{k_1 d} e^{-\frac{\kappa_1}{\sqrt{a}}}}{4 \sqrt{k_1}} \right)
\]

\[
= \left( \frac{-1 + \left( 1 + \frac{k_1}{d} \right) e^{-\frac{\kappa_1}{\sqrt{a}}} + a \sqrt{k_1 d} e^{-\frac{\kappa_1}{\sqrt{a}}}}{4 \sqrt{k_1}} \right)
\]

and substituting (B29) and (B31) into (B30) and we finally have a third order Taylor approximation to (B3):

\[
c(a; c_0) = \frac{2k_1 c_0}{a k_1} \left( \frac{1}{1!} \left( 1 + \frac{k_1}{d} \right) + a \sqrt{k_1 d} e^{-\frac{\kappa_1}{\sqrt{a}}} \right) c + \frac{2k_1 c_0}{a k_1} \left( \frac{3}{3!} \left( \frac{3}{16d k_1 \sqrt{k_1 d}} \left( e^{-\frac{\kappa_1}{\sqrt{a}}} \left( a \frac{k_1 (k_1 a^3 - 3ad)}{d} - 1 \right) + 1 \right) \right) \right) c^3
\]

And finally:

\[
c(a; c_0) = \frac{k_1 c_0}{2a k_1 \sqrt{k_1 d}} \left( -1 + \left( 1 + \frac{k_1}{d} \right) e^{-\frac{\kappa_1}{\sqrt{a}}} \right) c + \frac{k_1 c_0}{16a k_1 \sqrt{k_1 d}} \left( e^{-\frac{\kappa_1}{\sqrt{a}}} \left( \frac{k_1 (k_1 a^3 - 3ad)}{d} - 1 \right) + 1 \right) c^3
\]

\[
\text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ } \text{ }
Appendix C
CARTESIAN TO POLAR COORDINATE OF THE 1D MODEL EQUATION.

We consider an arbitrary function of two variable function $\phi = \phi(x, y)$, such that $\phi \in C^2$, that is $\phi$ is continuous differentiable function in its first and second derivatives of its Cartesian variables $x, y$. From our one-dimensional model equation:

$$\frac{\partial u}{\partial t} = D\Delta u + c(\nabla u; c_0) \frac{du}{dx} - k_1 u = p(x); \quad k_1, k_2, D \in \mathbb{R}^+,$$  \hspace{1cm} (C.1)

we will show a conversion from Cartesian to polar coordinates, however first let us consider the differential operator:

$$\Delta = \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right),$$  \hspace{1cm} (C.2)

referred to as the Laplacian. The transformation into the polar coordinate system, that is $x, y \rightarrow r, \theta$, were $r$ is the radial component and $\theta$ is the angular rotation about an arbitrary pole, which for simplicity can be taken as the origin, is achieved by relating the Cartesian system to the polar systems $(r, \theta)$:

$$x = r\cos(\theta)$$
$$y = r\sin(\theta),$$  \hspace{1cm} (C.3)

and conversely by

$$r = \sqrt{x^2 + y^2}$$
$$\theta = \arctan\left(\frac{y}{x}\right).$$  \hspace{1cm} (C.4)

The important point to note is that when performing the transformation, the variables are function of each other, that is we need to keep in mind that $\theta(x, y), r(x, y)$ and $x(r, \theta), y(r, \theta)$ when we are taking derivates. Thus applying the chain rule from the Calculus on $\phi(\theta, r)$ with respect to the Cartesian variables $x$ and $y$ we find:

$$\frac{\partial \phi}{\partial x} = \frac{\partial \phi}{\partial \theta} \frac{\partial \theta}{\partial x} + \frac{\partial \phi}{\partial r} \frac{\partial r}{\partial x}$$  \hspace{1cm} (C.5)

and

$$\frac{\partial \phi}{\partial y} = \frac{\partial \phi}{\partial \theta} \frac{\partial \theta}{\partial y} + \frac{\partial \phi}{\partial r} \frac{\partial r}{\partial y}$$  \hspace{1cm} (C.6)
Using the variable relationships (C.3) and (C.4):

\[
\frac{\partial \theta}{\partial x} = \frac{\partial}{\partial x} \left( \arctan \left( \frac{y}{x} \right) \right) = \frac{1}{1 + (y/x)^2} \frac{\partial}{\partial x} \left( \frac{y}{x} \right)
\]

\[
= -\frac{y}{x^2 + y^2}
\]

\[
= -\frac{\sin(\theta)}{r}
\]

\[
\frac{\partial \theta}{\partial y} = \frac{\partial}{\partial y} \left( \arctan \left( \frac{y}{x} \right) \right) = \frac{1}{1 + (y/x)^2} \frac{\partial}{\partial y} \left( \frac{y}{x} \right)
\]

\[
= \frac{x^2 - 1}{x^2 + y^2} x
\]

\[
= \frac{x}{x^2 + y^2}
\]

\[
= \frac{\cos(\theta)}{r}
\]

(C.7)

And

\[
\frac{\partial r}{\partial x} = \frac{\partial}{\partial x} \left( \sqrt{x^2 + y^2} \right) = \frac{x}{\sqrt{x^2 + y^2}} = \frac{r\cos(\theta)}{r} = \cos(\theta)
\]

(C.8)

\[
\frac{\partial r}{\partial y} = \frac{\partial}{\partial y} \left( \sqrt{x^2 + y^2} \right) = \frac{y}{\sqrt{x^2 + y^2}} = \frac{r\cos(\theta)}{r} = \sin(\theta)
\]

