Glassy Dynamics in Models of Confluent Tissue with Mitosis and Apoptosis

Michael Czajkowski,¹ Daniel M. Sussman,² M. Cristina Marchetti,³ and M. Lisa Manning²

¹ Physics Department, Georgia Institute of Technology

² Physics Department and Soft & Living Matter Program, Syracuse University

³ Department of Physics, University of California at Santa Barbara

(Dated: March 30, 2019)

Recent work on particle-based models of tissues has suggested that any finite rate of cell division and cell death is sufficient to fluidize an epithelial tissue. At the same time, experimental evidence has indicated the existence of glassy dynamics in some epithelial layers despite continued cell cycling. To address this discrepancy, we quantify the role of cell birth and death on glassy states in confluent tissues using simulations of an active vertex model that includes cell motility, cell division, and cell death. Our simulation data is consistent with a simple ansatz in which the rate of cell cycling and the rate of relaxation of the tissue in the absence of cell cycling contribute independently and additively to the overall rate of cell motion. Specifically, there is one regime where cell cycling dominates the dynamics and the tissue always fluidizes, and a second glassy regime where the tissue exhibits caging behavior with subdiffusive cell displacements.

I. INTRODUCTION

A number of experiments conducted over the past decade [1–3] have indicated the existence of glass-like states in confluent epithelial tissues. Evidence for glassy behavior includes the observation of dynamical heterogeneities characteristic of supercooled liquids [1] and caging effects in motile cell trajectories [3]. In these living tissues, the constituent cells will often undergo mitosis (cell division) and apoptosis (programmed cell death) in a regulated cell life cycle. These cell life events are necessarv for the survival and function of many tissues [4, 5], and they clearly differentiate these dense cellular systems from the molecular and polymeric glasses that are so well studied. Introducing and removing material elements from any system should impact the dynamics, and it is therefore important to examine the effect of cell division and cell death on glassy tissue dynamics.

Recent simulations have suggested that the presence of cell division and death should generically fluidize a tissue [6–9]. Employing a particle-based model, for instance, Matoz-Fernandez et. al. [6] have suggested that even at small rates k_{δ} of cell division and death per cell, one will not observe the sub-diffusive behavior of cell trajectories which we take to be a hallmark of glassy systems. This, therefore, poses a serious question for the experimental and theoretical studies of these systems: why are caging behavior (and other signatures of glassy dynamics) observed at all?

Earlier work by Ranft et. al. [8] at the continuum level considered the effect of cell division and death events on a model of 3-dimensional elastic tissue. Their analytical and numerical findings suggest that cell dynamics will be diffusive and controlled by k_{δ} in the limit of long times and infinitely large tissues. However, both this work and many other previous investigations have focused on cell division and death as the dominant driving force behind tissue dynamics. Epithelial cells can also escape the cages set by their neighbors through motility driven by traction forces on a substrate. Such forces play important roles in

the development and function of organisms [10–14], and even in the absence of cell division and cell death, the timescale over which cells may escape their cages changes by orders of magnitude with variation of propulsive forces and intercellular tensions [14–18]. The full dynamics of tissues which experience motile forces in addition to mitosis and apoptosis should therefore have contributions from each of these sources of internal driving. In particular, it is natural that the observed relaxation time of a tissue will be set by a competition between the timescale of cell cycling events and the timescale of cage escape due to the motile cells encountering mechanical energy barriers.

A complicating factor in previous investigations of model tissues with both active dynamics and cell cycle events is that they have focused on particle-based modes and they often couple the details of the cell cycle events to the current configurational state of the tissue [6–8] (for instance, triggering cell divisions when a particle is surrounded by a sufficient amount of space, or triggering cell deaths when the local density or contact number is too high). While such couplings may be biologically motivated, they also make it difficult to separate out the mechanically fluidizing effects of cell division and death on the glassy behavior of the model tissues.

In this work, we focus on models of confluent tissues, where there are no gaps or overlaps between cells. Such tissues are well-described by so-called "vertex models" that have been used extensively to understand patterning and rigidity in tissues [19–24]. Although there has been substantial recent work on "Voronoi" versions of such models [18, 25], here we introduce an "active vertex model" which naturally incorporates cell motile forces similar to the previously studied Self-Propelled Voronoi (SPV) model, but without the restriction that cell shapes be described by Voronoi volumes.

The remainder of this work is organized as follows. In Sec. II we describe the new Active Vertex model and characterize its dynamical behavior in the absence of cell division and cell death. In Sec. III we extend this model

to include a single controlled rate of mitosis and apoptosis, and we demonstrate that the phase diagram for this model has two dynamical regimes: a fluid-like regime dominated by cell cycle events and a glassy regime dominated by the inherent slow dynamics of the model. These combined regimes are well-described by a simple ansatz in which the overall relaxation time is an additive independent sum of the inherent rate of cell motility and the rate of cell cycling, revealing a universal crossover between cell-cycle-dominated and motility-dominated dynamics. In Sec. IV we investigate the displacement fields associated with individual mitosis and apoptosis events. Each event produces an Eshelby-like displacement field of tissue cells, in addition to a mechanical noise stemming from the disordered geometry of the tissue. Quantifying these displacements allows us to construct a prediction for the cell diffusion across a broad range of cell cycling rates, confirming that the effect of cell cycling on diffusion is similar to that predicted by previous continuum models [6, 8], although the non-affine motions ignored by those models also contribute to overall cell motion. We close in Sec. V with a discussion of how our results compare to previous work, and we suggest future experiments to guide future developments.

II. A VERTEX MODEL FOR MOTILE TISSUES

Numerical studies of the influence of motility on tissue dynamics have been conducted in particulate models [15], the Potts model [26, 27], and Voronoi / vertex models [14, 18, 28, 29]. These investigations have generally indicated that motile forces tend to promote fluidity and enhance diffusion. For instance, using the SPV model – in which cells are described as polygons obtained from a Voronoi tiling of the plane that self-propel analogously with self-propelled particle models – Bi et. al. [18] obtained a transition line separating glassy states from fluid-like states where cells frequently exchange neighbors.

