

Cover Page



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## Vascular networks due to dynamically arrested crystalline ordering of elongated cells

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background graphic: ordering in networks formed with elongated cells and chemotaxis (Figure 2.2B)

### Abstract

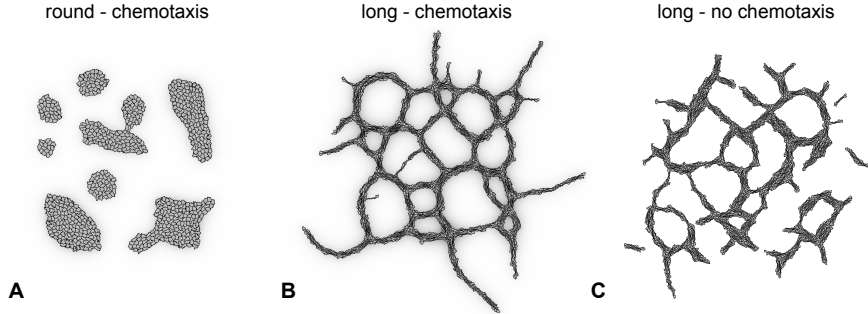
Recent experimental and theoretical studies suggest that crystallization and glass-like solidification are useful analogies for understanding cell ordering in confluent biological tissues. It remains unexplored how cellular ordering contributes to pattern formation during morphogenesis. With a computational model we show that a system of elongated, cohering biological cells can get dynamically arrested in a network pattern. Our model provides a new explanation for the formation of cellular networks in culture systems that exclude intercellular interaction via chemotaxis or mechanical traction.

### 2.1 Introduction

By aligning locally with one another, cells of elongated shape form ordered, crystalline configurations in cell cultures of, e.g. fibroblasts [72, 73], mesenchymal stem cells [73], and endothelial cells [74]. Initially the cells form small clusters of aligned cells; the clusters then grow and the range over which cells align increases with time [73, 75]. To study the emergence of such crystalline cellular ordering, it is useful to make an analogy with liquid crystals [73]. For example, a “cellular temperature” can be defined to describe the cell-type specific persistence (low cellular temperature) or randomness (high cellular temperature) of cell motility, where cells of high cellular temperature (e.g., fibroblasts) are less likely to form crystalline configurations than cells of low temperature (e.g., mesenchymal stem cells) [73]. It was similarly proposed that collective cell motion in crowded cell sheets can be understood as system approaching a glass transition [76, 77]. Although these studies provide useful insights into the ordering of cells in confluent cell layers, it remains unexplored how crystallization and glass-like dynamics contribute to the formation of more complex shapes and patterns during biological morphogenesis.

Cells’ organizing into network-like structures, as it occurs for example during blood vessel development, is a suitable system to study how cellular ordering participates in pattern formation. In cell cultures after stimulation by growth factors (VEGFs, FGFs), endothelial cells elongate and form vascular-like network structures [78–80]. The mechanisms that drive the aggregation of endothelial cells and their subsequent organization into network is a subject of debate. Most models assume an attractive force between cells, either due to chemotaxis [19, 20, 25–29, 49, 81] or due to mechanical traction via the extracellular matrix [21–24, 82, 83]. *In vitro* experiments show that astroglia-related rat C6 cells and muscle-related C212 cells can form network-like structures on a rigid culture substrate [30], which excludes formation of mechanical or chemical attraction between cells. Therefore a sec-

ond class of explanations proposed that cells form networks by adhering better to locally elongated configurations of cells [30] or elongated cells [31]. Here we show that, in absence of mechanical or chemical fields such mechanisms are unnecessary: elongated cells organize into network structures if they move and rotate randomly, and adhere to adjacent cells. As the cells align locally with one another, a network pattern appears. Additional, long-range cell-cell attraction mechanisms, e.g., chemotaxis or mechanotaxis, act to stabilize the pattern and fix its wave length.



**Figure 2.1:** Effect of chemotaxis and cell shape on pattern formation. **A** round, chemotacting, and adhesive cells (10,000 MCS), **B** elongated, chemotacting and adhesive cells (10,000 MCS), and **C** elongated, non-chemotacting and adhesive cells (250,000 MCS). In all panels 700 cells are seeded on the center 500x500 pixels of an 800x800 lattice.

