

Theoretical Biophysics: Projects

D.W. Heermann

Winter term 2010-2011

Course Organizational Information

- Study groups of 2-3 people
- The groups should meet about once a week
- Report due the last week of the semester
- Aim:
 - Develop intuition into biophysical problem solving
 - Develop skill in modelling
 - Get first experience of what research is all about
- Send me an email with your choice of a project you would like to tackle (indicate your preference with whom you would like to work with). Email due by the end of the week (Oct 15.)
- I will assemble groups and connect people
- Credits
 - Option I: Grading will be based on the handed in report
 - Option II: Oral exam (30 min)
- Songling Li will assist you

Course Organizational Information

- Hand-written submissions will not be accepted.
- Organize the report clearly.
- Integrate the graphs, diagrams, and other results into the report.
- I expect a report in a style of writing just like a journal article or a thesis.
- You must provide explicit written references to the sources you used.
- Use Latex.
- NO LATE REPORT ACCEPTED.

Capillary network formation during tissue differentiation. A mechano-biological model

Capillary network formation during tissue differentiation. A mechano-biological model

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Abstract—Angiogenesis, the formation of new capillaries from pre-existing vessels, plays a critical role during bone regeneration and repair. In addition to an appropriate mechanical environment, sufficient supply of oxygen and nutrients is critical for bone formation.

Mechano-biological models have been previously used to predict the time course of the differentiation process with the mechanical environment as the only regulator of cell activity. Here we propose a mechano-biological model for bone differentiation where cell activity is regulated by both the local mechanical environment and the local vascularity. Results show a significant effect of the morphology of the new capillary network on bone formation and heterogeneous distribution of cells similar to those seen in histological studies.

Keywords—Angiogenesis, blood vessel formation, capillary network, tissue differentiation, mechano-biology, mechanobiology.

1 INTRODUCTION

During the last decade, the effect of mechanical loads on bone formation and bone repair has attracted much attention. Several mechano-regulation theories have been proposed relating mechanical loads and tissue differentiation [1, 2, 3] which have been implemented in computer models to successfully reproduce some of the main aspects of the bone regeneration process [4, 5, 6, 7, 8].

However, none of the above mentioned models have considered angiogenesis. This refers to the formation of new capillaries from pre-existing vessels and plays a fundamental role during bone formation. Blood vessels supply oxygen and nutrients which are essential for the proliferation and survival of cells of high metabolic demand, such as osteoblasts. Osteogenic precursor cells in regions of poor vascularity have been shown to follow a chondrogenic rather than an osteogenic pathway [9] and inhibition of blood vessel growth has been shown to inhibit fracture healing in rats [10].

Genot et al. [11] took account of this process in a mechanical model for tissue differentiation to simulate fracture healing. They used endothelial cell density as a continuous variable that evolved according to a set of reaction-diffusion equations. Although their work represents the only attempt to include angiogenesis in a

mechano-biological model, their approach does not provide detailed information of the morphology of the capillary network. This may be important because oxygen diffusion is limited to 100–200 μm from capillary vessels [12] and therefore, the organization of the capillaries in the regenerating tissue may play a fundamental role during tissue differentiation.

The objective of this work was to develop a mechano-biological model for blood vessel growth and tissue differentiation to examine the influence of capillary network formation during tissue repair. We hypothesized that capillary network dynamics influences the regeneration of bone. We tested this hypothesis in a model of a bone repair interface under shear. If the hypothesis is confirmed, it would suggest that mechano-biological simulations of tissue differentiation could be improved by considering the effect of oxygen supply by the capillary network.

2 MATERIALS AND METHODS

The computer simulation combines a stochastic model for cell migration and proliferation [13] with a mechano-regulation theory for tissue differentiation [1]. A lattice model was used for the simulation of cell activity [7] while the local mechanical environment was determined using finite element analysis. A regular lattice was superimposed to the finite element model such that 1000 lattice points were contained inside each element (Fig. 1). Each lattice point represented a possible position for a cell to occupy, with a distance between lattice points of 50 μm . In this lattice, endothelial cells (ECs) formed the new capillary network while, simultaneously, mesenchymal stem cells (MSCs), osteoblasts, chondrocytes and neurons migrated, proliferated and differentiated based on the local mechanical environment and the local vascularity.