While the Voronoi description is theoretically appealing, the restriction of cell shapes to local Voronoi volumes is not always desirable. More seriously, Voronoi models are not easily extendable to describe tissues with free boundaries, as would be needed to describe woundhealing geometries in 2D or isolated spheroids in 3D. To avoid these issues, while simultaneously preserving the spirit of the self-propelled Voronoi model, we have constructed a model for a motile tissue that preserves the freedom of polygonal cell shapes. In this work we restrict our attention to the generic case where every vertex is 3fold coordinate, although this restriction could be easily relaxed to study the sorts of open boundaries described above. To our knowledge, the version of the vertex model presented herein has not been explored in previous work. Therefore, we include a more detailed description of the model, as well as quantification of its dynamical states in the absence of cell cycle events.

A. Shape forces in the Vertex Model

In the Vertex Model [19–22] cells are modeled as irregular polygons tiling the plane, but in contrast to Voronoi models, the degrees of freedom are the positions of the vertices of the spatial tiling. Previous studies using this model have typically involved searching for geometric states which minimize the tissue energy,

$$E_{Shape} = \sum_{a} E_{a} = \sum_{a} \left[\kappa_{A} (A_{a} - A_{0a})^{2} + \kappa_{P} (P_{a} - P_{0a})^{2} \right],$$
(1)

where the sum runs over all cells a and κ_A and κ_P are elastic moduli. This energy drives the area A_a and perimeter P_a of cell a toward a target area A_{0a} and a target perimeter P_{0a} respectively. The area term in Eq. 1 comes from the resistance to fluctuations in cell height. The perimeter term may be seen as arising from the competition between cell-cell adhesive interactions and the non-linear tension created by the cortical actin network. While the target shape properties may vary from cell to cell, in the absence of cell division and death we presume cells identical with uniform A_0 and P_0 values. It is important to briefly note the importance of the dimensionless target "shape index" $s_0 = P_0/\sqrt{A_0}$, that has proven to be a useful tuning parameter controlling rigidity in both static [23] and motile [18] tissues.

B. The Active Vertex Model

Although the simplest implementation of activity in a Vertex Model would be to allow each vertex to be self-propelled, we would like to model the behavior of a polarized motile cell moving coherently along a specific direction. Therefore, we extend this model to include self propulsion of cell a in the direction

$$\hat{\mathbf{n}}(\theta_a) = \cos(\theta_a)\hat{\mathbf{x}} + \sin(\theta_a)\hat{\mathbf{y}}, \qquad (2)$$

where the cell propulsion angle θ_a is governed by

$$\partial_t \theta_a = \eta_a \,, \tag{3}$$

with η_a being a white gaussian noise defined by

$$\langle \eta_a(t) \rangle = 0 \tag{4}$$

and

$$<\eta_a(t_1)\eta_{a'}(t_2)>=2D_r\delta(t_1-t_2)\delta_{aa'}$$
. (5)

In this model, each cell attempts to carry out a persistent random walk, which is a reasonable assumption as this is commonly observed for cells isolated from the surrounding tissue and in the absence of external signals [30, 31]. Assuming that cell dynamics take place in the overdamped limit, vertex μ will follow an equation of motion,

$$\partial_t \mathbf{r}^{\mu} = -\frac{1}{\gamma} \frac{\partial}{\partial \mathbf{r}^{\mu}} E_{AVM} \,, \tag{6}$$

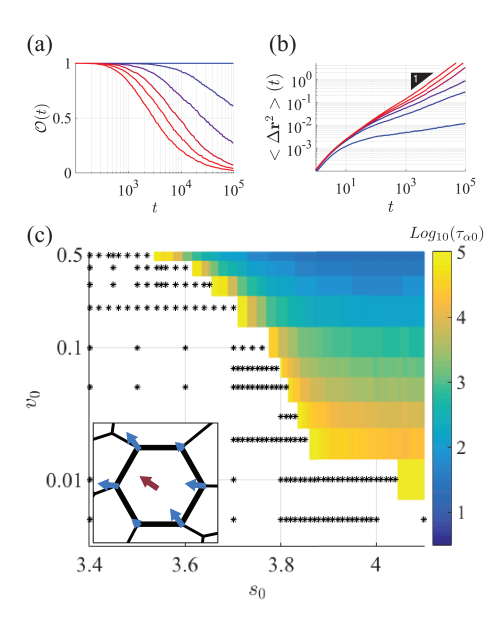


FIG. 1. Quantification of phase behavior in the Active Vertex Model (AVM) with $D_r = 1$ and $N_{cells} = 300$. In (a), the timescale $\tau_{\alpha 0}$ of the decay of the overlap function grows beyond the length of simulations as the vertex model tuning parameter s_0 is decreased while $v_0 = 0.05$ is held fixed. In (b), the mean square displacement of cell centers for the same phase points indicates this increasing timescale is associated with subdiffusive caging behavior. In (c), $\tau_{\alpha 0}$ is shown on a logarithmic colorscale for a representative set of points in the $s_0 - v_0$ plane. Black stars indicate phase points where $\tau_{\alpha 0}$ is too long to be resolved with our data. The inset shows the implementation of motile forces prescribed by Eq. (7).

where γ is the substrate friction. Here and throughout we use superscripted greek letters to refer to vertex positions, and subscripted latin letters to refer to cell positions APPENDIX NEEDS TO REFLECT THIS... if we like it as a solution? The total effective energy,

$$E_{AVM} = E_{Shape} - \gamma v_0 \sum_{cells-a} \hat{\mathbf{n}}(\theta_a) \cdot \mathbf{r}^a, \qquad (7)$$

now captures the cellular self-propulsion forces (of magnitude v_0) in addition to the standard shape energy from Eq. 1. In our AVM simulations we define \mathbf{r}^a as the cell centroid, or geometric center,

$$\mathbf{r}^{a} = \frac{1}{6A_{a}} \sum_{\nu} \left(\mathbf{r}_{a}^{\nu} + \mathbf{r}_{a}^{\nu+1} \right) \left(\mathbf{r}_{a}^{\nu} \times \mathbf{r}_{a}^{\nu+1} \right) \cdot \hat{\mathbf{k}}, \quad (8)$$

where the sum goes counterclockwise over the N_a vertices on cell a and $\mathbf{r}_a^{N_a+1}=\mathbf{r}_a^1$.