## 2.2 Model description

To model the collective movement of elongated cells, we use the cellular Potts method (CPM), also known as the Glazier-Graner-Hogeweg model [84, 85], a lattice-based, Monte-Carlo model that has been used to model developmental mechanisms including somitogenesis [86, 87], convergent extension [88] and fruit fly retinal patterning [89]. The CPM represents cells as connected patches of lattice sites with identical spin  $\sigma \in \mathbb{N}$ ; lattice sites with spin  $\sigma = 0$  represent the extracellular matrix (ECM). To simulate stochastic cell motility, the CPM iteratively displaces cell-cell and cell-ECM boundaries by attempting to copy the spin of a randomly selected site into a randomly selected adjacent lattice site  $\vec{x}$ , monitoring the resulting change  $\Delta H$  of a Hamiltonian,

$$H = \sum_{(\vec{x}, \vec{x}')} J(\sigma(\vec{x}), \sigma(\vec{x}')) (1 - \delta(\sigma(\vec{x}), \sigma(\vec{x}'))) + \sum_{\sigma} \lambda_A (a(\sigma) - A)^2 + \sum_{\sigma} \lambda_L (l(\sigma) - L)^2.$$

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A copy attempt will always be accepted if  $\Delta H \leq 0$ , if  $\Delta H > 0$  a copy attempt is accepted with the Boltzmann probability  $P(\Delta H) = \exp(-\Delta H/\mu(\sigma))$ , with  $\mu(\sigma)$  a “cellular temperature” to simulate cell-autonomous random motility. For simplicity, we here assume that all cells have identical temperature  $\mu$ . The time unit is a Monte Carlo step (MCS), which corresponds with as many copy attempts as there are lattice sites.

The first term of Equation 2.2 defines an adhesion energy, with  $(\vec{x}, \vec{x}')$  a pair of adjacent lattice sites and the Kronecker delta returning a value of 1 for pairs at cell-cell and cell-ECM interfaces, or zero otherwise. In the model two contact energies are defined:  $J_{\text{cell,cell}}$  for  $\sigma > 0$  at both lattice sites, and  $J_{\text{cell,ECM}}$  for  $\sigma = 0$  at one lattice site. The second and third term are shape constraints that penalize deviations from a target shape, with  $A$  and  $L$  a target area and length, and  $a(\sigma)$  and  $l(\sigma)$  the current area and length of the cell;  $\lambda_A$  and  $\lambda_L$  are shape parameters. We efficiently estimate  $l(\sigma)$  by keeping track of a cellular inertia tensor as previously described [25].

In a subset of simulations, we further assume that cells secrete a diffusing chemoattractant  $c$ , which we describe with a partial differential equation:

$$\frac{\partial c(\vec{x}, t)}{\partial t} = D\nabla^2 c(\vec{x}, t) + s(1 - \delta(\sigma(\vec{x}), 0)) - \epsilon \delta(\sigma(\vec{x}), 0),$$

with diffusion constant  $D$ , secretion rate  $s$  and decay rate  $\epsilon$ . After each MCS, a forward Euler method solves Equation 2.2 for 15 steps with  $\Delta t = 2$  s and zero boundary conditions. To model the cells’ chemotaxis up concentration gradients of the chemoattractant, during each copy attempt from  $\vec{x}$  to  $\vec{x}'$  we increase  $\Delta H$  with a  $\Delta H_{\text{chemotaxis}} = \lambda_c (c(\vec{x}) - c(\vec{x}'))$ , with  $\lambda_c$  a chemotactic sensitivity [90].

We use the following parameter settings, unless specified otherwise:  $\mu = 1$ ,  $J_{\text{cell,cell}} = .5$ ,  $J_{\text{cell,ECM}} = .35$ ,  $\lambda_A = 1$ ,  $\lambda_L = .1$ ;  $\lambda_c = 10$ ,  $A = 100$  l.u.<sup>2</sup>,  $L = 60$  l.u.;  $D = 10^{-13}$  m<sup>2</sup>s<sup>-1</sup>,  $\epsilon = 1.8 \cdot 10^{-4}$  s<sup>-1</sup>,  $s = 1.8 \cdot 10^{-4}$  s<sup>-1</sup>; where l.u. denotes a lattice unit which corresponds with 2  $\mu\text{m}$ . Adjacent lattice sites  $\vec{x}'$  are defined as the eight nearest neighbors of lattice site  $\vec{x}$ . Unless stated otherwise, a simulation is initialized with 175 cells randomly distributed on a 220x220 area at the center of a 400x400 lattice.