Now substituting

(C.7) and (C.8) into (C.5) and (C.6) we have the first derivative with respect to \(x\) in polar coordinates:

\[
\frac{\partial \phi}{\partial x} = \frac{\partial \phi}{\partial r} \cos(\theta) - \frac{\partial \phi}{\partial \theta} \frac{\cos(\theta)}{r},
\]

(C.9)

and with respect to \(y\) in polar coordinates:

\[
\frac{\partial \phi}{\partial y} = \frac{\partial \phi}{\partial r} \sin(\theta) + \frac{\partial \phi}{\partial \theta} \frac{\sin(\theta)}{r}.
\]

(C.10)

For the second derivatives we can make the observation that in (C.5) and (C.6) the chain rule would produce same for result for any \(\phi(r, \theta)\) we might choose. Therefore to find the second polar
derivative we need only substitute $\phi(r, \theta) = \partial \phi / \partial x$ into (C.5) and $\phi(r, \theta) = \partial \phi / \partial y$ into (C.6) and then it follows that:

$$\frac{\partial^2 \phi}{\partial x^2} = \frac{\partial}{\partial x} \left( \frac{\partial \phi}{\partial x} \right) + \frac{\partial}{\partial r} \left( \frac{\partial \phi}{\partial r} \right)$$

$$= \left( \frac{\partial}{\partial r} \left( \frac{\partial \phi}{\partial r} \right) \cos(\theta) - \frac{\partial \phi}{\partial r} \sin(\theta) \right) \cos(\theta) - \left( \frac{\partial}{\partial r} \left( \frac{\partial \phi}{\partial r} \right) \cos(\theta) - \frac{\partial \phi}{\partial r} \sin(\theta) \right) \sin(\theta) \right)$$

$$= \frac{\partial^2 \phi}{\partial r^2} \cos^2(\theta) - \frac{\partial^2 \phi}{\partial \theta \partial r} + \frac{\partial \phi}{\partial \theta} \sin(\theta) \cos(\theta)$$

$$- \frac{\partial \phi}{\partial r} \sin(\theta) \cos(\theta) + \frac{\partial^2 \phi}{\partial \theta^2} - \frac{\partial \phi}{\partial \theta} \sin(\theta) \cos(\theta)$$

$$= \cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2} - \frac{2}{r} \sin(\theta) \cos(\theta) \frac{\partial^2 \phi}{\partial \theta \partial r} + \frac{2}{r} \sin(\theta) \cos(\theta) \frac{\partial^2 \phi}{\partial \theta^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial r^2}}{r^2} \quad (C.11)$$

and

$$\frac{\partial^2 \phi}{\partial y^2} = \frac{\partial}{\partial y} \left( \frac{\partial \phi}{\partial y} \right) + \frac{\partial}{\partial r} \left( \frac{\partial \phi}{\partial r} \right)$$

$$= \left( \frac{\partial}{\partial \theta} \left( \frac{\partial \phi}{\partial \theta} \right) \sin(\theta) + \frac{\partial \phi}{\partial \theta} \cos(\theta) \right) \cos(\theta) - \left( \frac{\partial}{\partial \theta} \left( \frac{\partial \phi}{\partial \theta} \right) \sin(\theta) + \frac{\partial \phi}{\partial \theta} \cos(\theta) \right) \sin(\theta) \right)$$

$$= \frac{\partial^2 \phi}{\partial \theta \partial \theta} \sin(\theta) \cos(\theta) + \frac{\partial \phi}{\partial \theta} \sin(\theta) \cos(\theta)$$

$$- \frac{\partial \phi}{\partial \theta} \cos(\theta) \sin(\theta) + \frac{\partial^2 \phi}{\partial \theta^2} \sin(\theta) \cos(\theta)$$

$$= \sin^2(\theta) \frac{\partial^2 \phi}{\partial r^2} + \frac{2}{r} \sin(\theta) \cos(\theta) \frac{\partial^2 \phi}{\partial \theta \partial r} - \frac{2}{r} \sin(\theta) \cos(\theta) \frac{\partial^2 \phi}{\partial \theta^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \frac{\cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2}}{r^2} \quad (C.12)$$

If we now combine (C.11) and (C.12), noting that terms with mixed trigonometric functions cancel, we find:

$$\frac{\partial^2 \phi}{\partial x^2} + \frac{\partial^2 \phi}{\partial y^2} = \cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2}$$

$$= \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial r^2}}{r} + \frac{\cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2}}{r} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \frac{\cos^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} + \frac{\cos^2(\theta) \frac{\partial^2 \phi}{\partial r^2}}{r^2} + \frac{\sin^2(\theta) \frac{\partial^2 \phi}{\partial \theta^2}}{r^2} \quad (C.13)$$

And finally from the trigonometric identity $\cos^2(\theta) + \sin^2(\theta) = 1$ we obtain the two dimensional Laplacian in polar coordinates:

$$\Delta = \left( \frac{\partial^2}{\partial r^2} + \frac{1}{r} \frac{\partial}{\partial r} + \frac{1}{r^2} \frac{\partial^2}{\partial \theta^2} \right). \quad (C.14)$$

Therefore it is as simple matter of substituting the (C.9)and (C.14) into (C.1) and we find the polar coordinate transformation:

$$\frac{\partial u}{\partial t} = D \left( \frac{\partial^2 u}{\partial r^2} + \frac{1}{r} \frac{\partial u}{\partial r} + \frac{1}{r^2} \frac{\partial^2 u}{\partial \theta^2} \right) + c \left( \cos(\theta) \frac{\partial u}{\partial r} - \frac{\sin(\theta) \partial u}{r \partial \theta} \right) - k_1 u(r, \theta) = p(r, \theta). \quad (C.15)$$
The last thing to consider is the forcing/production term $p(r, \theta)$.  


BIBLIOGRAPHY


using pseudopod-based feedback".