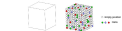


Figure 1. Lattice contained inside each finite element for the simulation of cell activity. Each lattice point represents a possible position for a cell to occupy.

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J. Vander Sloten, P. Verdonck, M. Nysen, J. Hauelsen (Eds.):
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- Using evolvable genetic cellular automata to model breast cancer

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ORIGINAL PAPER

Using evolvable genetic cellular automata to model breast cancer

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Abstract Cancer is an evolutionary process. Mutated cells undergo selection for abnormal growth and survival creating a tumor. We model this process with cellular automata that use a simplified genetic regulatory network simulation to control cell behavior and predict cancer etiology. Our genetic model gives us the ability to relate genetic mutation to cancerous outcomes. The simulation uses known histological morphology, cell types, and stochastic behavior to specifically model ductal carcinoma in situ (DCIS), a common form of non-invasive breast cancer. Using this model we examine the effects of hereditary predispositions on DCIS incidence and aggressiveness. Results show that we are able to reproduce in vivo pathological features to hereditary forms of breast cancer: earlier incidence and increased aggressiveness. We also show that a contributing factor to the different pathology of hereditary breast cancer results from the ability of progenitor cells to pass cancerous mutations on to offspring.

Keywords Genetic cellular automata · DCIS · Progenitor hierarchy · Ductal simulation · Hereditary genetic predisposition · Hereditary breast cancer

1 Introduction

One in eight women will be diagnosed with breast cancer in their lifetime [21]. Carcinogenesis is an evolutionary phenomenon known to result from genetic mutations effecting cellular reproduction and survival [4, 12]. Breast cancer cells that are able to abnormally reproduce and survive, undergo a selection process that

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Using evolvable genetic cellular automata to model breast cancer

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Cancer Stem Cell Tumor Model Reveals Invasive Morphology and Increased Phenotypical Heterogeneity

Integrated Systems and Technologies: Mathematical Oncology

Cancer Stem Cell Tumor Model Reveals Invasive Morphology and Increased Phenotypical Heterogeneity

Andrea Sottoriva¹, Joost J.C. Verhoeff², Tijana Borovski³, Shannon K. McWeeney^{3,4}, Lev Naumov⁵, Jan Paul Medema⁶, Peter M.A. Slood⁷, and Louis Vermeulen¹

Abstract

The recently developed concept of cancer stem cells (CSC) sheds new light on various aspects of tumor growth and progression. Here we present a mathematical model of malignancies to investigate how a hierarchical organized cancer cell population affects the fundamental properties of solid malignancies. We establish that tumors modeled as a CSC contain more functionally versatile tumor subpopulations and show invasive behavior, whereas tumors without a CSC hierarchy do not. These findings are corroborated by *in vivo* studies. In addition, we provide evidence that the CSC model is accompanied by highly altered evolutionary dynamics compared with the ones predicted by *in vitro* *in vivo* models, underscoring the tumor model. Our results indicate that the CSC model allows for significantly higher tumor heterogeneity, which may affect therapy outcome. Moreover, we show that therapy which fails to target the CSC population is not only unsuccessful in curing the patient, but also promotes malignant features in the recurring tumor. These include rapid expansion, increased invasion, and enhanced heterogeneity. *Cancer Res* 70(1): 4656-4666, 2010.

Introduction

Malignancies arise after sequential accumulations of mutations in oncogenes and tumor suppressor genes (1). During the process of malignant transition, fundamental regulatory mechanisms are lost. The cancerous cell population has unlimited growth potential, evades the immune system and apoptosis, and often acquires the ability to breach tissue boundaries and expand into foreign environments (1). The cancer stem cell (CSC) concept sheds new light on all of these features (2, 3). In this study, we investigate the influence of a hierarchical organization of malignant cells on tumor growth, evolution, invasion, and metastasis (4) using a multi-scale cellular automaton-based computer model (5).