### 2.3 Results

As Figure 2.1 shows, and in agreement with previous reports [25], if we allow for chemotaxis, rounded cells accumulate into rounded clusters (Figure 2.1A) and elongated cells aggregate into networks (Figure 2.1B). Interestingly, however, chemotaxis is not required for network formation: cell-cell adhesion between elongated cells suffices for forming networks (Figure 2.1B). Movies corresponding with Figure 2.1B and C (See Supplemental Movie S1, model without chemotaxis, and Movie S2, model with chemotaxis) suggest that the

gradual alignment of cells with their neighbors is key to network formation and network evolution. To characterize this cell alignment, we define  $\theta(\vec{x}, r)$  as the angle between the direction of the long axis  $\vec{v}(\sigma(\vec{x}))$  of the cell at  $\vec{x}$ , and a local director  $\vec{n}(\vec{x}, r)$ , a weighted local average of cell orientations defined at radius  $r$  around  $\vec{x}$ :

$$\vec{n}(\vec{x}, r) = \langle \vec{v}(\sigma(\vec{y})) \rangle_{\{\vec{y} \in \mathbb{Z}^2: |\vec{x} - \vec{y}| < r\}}.$$

Figure 2.2A and B depict the value of  $\theta(\vec{x}, 3)$  for simulations without chemotaxis (Figure 2.2A) and with chemotaxis (Figure 2.2B), with dark gray values indicating values of  $\theta(\vec{x}, 3) \rightarrow \pi/2$ . Network branches are separated by large values of  $\theta(\vec{x}, 3)$ , indicating that within branches cells are aligned, whereas branch points are “lattice defects” in which cells with different orientations meet.

Supplemental Movies S3 and S4 ( $\theta(\vec{x}, 3)$ ) show how the cells align gradually over time in the absence and presence of chemotaxis. To characterize the temporal development of cell alignment in more detail, we use an orientational order parameter [91]:

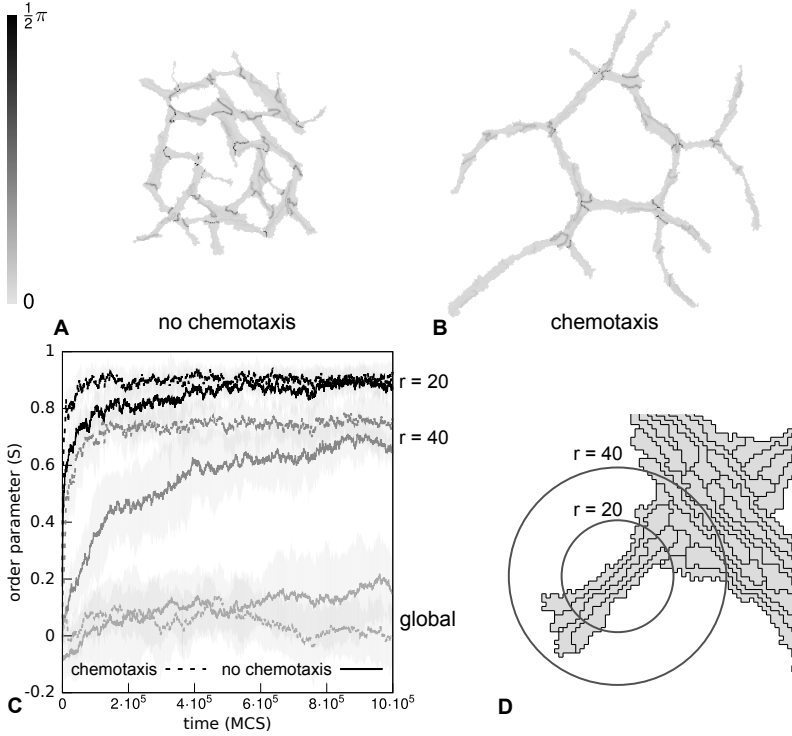
$$S(r) = \left\langle \cos(2\theta(\vec{X}(\sigma), r)) \right\rangle_{\sigma}$$

with  $\vec{X}(\sigma)$  the center of mass of cell  $\sigma$ .  $S$  ranges from 0 for randomly oriented cells to 1 for cells oriented in parallel.