While it is uncommon to think of such a self-propulsion force as coming from an energy [32], writing things in this way permits an analogy with the Voronoi model. In the SPV, the degrees of freedom are the Voronoi centers and we might imagine constructing the same energy Eq. 7, using these Voronoi centers as the $\{\mathbf{r}^a\}$. It is easy to see that using these as the degrees of freedom in the overdamped equation of motion,

$$\partial_t \mathbf{r}^a = -\frac{1}{\gamma} \frac{\partial}{\partial \mathbf{r}^a} E$$

$$= v_0 \hat{\mathbf{n}}(\theta_a) - \frac{1}{\gamma} \frac{\partial}{\partial \mathbf{r}^a} E_{shape} ,$$
(9)

indeed leads to the standard SPV dynamics [18, 25, 33]. We are therefore using the closest energetic analog that implements SPV dynamics in a vertex model. In the AVM we will instead apply shape-based and motility-based forces to the vertices. These forces may be precisely defined by taking the derivatives in Eq. 6, as explained in greater depth in Appendix A 1.

C. Glassy dynamics in the Active Vertex Model

Since this model for confluent tissue dynamics has not been explored in any previous work, we first examine its behavior and search for glassy states. We quantify the dynamical state of the system by using two familiar metrics: the mean square displacement and the self-overlap function. The self-overlap function is defined as

$$\mathcal{O}(t) = \frac{1}{N_{cells}} \sum_{a} \Theta \left(b - |\mathbf{r}^{a}(t) - \mathbf{r}^{a}(0)| \right) , \qquad (10)$$

where Θ is the heaviside function and b represents the size of a typical cage in natural units. We set b to 0.5 for the purposes of this work. All lengths are measured in units of $1/\sqrt{\rho}$, where ρ is the system number density. The function \mathcal{O} has a value of 1 at t=0 and will decay toward zero as cells travel past this caging distance b. The structural relaxation time $\tau_{\alpha 0}$ is estimated as the point at which the self-overlap decays below 1/e of its initial value and is an estimate of the uncaging time.

As indicated in the introduction, in this work we use the term "glassy" to primarily refer to the presence of a subdiffusive regime in the mean squared displacements, and to the observation of uncaging times $\tau_{\alpha 0}$ that are orders of magnitude longer than expected from free diffusion. The uncaging time averaged over 10 systems of $N_{cells}=300$ is displayed in Fig. 1. Its behavior is consistent with previous work on the Self-Propelled [18] and Thermal [29] Voronoi models. Less motile tissues at lower s_0 (higher cortical tension) are indeed glassy and have long relaxation times. The tissue may be effectively fluidized by either increasing v_0 (higher effective temperature) or increasing s_0 (higher cell-cell adhesion).

Although the overlap function shown in Fig. 1 appears qualitatively similar to those observed in other glassy

materials, both it and the mean-squared displacement shows an unusual behavior. In most glassy materials, the mean-squared displacement levels off to a subdiffusive regime where $\Delta r^2 \sim t^0$, followed later by a diffusive regime $\Delta r^2 \sim t^1$. Here we find anomalous sub-diffusive behavior with a non-standard exponent, $\Delta r^2 \sim t^\alpha$, where $0 < \alpha < 1$, is found to persist across a wide range of timescales. In this same parameter regime, we also see multiple timescales in the shear stress autocorrelation function. These results are highlighted in Appendix A 2. Such properties are highly unusual in the context of standard glasses, and may be related to the unusual zero-temperature rigidity transition in such models [28]. Understanding the origins of this behavior will be an interesting avenue for future work.

As we are interested in characterizing the simple long-time diffusion of cells in our model, this unusual behavior is problematic. To acknowledge this, throughout the manuscript we highlight any data presented from simulations which remain sub-diffusive on long timescales. In Section III we will see that the parameter regimes of the active vertex model which are diffusive at long times have dynamics which are easily interpreted as a competition between glassy relaxation and relaxation driven by cell division and death.

III. SIMULATION OF MITOSIS AND APOPTOSIS EVENTS

We would now like to incorporate the influence of cell cycle events in these vertex model simulations. To identify an appropriate procedure, we first note that division and death events can change the total number of cells and the cell areal density, with potentially drastic changes in tissue dynamics. To minimize such effects, we choose to work in a constant-number ensemble. In our simple implementation, each instance of apoptosis will be accompanied by a mitosis event somewhere else in the tissue. In addition, the implementation of the individual cell cycle events is chosen to preserve the sum $A_0^{(T)} = \sum_a A_{0a}$ at all times.

In apoptosis, a cell will abruptly contract to a small size and then extrude itself, effectively disappearing from the 2d monolayer [34]. Shown in Fig.2(d-f), our simple realization of apoptosis on cell a begins with setting $A_{0a} \rightarrow 0$ and $P_{0a} \rightarrow 0$. This change induces the rapid contraction of cell a to small size and generally will lead to a final triangular shape. The simulation then detects triangular cells that are smaller than some threshold area A_{min} and deletes them.