CSCs. Malignancies are highly heterogeneous tissues containing largely diverse cancer cell populations as well as cancer

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Major Findings

This research shows that the hierarchical organization of malignant cells, as advocated in the CSC concept, has major implications for tumor biology. CSC-driven tumor growth intrinsically accelerates tumor invasion, influences clonal selection, and has crucial consequences for the development of successful cancer therapies.

cells such as fibroblasts and macrophages (1). The dominant genetic view of malignancies explains the heterogeneity in cancer cells with the presence of genetically diverse cells emerging from a highly heterogeneous population of clonal genetic lesions by cancer cells, each arising separately for reasons, resulting in highly diverse genetic lesions on tumor formation (1). In this article, we refer to the view of malignancies as the classical model. Although this model greatly contributes to our understanding of malignancies, recent experimental evidence suggests an additional layer of complexity. The heterogeneity present in tumor cells, as well as the diversity in differentiation grade of genetically identical cells (6, 7), for example in glioblastoma multiforme, cells with an identical phenotype originating the cell surface marker CD133 and have the capacity to self-renew, differentiate, and transplant the malignancy into severe combined immunodeficiency mice (7). Both numerical and size of differentiated cells are features shared with normal stem cells, and therefore, cells with such features in malignancies are defined as CSCs (8, 9).

Cancer Stem Cell Tumor Model Reveals Invasive Morphology and Increased Phenotypical Heterogeneity

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Reaction-diffusion model for pattern formation in E.coli swarming colonies with slime

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Reaction-diffusion model for pattern formation in *E. coli* swarming colonies with slime.

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A new experimental colonial pattern and pattern transition observed in *E. coli* MG1622 swarming cells grown on semi-solid agar are described. We present a reaction-diffusion model that, taking into account the slime generated by these cells and its influence on the bacterial differentiation and motility, reproduces the pattern and especially predicts the observed changes when the colonial substrate motility is hindered. In spite of having used non-hyperplasticized swimming cells, under these experimental conditions *E. coli* MG1622 can very rapidly colonize a surface, with a low branching rate, thanks to a strong flag production and a local increased density of motile, lubricating cells.

PACS numbers: 87.20.Hf, 87.20.Hk, 87.20.Hg, 87.20.Hv

I. INTRODUCTION

The macroscopic pattern exhibited by a bacterial colony and its dependence on certain environmental parameters can give us some clues about the coordination, self-organization strategy followed by the community of cells. The biological interest in this interdisciplinary area is in pointing out under controlled laboratory conditions the cooperative mechanisms (intercellular interactions, motility and communication) that these bacteria, which are traditionally considered as solitary life forms, may have developed to adapt to changing environments. It is an extremely fertile and interesting area for close collaboration between physicists and biologists because it helps both to understand the transition from individual (non-colonial, a bacterium) to collective (multi-colonial, a colony) behavior [1].

Bacterial colony pattern formation on semi-solid agar surfaces has been studied extensively by microbiologists and physicists [1, 2, 3, 4, 5, 6, 7, 8]. Through these works undertaken with colonies arising from different species and strains of bacteria, the following consensus has been reached. Bacteria can exist in liquid medium without difficulty but in environments with adverse conditions for the swimming motility, they need to develop mechanisms to become more mobile. On semi-solid agar medium, when the viscosity is high and the motility of short-swimmer cells very low, some may differentiate into very elongated, multiciliated and probably flagellated swimmer cells that can move more [2]. The initiation of swarms differentiation seems to be strictly correlated with physico-chemical factors such as surface contact and oxygen sensing response (cell density sensing mechanism) [9]. Swimming cells have the ability to contract water from the agar and produce a lubrication fluid (slime). The flagella driven motility of swimmer cells together with the extracellular slime helps to overcome the surface friction [3]. After some time signaling swimmer cells have

been observed to cease movement, separate and produce groups of swimmer cells (de-differentiation process) [3]. The chemotactic system is essential for swimming motility in *E. coli* [6]. With the chemotactic mechanism bacteria can orient their motility in response to a gradient (positive or negative) of a certain chemical field (a nutrient or a field produced by the bacterial cells such as chemical signals or a pH change) [7]. Based on these experimental observations great effort has been devoted to model the cooperative behavior of a cell community [9, 10, 11, 12, 13].