Figure 2.2C shows the evolution of the global orientational order parameter  $\lim_{r \rightarrow \infty} S(r)$  and of the local orientational order parameters  $S(20)$  and  $S(40)$ . Both with chemotaxis (dashed lines) and without (solid lines),  $S(20)$  grows more quickly and reaches higher ordering than  $S(40)$ . The reason for this is that in cells of length 50 – 60 l.u.,  $S(20)$  (covering cells up to a radius  $r = 20$  from the cell’s center of mass) only detects lateral alignment of cells, whereas a radius  $S(40)$  also detects linear line-up of cells (Figure 2.2D). Thus cell-cell adhesion of long cells quickly aligns cells with the left and right neighbors, while it aligns them more slowly with those in front and behind. This results in networks with short branches of aligned cells. Interestingly, chemotaxis aligns cells more rapidly, both along the short and long sides of cells, resulting in networks with much longer branches than with adhesion alone.

Next we analyze the mechanisms that drive the orientational ordering in the cell networks. Visual inspection of the simulation movies suggests that single cells move and rotate much more rapidly than locally aligned clusters of cells. A network of locally aligned cells forms rapidly from initially dispersed cells. Merging of branches seems to be a much slower process, and potentially prevents a further evolution to global nematic order. To quantify these observations we measured the translational and rotational diffusion coefficients of cells as a function of the size of the network branch to which it

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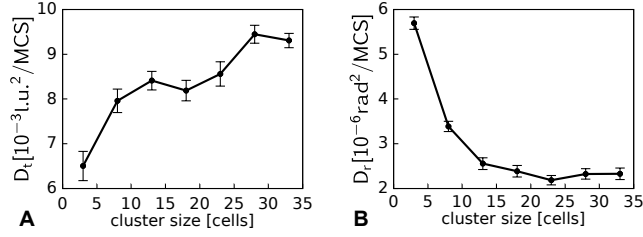


**Figure 2.2:** Crystalline cell ordering during network formation. **A-B**  $\theta(\vec{x}, r)$  with  $r = 3$  for a simulation with chemotaxis (**A**) and without chemotaxis (**B**) after 25,000 MCS. **C** Temporal evolution of orientational order parameter  $S(r)$  for  $r = 20$  (black curves),  $r = 40$  (gray curves) and  $r \rightarrow \infty$  (light gray) without chemotaxis (solid) and with chemotaxis (dashed). Order parameter is averaged over 10 simulation repeats (gray shadows represent standard deviation). **D** cells included in the order parameter with  $r = 20$  and  $r = 40$ .

belongs. We loosely define a network branch, or *cluster* of aligned cells as a connected set of at least two cells with relative orientations  $< 5^\circ$ , *i.e.*, in Figure 2.2A and B dark gray values separate the clusters. To detect clusters computationally, we first identify the connected sets for which  $\theta(\vec{x}, 3) \leq 5^\circ$ , which are surrounded by lattice sites of  $\sigma = 0$  or sites with  $\theta(\vec{x}, 3) > 5^\circ$ . We then eliminate connected sets of fewer than fifty lattice sites. The CPM cells sharing at least 50% of their lattice sites with one of the remaining sets form a cluster. The translational diffusion coefficient,  $D_t$ , derives from the mean square displacement (MSD) of a set of cells:

$$\left\langle |\vec{X}(\sigma, t) - \vec{X}(\sigma, 0)|^2 \right\rangle_\sigma = 4D_t t.$$

Similarly, the rotational diffusion coefficient,  $D_r$ , derives from the mean square