Similarly, in the process of mitosis a cell will expand, eventually reaching a threshold size checkpoint and dividing into two complete cells [35–37], all the while maintaining tissue cohesion [38]. While this process typically spans a much longer time period than the event of a cell death, we model this growth process as similarly instantaneous to preserve the constant A_{0T} . Shown in Fig 2,

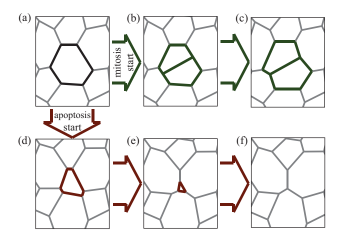


FIG. 2. The process for a single mitosis event in our model is shown sequentially in (a-c). Starting again with the initial cell in (a), the model steps for apoptosis are shown sequentially in (d-f).

in the division of cell a opposite edges of the cell are chosen at random, thus determining the axis of the division. A vertex is then added to the center of each edge and these new vertices are connected with a new edge. Naturally the product is two closed polygonal cells out of one, and the shape parameters of these new cells are then set to the uniform value of the other cells in the tissue. These cells are then allowed to expand dynamically in the simulation. Similar division dynamics were studied in a quasi-static system by others, including Farhadifar et. al. [19].

In practice, such constant number simulations allow us to enforce a global cell cycle event rate k_{Δ} , which is implemented as a Poisson process. This sets the effective cycle rate per cell as $k_{\delta} = k_{\Delta}/N_{cells}$. Note that k_{δ} is a more appropriate parameter for real biological systems whose cell cycle timing does not depend strongly on the extent of the surrounding tissue. We therefore use $k_{\delta} = 1/\tau_{\delta}$ as our tuning parameter.

In simulations with cell division and death, complications arise when trying to extract cell trajectory data. This is because many trajectories will start and end during the course of the simulation. Rather than parse this network of trajectories, we follow the method used in [6] and exclude a small (%10) subset of the cells from the cell cycle events. Using these "tracer" cells to obtain our dynamical data is both convenient and affords the closest comparison with the previous literature [6].

Fast rates of cell division tend to cause tissues to reach rare topological states which introduce very stiff coefficients in the ODEs and are not easily handled by our numerical integrator MICHAEL is checking this. We therefore restrict our search to $\tau_{\delta} > 3000$.

A. Tissue dynamics in the presence of mitosis and apoptosis

With the simulation protocol established, we again search for signatures of glassy behavior. Tissue dynamics is quantified using uncaging time τ_{α} , defined as the time required for the observed overlap to decay past $1/e \sim 0.368$. The notation intentionally differs from that in Section II C, to distinguish $\tau_{\alpha 0}$ as the uncaging time in the limit of zero apoptosis and zero mitosis $(k_{\delta} \to 0)$.

Our search for glassy behavior is guided by the simple expectation that the dynamics is determined by a competition between the timescale τ_{δ} of division and death events and the timescale of motility driven cage escape $\tau_{\alpha 0}$. In particular, we expect to recover the dynamics of the "bare" (free of cell division and death) AVM for sufficiently small values of τ_{δ} . the following two sentences may need to be removed This can be seen in Fig. 3 (b) and (c), where values of k_{δ} leads the msd and the self-overlap to approach a limiting form which matches the data from Fig. 1. This is useful for identifying the limits of when cell division and cell death events have an impact on mechanics.

That the influence of mitosis and apoptosis will be negligible in the limit $\tau_{\delta} \gg \tau_{\alpha 0}$ where motility operates on a much faster timescale, is reasonable. One may expect to find the opposite behavior in the limit $\tau_{\delta} \ll \tau_{\alpha 0}$, where motility-based dynamics should play a negligible role. To check this, we vary s_0 , because $\tau_{\alpha 0}$ decreases as s_0 increases, in Fig. 3. When $\tau_{\alpha 0}$ is large and τ_{δ} is small (k_{δ} is high) we observe structural relaxation times τ_{α} which are approximately independent of s_0 and by proxy, $\tau_{\alpha 0}$.

B. Simple model for interaction between τ_{δ} and $\tau_{\alpha 0}$

Building on the limiting behaviors described above we develop a very simple model for the interplay between cell death and division and glassy dynamics. Specifically, we assume that the overlap decay rate $\frac{1}{\tau_{\alpha}}$ of a dividing tissue is determined by the weighted sum of the bare cage escape rate $\frac{1}{\tau_{\alpha 0}}$ and the rate of division and death events according to

$$\frac{1}{\tau_{\alpha}} = \frac{1}{\tau_{\alpha 0}} + \frac{C_2}{\tau_{\delta}} \,, \tag{11}$$

where C_2 captures the displacements of surrounding cells resulting from the division and death events. The strong assumption we have made here is that the two rates add in series and are not strongly correlated. Our model predicts that the quantity $\frac{\tau_{\alpha}}{\tau_{\alpha 0}}$ will be a function only of $\frac{\tau_{\delta}}{\tau_{\alpha 0}}$. As shown in Fig. 4, this works fairly well, indicating the generic presence of a regime with fast divisions which dominate the dynamics and τ_{α} is proportional to τ_{δ} (left side of Fig. 4), along with a regime of slow divisions where τ_{α} becomes independent of τ_{δ} (right side of Fig. 4). To understand what controls the crossover, we