Here we describe the new pattern and pattern transition observed in a swimming colony of *E. coli* MG1622 [3], a laboratory domesticated wild type strain that produces a lubricating fluid or slime when the substrate motility is hindered this colony expands irregularly with few relatively thick, dense, branched and very rare secondary branches, whereas for conditions of improved colonial motility the pattern is compact and filled with structures of higher cell densities. To study the coordination, self-organized, submillimeter release of this colony and based on experimental microscopic observations, we will define a mathematical model of the cooperative behavior of this colony which is able to predict the pattern transitions when certain control parameters are changed.

This work is organized as follows: in Section II we will introduce the experimental results and our hypotheses for the non-septate reader. Then in Section III, our hypothesis for the collective dynamics will be explained, followed by a mathematical formulation. Results of the computer simulation in comparison with experiments will be posed and discussed in Section IV. The summary and conclusions are given in Section V.

II. EXPERIMENTAL OBSERVATIONS

The development of an *E. coli* MG1622 colony on a semi-solid surface (0.2% "Difco" Agar) under certain nutritional (1% "Difco" Tryptone, 0.2% "Difco" yeast extract, 0.2% NaCl and 0.2% D(+)-fructose) and environmental (at 37°C and an average 22% relative humidity) conditions has been studied in detail with regard and

Reaction-diffusion model for pattern formation in E.coli swarming colonies with slime

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Mapping the phase diagram of the writhe of DNA nanocircles using atomistic molecular dynamics simulations

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Mapping the phase diagram of the writhe of DNA nanocircles using atomistic molecular dynamics simulations

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ABSTRACT

We have investigated the effects of duplex length, sequence, salt concentration and superhelical density on the conformation of DNA nanocircles containing up to 178 base pairs using atomistic molecular dynamics simulation. These calculations reveal that the partitioning of twist and writhe is governed by a delicate balance of competing energetic terms. We have identified conditions which favour circular, positively or negatively writhe and denatured DNA conformations. Our simulations show that AT-rich DNA is more prone to denaturation when subjected to torsional stress than the corresponding GC containing circles. In contrast to the behaviour expected for a simple elastic rod, there is a distinct asymmetry in the behaviour of over and under-wound DNA nanocircles. The most biologically relevant negatively writhe state is more elastic than the corresponding positively writhe conformation, and is only observed for larger circles under conditions of high electrostatic screening. The simulation results have been summarised by plotting a phase diagram describing the various conformational states of nanocircles over the range of circle sizes and experimental conditions explored during the study. The changes in DNA structure that accompany supercoiling suggest a number of mechanisms whereby changes in DNA topology *in vivo* might be used to influence gene expression.

INTRODUCTION

DNA topology *in vivo* is extremely diverse. While regions of the genetic material must be accessible during transcription, replication and repair, the bulk must be

compact to fit within the nucleus. In bacteria, circular plasmids are condensed by supercoiling the DNA into highly writhe superhelical structures. The importance of higher order DNA structures is illustrated by the elaborate machinery for the regulation of DNA topology that exists within the cell. Examples of topoisomerase and gyrase enzymes alter the level of supercoiling by transiently introducing single or double strand breaks and changing the number of times the two strands of the duplex are wrapped around each other (1). Variation in DNA topology influences promoter activity and is consequently involved in the regulation of gene expression and replication (2). For example, the shock response of bacterial cells placed under environmental pressure (such as starvation or thermal stress) is accompanied by dramatic changes in supercoiling, which enable the cell to rapidly modify transcription levels across the whole genome, whereas adaptation through genetic mutations can only occur over a much longer timescale (3,4).