**Figure 2.3:** Relation between cluster size and cell displacement. Clusters are calculated for each morphology between 500 and 25,000 MCS (100 simulation repeats), with an interval of 500 MCS; see text for details. The error bars represent the standard error of the linear fits used to estimate diffusion coefficients.

rotation (MSR) of a set of cells:

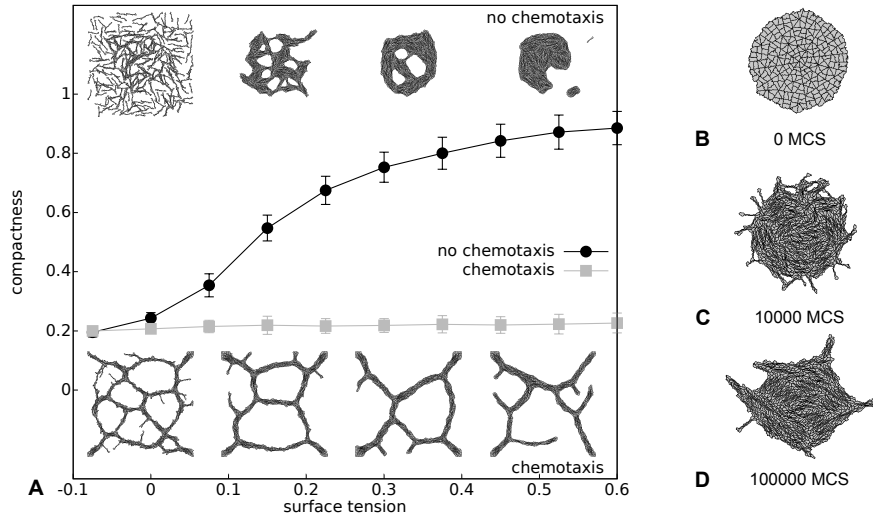
$$\langle (\alpha(\sigma, t) - \alpha(\sigma, 0))^2 \rangle_\sigma = 2D_r t,$$

with  $\alpha(\sigma, t) - \alpha(\sigma, 0)$  the angular displacement of a cell between time 0 and  $t$ . During a simulation, cells may move between clusters, and clusters can merge. Therefore, to calculate  $D_t$  and  $D_r$  of cells as a function of cluster size, for 100 simulations of 250,000 MCS we measured trajectories of each individual cell with one data point per 500 MCS, and kept track of the size of the cluster it was classified into at each time point. We defined cluster size bins, with the first bin collecting all clusters consisting of two to five cells, and the next bins running from 6 to 10, 11 to 15, etc. We split up the trajectories into chunks of 10 consecutive data points, during which the cells stayed within clusters belonging to one bin. To calculate  $D_t$  and  $D_r$  we performed a least square fitting on the binned MSD and MSR values for these trajectory chunks.

The translational diffusion,  $D_t$ , increases slightly with cluster size (Figure 2.3A). This may reflect that the probability of hopping between small clusters will be larger than the probability of hopping between larger clusters, resulting in an overrepresentation of slow cells in the small clusters. Interestingly, the rotational diffusion  $D_r$  drops with the cluster size (Figure 2.3B), indicating that cells in large clusters rotate more slowly. These results suggest that the rotation of cells in big clusters is limited, which reduces the probability that two clusters rotate and merge into a single larger cluster. Therefore, if the size of clusters increases, their rotation speeds drop as does the probability of cluster fusion. Thus, although further alignment of clusters would reduce the pattern energy  $H$  (Equation 2.2), the pattern evolution essentially freezes.

To corroborate our hypothesis that network patterns are transient patterns that increasingly slowly evolve towards nematic order, we looked for model parameters that could speed up pattern evolution. Figure 2.4A shows the ef-

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**Figure 2.4:** A Relation between compactness and surface tension, with and without chemotaxis. The compactness was calculated at 100,000 MCS and averaged over 100 simulations (error bars represent standard deviation). Simulations were initialized with 350 cells on 260x260 path on the center of a 420x420 lattice. **B-D** evolution of a simulation initialized with a 128-cell blob on the center of a 420x420 grid.