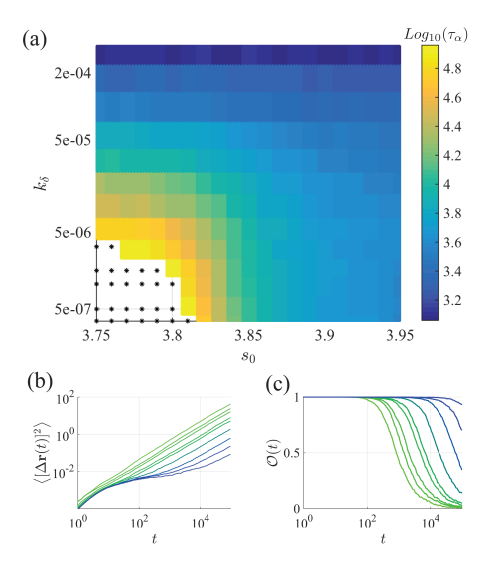


FIG. 3. Dynamical measures of tissues with mitosis and apoptosis for $v_0 = 0.05$ and $D_r = 1$. In (a), the measured overlap decay τ_{α} as a function of s_0 and k_{δ} (the cell cycling rate) is displayed as a colormap, indicating generically that k_{δ} tends to fluidize the tissue. Black stars indicate where τ_{α} is beyond the length of our simulations. In (b) and (c), mean square displacement and overlap curves are plotted for constant $s_0 = 3.76$ over a series (from green to blue) of $\tau_{\delta} = 3k, 6k, 9k, 20k, 30k, 90k, 300k, 900k, 3M$. The following sentence will need to be removed These characteristic examples show that the dynamics approaches the expected values from the "bare" AVM at long enough τ_{δ} . please note what N the data here corresponds to

quantify the impact of cell division and cell death events on the motion of surrounding cells in the next section.

IV. FLOW AND FLUIDIZATION FROM INDIVIDUAL MITOSIS AND APOPTOSIS EVENTS

To quantify the effect of a single cell cycling (division or death) event on the motion of the surrounding tissue, we perform a special set of "single-event" simulations. In these, the standard AVM is run for a short equilibration time before a single cell is chosen at random to undergo either apoptosis or mitosis in a randomly chosen direction. We then monitor the motion of the surrounding tissue cells and average over 1000 realizations.

We measure the individual cell displacement vectors $\{\mathbf{u}_i\}$ and use them to construct a few useful quantities. In order to resolve coherent spatial data, the cells are first binned based on their distance from the event (and in the case of cell division based on their angle relative to the division axis orientation). Within each bin, we calculate two quantities: the vector averaged displacement $\mathbf{u}_{(m,a)}(r,\theta)$ and the vector standard deviation of this average, $\mathbf{w}_{(m,a)}(r,\theta)$, where the subscripts m and a refer to mitosis and apoptosis events, respectively.

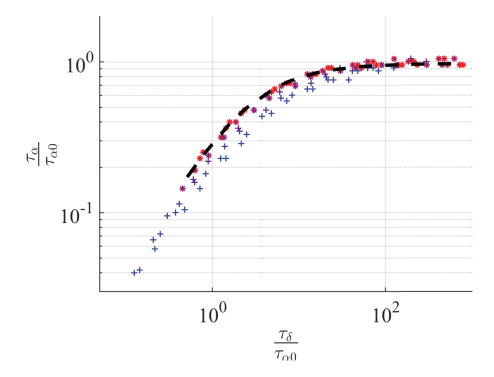


FIG. 4. Observed uncaging times for the AVM with division and death at a range of rates τ_{δ} and (from blue to red) $s_0 = (3.82, 3.825, 3.83, 3.84, 3.85, 3.875, 3.9, 3.925, 3.95)$. The τ_{α} values come close to collapse with simple rescaling using $\tau_{\alpha 0}$ and suggest a universal crossover between motility-dominated and division-death dominated dynamics. Data is represented by "+" symbol where diffusion is never observed in the corresponding "bare" AVM and is represented by "*" otherwise. The black dashed line represents the prediction via Eq. 11 for $s_0 = 3.95$, which matches well with the simulation data. All data is from simulations with $N_{cells} = 3000$, $D_r = 1.0$ and $v_0 = 0.05$.

As noted by Puosi et. al. [39] for the similar case of shear transformations in thermal sphere packings, the vector averaged $\mathbf{u}_{(m,a)}(r,\theta)$ captures the affine elastic response of the surrounding medium. I feel like rather than go into this discussion, we can just say what we mean. This sentence doesn't make sense to me. Michael, could you re-write to be more specific about it? Note that the term affine is used here for historical reasons, despite being distinct from the mathematical concept of an affine transformation where the Jacobian of the mapping is spatially uniform. For systems which undergo simple shear these two notions of affine do coincide. However, we are considering point-like sources of deformation which do not generally result in a uniform Jacobian. This can be seen in the canonical example of the Eshelby inclusion. A straightforward calculation of the response of a homogeneous elastic medium to localized strains [39] suggests that the affine field associated with either a cell death or cell division event should fall off as r^{-1} in two dimensions. This is consistent with the numerical observations in our model, shown in Fig. 5 (a), (b) and (c). In particular, the dashed line in Fig. 5(c) shows the expected scaling of r^{-1} , which is in reasonable agreement with the data.

The "non-affine" piece captured in $\boldsymbol{w}_{(m,a)}(r,\theta)$ quan-

tifies the additional cell displacements that are expected to arise in the vicinity of the event due to disorder in the tissue structure. It is important to note that this quantity captures both local contributions from the mitosis (apoptosis) event, as well as contributions everywhere from active, motility driven, cell motions. To separate the contribution of non-affine elastic displacements from the noise generated by the active forces in the AVM, we define

$$\delta u_{(m,a)}(r)^2 = |\mathbf{w}_{(m,a)}|^2(r) - |\mathbf{w}_{(m,a)}|^2(\infty), \qquad (12)$$

where $|w_{(m,a)}|^2(\infty)$ is the far-field plateau in the non-affine displacement field generated by active noise, shown in the inset to Fig. 5 (d). The resulting rotationally averaged elastic non-affine displacement field, $\delta u_{(m,a)}(r)$, is plotted in Fig. 5(d) for a mitosis event.