Closed circular DNA can form superhelical structures whenever the topological quantity known as the linking number Lk (5) deviates from its value as a topologically relaxed duplex Lk_0 . The quantity Lk_0 is simply the number of base pairs (bp) in the circle divided by the helical repeat. The linking number is constrained to be integer in closed circular structures and cannot be altered without cutting either one or both DNA strands. The linking number difference $\Delta Lk = Lk - Lk_0$ is commonly normalised to the size of the circle and expressed as the superhelical density σ where:

$$\sigma = \frac{\Delta Lk}{Lk_0} \quad 1$$

The topological property, the linking number, is related to two geometrical parameters of the duplex; the helical twist (T) and the writhe (W), where the writhe is a measure of the contortion of the DNA axis.

$$Lk = Tw + Wr \quad 2$$

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Mapping the phase diagram of the writhe of DNA nanocircles using atomistic molecular dynamics simulations

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How the chromatin fiber deals with topological constraints

PHYSICAL REVIEW E 71, 031910 (2005)

How the chromatin fiber deals with topological constraints

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In the nuclei of eukaryotic cells, DNA is packaged through several levels of compaction in an orderly, reversible way that enables the correct regulation of gene expression. The functional dynamics of this assembly involves the unwinding of the so-called 30-nm chromatin fiber and accordingly requires strong topological constraints. We present a general method for computing both the twist and the writhe of any winding pattern. An explicit derivation is implemented for the chromatin fiber which provides the linking number of DNA in eukaryotic chromosomes. We show that there exists one and only one unwinding path which satisfies both topological and mechanical constraints that DNA has to deal with during decondensation/condensation processes.

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PACS number(s): 87.16.50

In the nuclei of higher eukaryotes, e.g., animals or plants, meters of DNA are packaged by means of proteins into a nucleus of a few micrometers diameter, providing an extreme level of compaction. Coding sequences (genes) are therefore dispersed in a mass of folded DNA and proteins (chromatin) and should be retrieved at will in order to enable a correct genetic expression and therefore the cell survival. This leads to the need for an orderly and dynamically retrievable structure, which is actually achieved by means of a chromatin partition into functional domains, each containing one or a group of genes. In each domain, DNA is folded in a hierarchical structure, including several winding levels. It is first wrapped around spools of proteins that forming a "beads on a string" assembly, which is in turn folded into a 30-nm chromatin fiber. This fiber is further organized into a three-dimensional cross-linked network [1]. In this network, two neighboring nodes are connected by a chromatin fiber loop whose typical length is about 5000 base pairs (bp). In order to provide the transcription machinery with an access to specific genomic regions, the corresponding loop has to be actively decondensed, via a reversible unwinding process that elongates the fiber [2]. The dynamics of this process involves strong mechanical and topological constraints, the former due to DNA elasticity [3], the latter due to the conservation of the linking number Lk of DNA [4] in a loop during the unwinding. Of course, topological constraints could be released by the intervention of topoisomerases, but it has been shown *in vivo* that chromatin decondensation could take place even when these enzymes were inhibited [5]. Moreover, the classic experiment on *in vitro* λ 40 SV40 minichromosomes clearly demonstrates that the linking number is unaffected by the decondensation process *in vivo* [6].

In this paper, we address the issue of how evolution has dealt with the extremely difficult problem of finding an efficient winding pattern fulfilling both mechanical and topological constraints at a time. To answer this question, we start