fect of surface tension ( $\gamma_{\text{cell,ECM}}$ ) on the ability of cells to form networks after 100,000 MCS, as expressed by the configuration's  $compactness C = \frac{A_{\text{cells}}}{A_{\text{hull}}}$ , where  $A_{\text{hull}}$  is the area of the convex hull of the largest connected group of cells, and  $A_{\text{cells}}$  is the summed area of the cells inside the hull. A value of  $C \rightarrow 1$  indicates a spheroid of cells, where for networks  $C$  would tend to zero. For values of  $\gamma_{\text{cell,ECM}} = J_{\text{cell,ECM}} - \frac{J_{\text{cell,cell}}}{2} > 0$ , the equilibrium pattern should minimize its surface area with the ECM. Indeed at increased surface tensions the cells settle down in spheroids or networks with only few meshes, although they initially still form network-like patterns (see Supplemental Movie S5, model without chemotaxis and  $\gamma_{\text{cell,ECM}} = 0.3$ ). To confirm that also for  $\gamma_{\text{cell,ECM}} = 0.1$  (i.e., the values used in Figs. 1-3) spheroids are stable configurations, we initialized our model with a spheroid (Figure 2.4B). Although initially some cells sprout (Figure 2.4C) from the spheroid due to their elongation, they then align gradually and the cell cluster remains spherical. No network formation was detected in simulations of 100,000 MCS (Figure 2.4D), suggesting that spheroids represent the global minimum of the Hamiltonian. Interestingly, in presence of chemotaxis networks form for a wide range of surface tensions (inset Figure 2.4A and [25]).

## 2.4 Discussion

Our analysis suggests that in the cellular Potts model elongated, adhesive cells can form networks in a parameter regime where a spheroid pattern is the minimal energy state. The cells initially align with nearby cells, thus forming the branches of the network. In order for the pattern to evolve further towards the minimal-energy spheroid pattern, the locally aligned clusters of cells must join adjacent branches, for which they must move and rotate. Our analysis of the rotational and translational diffusion of cells in Figure 2.3 shows that this becomes more difficult for cells belonging to larger clusters. Thus the networks evolve ever more slowly to the minimal energy state, and gets dynamically arrested in a network-like configuration, a phenomenon reminiscent of the glass transition, as e.g. observed in attractive colloid systems [92], collective cell migration of biological cells *in vitro* [76, 77], and colloid rod suspensions [93] in which gels can form from clusters of parallel rods [94–96].

Figure 2.4A suggests that the cellular Potts simulations undergo a glass transition as the surface tension drops: for high surface tension the system evolves towards equilibrium, for lower surface tensions the system becomes jammed in a network-like state. Thus our model provides a new explanation for the formation of vascular networks in absence of chemical or mechanical, long-range, intercellular attraction [30]. Interestingly, intercellular attraction via chemotaxis stabilizes the formation of networks in our simulations [25] and can drive sprouting from spheroids (not shown). This suggests that networks are an equilibrium pattern of our system in presence of intercellular attraction. Nevertheless the present analysis of arrested dynamics provides new insight into the system with intercellular attraction: chemotaxis reinforces local ordering over a distance proportional to the diffusion length of the chemoattractant producing networks of a scale independent of surface tension [25].

### 2.A Supplementary movies

An archive containing all supplementary movies can be found at

<http://persistent-identifier.org/?identifier=urn:nbn:nl:ui:18-22535>.

**Movie S1** Network formation with elongated cells that chemotact towards a chemoattractant they secrete themselves. This simulation was performed with 700 cells on a 800x800 lattice and ran for 10,000 MCS.

**Movie S2** Network formation with elongation cells that do not chemotact. This simulation was performed with 700 cells on a 800x800 lattice and ran for 250,000 MCS.

**Movie S3** Evolution of the alignment of elongated cells that chemotact to-

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wards a chemoattractant they secrete themselves. This simulation was performed with 175 cells on a 400x400 lattice and ran for 25,000 MCS.

**Movie S4** Evolution of the alignment of elongated cells that do not chemotact. This simulation was performed with 175 cells on a 400x400 lattice and ran for 25,000 MCS.

**Movie S5** Pattern formation with adhesive, elongated cells, without chemotaxis. This simulation was performed with 175 cells on a 400x400 lattice and ran for 100,000 MCS.