We note immediately from the data in Fig. 5(c) that the affine deformations show only minor variation with the change in s_0 and therefore depend little on the quiescent dynamics. In contrast, in Fig. 5(d) we see that the non-affine motions decrease as we go from more rigid tissue at low s_0 to more fluid tissue at $s_0 \sim 3.9$. The degree of variation is, however, still minor and in general the dynamical response to cell division and death events is not expected to be a good indicator of the underlying

dynamical state of the tissue.

A. Predicting Cell Displacements from Mitosis and Apoptosis

Guided by the displacements measured for single cell-death and cell-division events, we construct simplified models to estimate their influence on measurable quantities, such as D and τ_{α} . As pointed out by Ranft et. al. [8], the "affine" displacements identified above produce a constantly changing reference state in an elastic material. Therefore, while cells may oscillate in their cages, the cages themselves move as a result of each event via the displacement fields $\mathbf{u}_{(m,a)}(r,\theta)$. We would like to estimate the effect of this changing reference state on the mean square displacement of our tracer cells. These events will also be creating non-affine displacements on top of the affine motion.

To estimate the total diffusion produced by these different contributions (mitosis and apoptosis, affine and non-affine), we again make the simplifying assumption that they are uncorrelated. We can then write the total diffusion as a sum of the decoupled parts

$$D_{\delta} = D_{(a)}^{aff} + D_{(m)}^{aff} + D_{(a)}^{non-aff} + D_{(m)}^{non-aff}.$$
 (13)

Here, $D_{(m,a)}^{aff}$ captures the diffusion due to the affine $\mathbf{u}_{(m,a)}(r,\theta)$, while $D_{(m,a)}^{non-aff}$ captures diffusion due to the non-affine $\delta\mathbf{u}_{(m,a)}(r,\theta)$. Quantifying each of these diffusion constants will rely on the simplifying assumptions (1) that we may ignore randomness in the timing of the divisions and deaths and (2) that the effect from each event is felt instantaneously by the surrounding tissue. This enables us to simply sum up the contributions from the $n(t) = tN_{cells}/\tau_{\delta}$ events that will have taken place after a time t. Therefore, each term in Equation 13 may be estimated in terms of an average magnitude of motion per cell per event d_i as

$$D_i = \frac{N_{cells}}{4\tau_\delta} \left\langle d^2 \right\rangle_i \,, \tag{14}$$

where the 4 contributions from the rhs of Eq. 13 are now indexed by i. As shown in Appendix C, the values of $\langle d^2 \rangle_i$ may be then estimated from summation of the single-event data shown in Fig. 5(b) and (c). In practice, $\langle d^2 \rangle_i$ is computed as a spatial average

$$\left\langle d^2 \right\rangle_i = \frac{1}{A_T} \int d^2 \mathbf{x} \, d_i^2(r, \theta)$$
 (15)

where $d_i^2(r,\theta)$ represents the square displacements captured in our previously averaged values $\mathbf{u}_{(a,m)}^2(r,\theta)$ and $\delta \mathbf{u}_{(a,m)}^2(r,\theta)$. As these simulation data are measured on a radial grid in discrete bins, the integral in Eq. 15 will turn into a sum over bins, and will have a cutoff radius at $r \sim L/2$. While this excludes cell displacements in

the corners of the simulation box, this contribution is small and the predictions constructed without these corners validate their exclusion. As shown in Fig. 6-(a), each of these contributions to the diffusion D_i is essential to form a complete prediction of the observed diffusion. In Fig. 6 this prediction provides a reasonable estimate, deviating most significantly for tissues which display glassy signatures in the $(\tau_\delta \to \infty$ limit) bare-AVM. This breakdown is expected because the corresponding component D_0 in our prediction is no longer truly a diffusion, as the AVM displays anomalous subdiffusion in this regime as discussed in Appendix II.

This prediction for the diffusion constant may be extended to a prediction for the uncaging time τ_{α} , which performs similarly well. Relating these two quantities relies on the simple result $\mathbf{D} \cdot \tau_{\alpha} = c_B$ which applies to free overdamped Brownian motion in 2D. The brownian constant $c_B \sim 0.1363$ is straightforwardly derived as shown in Appendix. B. Using this relation for our prediction of τ_{α} then relies only on the assumption that on long time scales, the motion of cells will look roughly Brownian.

This is an assumption which, again, begins to break down as motion becomes glassy and cage-escapes become intermittent. In Fig. 4, the green line prediction matches well with (red) data. However, even where anomalous subdiffusive glassy signatures arise, our prediction captures this crossover between motility-dominated and cell division/death dominated dynamics reasonably well.

V. DISCUSSION

To study the influence of propulsive forces on the dynamical state of tissue, we have developed a new Active Vertex Model. This model generalizes the Vertex Model framework to incorporate cell motility in a way that naturally extends the activity implemented in self-propelled Voronoi models [18]. In Section II, we have shown that this new model reproduces qualitatively some of the dynamical results of the SPV model. As found previously by Bi et. al. [18], we locate "glassy" states where exhibit sub-diffusive mean square displacements. The transition from a fluid-like state to a solid-like state is achieved either by decreasing the propulsive forces, or by increasing the effective cell-cell interfacial tension.