by giving an analytical formula for the linking number of DNA in a generic bare fiber. We show that despite the fact that Lk is known to be a nonextensive quantity, it is yet possible to express it in terms of the mean linking number of the constitutive elements of the beads on a string assembly. This allows us to set about an exhaustive numerical exploration of Lk for all the possible fiber configurations. By analyzing the results of this exploration, we are able to infer the existence and the uniqueness of a relevant winding unwinding path satisfying all the constraints. We show furthermore that this engineering problem is solved at the local level by the design of the constitutive elements of the fiber. In a typical chromatin fiber, the beads, called nucleosome core particles (NCP), are spaced at intervals of ~ 200 bp. Each NCP contains ~ 150 DNA bp forming 12 turns of a left-handed superhelix [7] and two neighboring beads are connected by ~ 50 bp stretches of DNA, called linkers. The unit of a NCP and a linker is called a nucleosome. The number of DNA bp in a nucleosome is known as the repeat length. In order to describe the structural polydispersity of a chromatin fiber loop, we use the original two-angle model (Fig. 1) introduced by Windrock *et al.* [8]. In this model, when the linkers are assumed to be straight, the geometry of a fiber made up of N nucleosomes can be characterized by two sets of angles α_i and β_i ($i = 1, \dots, N$), specific to the

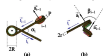


FIG. 1. Color online Schematic of the DNA winding pattern along two neighboring nucleosomes in the two-angle model. (a) View along the NCP axis z_0 . (b) View along the linker direction C_i . The angle α_i is the tilted angle (\vec{C}_i, \vec{z}_0) and β_i is the tilted angle $\arg(\vec{C}_i, \vec{z}_0)$ standing for the twist (twice) of the of the DNA linker. We also indicate the DNA radius $r = 1.0$ nm, NCP radius $R = 5.5$ nm, and NCP pitch $p = 2.0$ nm.

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● Development of regular cellular spacing in the retina: theoretical models

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Development of regular cellular spacing in the retina: theoretical models

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During development of the nervous system, neurons should be appropriately positioned to enable them to make the right functional contacts. Neurons do not immediately migrate to their correct location, but instead regular arrangements gradually emerge from randomly arranged cell populations. This phenomenon has been studied often in the retina, due to its relatively simple layered organisation. In this review, I highlight the principal mechanisms that are thought to be involved, and how mathematical modelling has helped to further our understanding of the role of these processes upon mosaic formation. Three developmental mechanisms are studied in detail, namely, lateral migration, cell fate and cell death. As a case study, I then consider which mechanisms might be involved in the formation of retinal ganglion cell mosaics.

1. Nature and formation of retinal mosaics

The retina is assembled as a stack of several cell layers, with the photoreceptors in the outermost layer and the retinal ganglion cells (RGCs) in the innermost layer (Fig. 1). Photoreceptors convert light into electrical activity which is then modulated by the interacting vertical and horizontal neural pathways within the retina. Once the activity reaches the RGCs, it is converted into spike trains which are sent along the optic nerve and into subcortical and cortical areas in the brain for higher processing. For a comprehensive review of retinal architecture and processing, see Rodieck (1998). The organisation of the retina obeys certain principles which make it relatively easy to study compared to other parts of the nervous system, such as neocortex. For example, there are five major classes of retinal neuron, each with clearly defined morphologies. Each class of retinal neuron tends to be found in one vertical layer, making cell identification easier. Following the terminology of Cook (1998), each class divides into subclasses of neurons which in turn can be classified into individual types of retinal neuron.

At the level of individual cell types, we typically find the spatial organisation of cells within a type to be highly non-random (Fig. 2a). This regular organisation of cell bodies is referred to as a ‘retinal mosaic’ due to the way that the cell body (and its dendrites) tiles the retina. This regular organisation makes sense from the point of view of processing the visual world: receptors are needed uniformly across the retina to ensure that the visual world is regularly sampled and that there are no ‘holes’ in visual space. In this review, I will discuss recent mathematical models that have helped us investigate how these retinal mosaics form during development. I will first review the neurobiological background about the mechanisms that are thought to be involved, and review several types of model, before focusing on a case study comparing how different models may account for the positioning of RGCs.

As well as gaining an insight into the formation of neural patterning in retina, the hope is that we may also learn something about the principles underlying development of the nervous system in general.

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Development of regular cellular spacing in the retina: theoretical models

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