We then extended the AVM to investigate the influence of cell death and cell division on tissue dynamics. Our investigation is guided by the expectation that the rate of cell division and death will compete with the "bare" uncaging rate from the AVM to determine the tissue dynamics. This simple picture allows us to identify evidence of glassy states even at finite rates of apoptosis and mitosis, which appears to be in a different regimes from the one explored in [6]. We again emphasize that another pertinent difference is the choice of whether or not to couple cell cycle events directly to the local structural state of the individual cells. This may resolve a standing discrepancy between existing experiments and

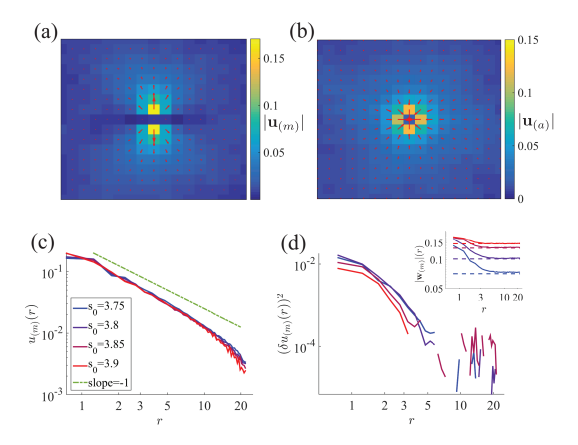


FIG. 5. Affine and non-affine displacements quantified in response to a single mitosis/apoptosis event. In (a) and (b), arrows show the affine displacements in response to a mitosis and apoptosis, respectively, for a tissue with $s_0=3.6,\ v_0=0.05$. Colormap indicates the magnitude of these affine vectors with grid spacing 1.5. Panels (c) and (d) show cell displacements as a function of distance to the cycling cell for $s_0=3.75,3.8,3.85,3.9$ (blue to red). Michael, add numbers for s0 here Panel (c) shows the magnitude of the rotationally averaged affine displacements as a function of distance from a mitosis event. The green dot-dashed line is the expected scaling 1/r. In (d), the values for the $\delta u_{(m,a)}(r)$ per Eq. 12 are plotted similarly, revealing a finite region of non-affine displacements. Inset shows that w plateaus at long distances to the expected value from mean-square displacement data in the bare AVM.

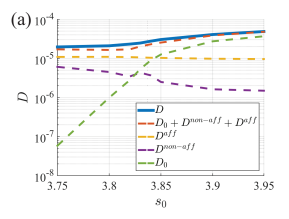
theory. While cell division and death events will generally lead to diffusive fluid-like behavior on the longest timescales, subdiffusive behavior is observed on intermediate timescales when the cell division rate and the bare uncaging rate are slow enough. These tissues are analogous to supercooled fluids and are thereby expected to have elastic responses on these intermediate timescales.

We further characterized the spatial distribution of cell displacements in the vicinity of each individual mitosis and apoptosis event. As expected from previous work for the case of area-preserving deformations [39], we have identified an average displacement field that matches the one obtained from elasticity theory, as well as "non-affine" displacements that capture noise. The elastic affine response decays spatially as 1/r, with a magnitude that does not depend significantly on tissue dynamics. The non-affine displacements, in contrast, vary in magnitude with model parameters, such as cell interfacial tension. Specifically, mitosis and apoptosis events tend to generate more rearrangements in rigid tissues than in

fluid ones.

Using the calculated affine and non-affine displacements resulting from these division and death events, we assess the relative importance of motile forces versus division and death in the tissue dynamics. From this data, we are able to construct predictions for both the diffusion coefficient and the uncaging time. These predictions are very accurate for systems where true diffusive exponents are measured and begin to break down as subdiffusive glassy signatures emerge. Even in these glassy states, however, the prediction gives a good qualitative description of the crossover from motility dominated to cell cycle dominated dynamics.

Taken together, our results provide a general perspective on the role of cell division and cell death in tissues. To reasonably good approximation, these cell cycle events appear to produce displacements which add linearly with the existing cell dynamics. This linear behavior appears independent of the approximate cell lifecycle $2\tau_{\delta}$ for the values probed here, but is expected to break



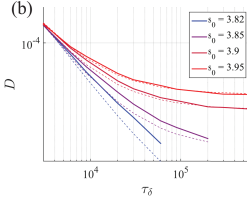


FIG. 6. Separate contributions to diffusion from affine and non-affine displacements and the bare diffusion add up to approximate the measured diffusion in (a). In (b), the validity of this prediction is tested for a range of division rates and a series of s_0 values. Simulations where the MSD does not ever become truly diffusive are not included. Criteria for diffusion here is that the MSD-exponent α be greater than the cutoff value 0.9.

down as this lifetime gets shorter than the speed of sound in the tissue and events begin to interact. In addition, we have not addressed the question of whether glassy signatures persist as the tissue size is increased.

In Appendix C1 we present evidence that the total diffusion due to affine displacements scales as Log(N) for our two-dimensional simulations. In disordered systems we expect, however, that the signal generated by the affine displacements will be comparable to the noise generated by non-affine motions at a characteristic length-scale, as has been previously highlighted for sheared particulate packings [40]. Beyond this lengthscale, we no longer expect the Log(N) scaling to hold, and so the system will remain glassy in the thermodynamic limit. Testing this hypothesis carefully is therefore an important avenue for future investigations.

So far, we have focused on dynamics that explicitly keep the tissue at fixed number density, in order to pinpoint the effect of cell divisions on tissue dynamics. However, in many experiments on epithelial monolayers, a common and interesting regime is where cell divisions outnumber cell deaths and so the number density increases rapidly. An important open question is how Active Vertex model parameters, such as s_0 or v_0 change during epithelial densification. Several experiments on pairs or triplets of cells indicate that many epithelial cells exhibit "contact inhibition of locomotion", where cells reduce traction forces and slow down upon contact [10, 41]. Such behavior could be modeled by decreasing v_0 as a function of time in densifying monolayers, and additional experiments to measure traction forces and fluctuations during such processes would be very valuable in constraining model predictions. Perhaps even more interestingly, it remains unclear how cells regulate their shape s_0 during such processes. One possibility is that cells attempt to maintain the same shape despite changes

to density, meaning that they must change their perimeter to match their decreasing area as a function of time. Alternatively, one could postulate that cells keep their preferred perimeter fixed while their area decreases dramatically, resulting in an increasing s_0 that would fluidize a tissue. This latter hypothesis seems inconsistent with existing experimental data, where tissues are generically found to solidify as they become more dense. Careful studies of cell shape coupled with careful studies of interfacial tensions [42] would help to constrain models and better test model predictions.

Finally, it is interesting to think about how cell division and death rates are themselves affected by tissue dynamics. For example, it is known that in many cell types cell division rates are governed by the magnitude of local stresses that build up in the tissue [37, 43]. In addition, the axis of cell division is also controlled by local stresses This creates the possibility for interesting feedbacks, where tissue dynamics is controlled by the rate of cell division and the rate of cell division is controlled by tissue dynamics. Understanding precisely how division and death affect dynamics is therefore very important for predicting how such feedback loops can control tissue growth and patterning. It would be interesting to explore how such feedback loops generate patterns in AVMs, and compare to experiment.

_

VI. APPENDICES

Appendix A: The Active Vertex Model

1. Forces in the Active Vertex Model

To our knowledge, the "Active Vertex Model" which is employed in our simulations has not been explored in any previous work. Therefore, we include here a more detailed description of the model implementation.

In this Active Vertex Model (AVM), the degrees of freedom are the positions of the vertices of the spatial tiling. The vertex labeled μ follows an overdamped equation of motion

$$\partial_t \mathbf{r}^{\mu} = -\frac{1}{\gamma} \frac{\partial}{\partial \mathbf{r}^{\mu}} E \,, \tag{A1}$$

where γ is the substrate friction. The effective energy

$$E = E_{shape} - \gamma v_0 \sum_{cells-a} \hat{\mathbf{n}}(\theta_a) \cdot \mathbf{r}^a, \qquad (A2)$$

captures the self-propulsion force (of magnitude v_0) of each cell a in direction θ_a in addition to the standard shape energy terms from Eq. 1. The geometric center (centroid) \mathbf{r}^a of cell a defined by

$$\mathbf{r}^{a} = \frac{1}{6A_{a}} \sum_{\nu} \left(\mathbf{r}^{\nu} + \mathbf{r}^{\nu+1} \right) \left(\mathbf{r}^{\nu} \times \mathbf{r}^{\nu+1} \right) \cdot \hat{\mathbf{k}}, \quad (A3)$$

captures the center of mass of a polygon of uniform mass density. Here, the sum goes counterclockwise over the N_a vertices on cell a and $\mathbf{r}^{N_a+1} = \mathbf{r}^1$. The polygon area may be expressed in similar terms as

$$A_a = \frac{1}{2} \sum_{\nu} (\mathbf{r}^{\nu} \times \mathbf{r}^{\nu+1}) \cdot \hat{\mathbf{k}}. \tag{A4}$$

The force (and therefore motion) on each vertex can be calculated by carrying out the derivatives in Eq. 6. Each vertex in this model is connected to 3 cells and each cell energy will contribute separate terms to the net motion of the vertex. If vertex μ is connected to cells a, b and c, then the motion breaks down into

$$\partial_{t}\mathbf{r}^{\mu} = -\frac{1}{\gamma} \left(\frac{\partial}{\partial \mathbf{r}^{\mu}} E_{a} + \frac{\partial}{\partial \mathbf{r}^{\mu}} E_{b} + \frac{\partial}{\partial \mathbf{r}^{\mu}} E_{c} \right) + v_{0} \left(\frac{\partial (\hat{\mathbf{n}}(\theta_{a}) \cdot \mathbf{r}^{a})}{\partial \mathbf{r}^{\mu}} + \frac{\partial (\hat{\mathbf{n}}(\theta_{b}) \cdot \mathbf{r}^{b})}{\partial \mathbf{r}^{\mu}} + \frac{\partial (\hat{\mathbf{n}}(\theta_{c}) \cdot \mathbf{r}^{c})}{\partial \mathbf{r}^{\mu}} \right).$$
(A5)

For simplicity, we may focus on the contributions from cell a. As has been identified in previous work [33], the shape energy produces tension-based and pressure-based forces on each vertex. The shape-based force on vertex μ from cell a reads

$$\frac{\partial}{\partial \mathbf{r}^{\mu}} E_a = -\frac{\Pi_a}{2} (\hat{\mathbf{n}}^{ab} l^{\mu\gamma} + \hat{\mathbf{n}}^{ac} l^{\mu\lambda}) - T_a (\hat{\mathbf{l}}^{\mu\gamma} + \hat{\mathbf{l}}^{\mu\lambda}), (A6)$$

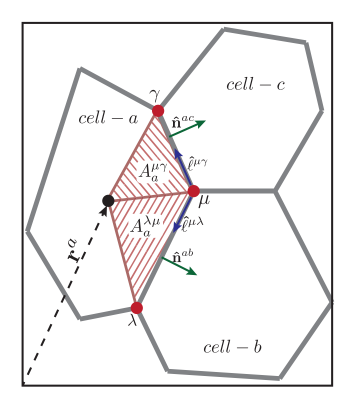


FIG. 7. Illustration of the vectors and geometric components involved in evaluating the forces on vertex μ due to the shape energy and the motility of cell a

where γ and λ index the vertices of cell a which are adjacent to μ , $l^{\mu\gamma}$ ($l^{\mu\lambda}$) and $\hat{\mathbf{l}}^{\mu\gamma}$ ($\hat{\mathbf{l}}^{\mu\lambda}$) are the length and direction of the edge connecting vertex μ to vertex γ (λ) and the unit vector $\hat{\mathbf{n}}^{ab}(\hat{\mathbf{n}}^{ac})$ points across the edge shared by cell a and cell b (c) as in Fig 7. The tension and pressure of cell a are respectively

$$T_a = \frac{\partial E_{shape}}{\partial P_a} \qquad \Pi_a = -\frac{\partial E_{shape}}{\partial A_a}.$$
 (A7)

Similar to the above, we now evaluate derivatives to understand the self propulsion forces which act on the vertices. Again, we will consider only the contributions from cell a. The derivative will take the form

$$\frac{\partial (\hat{\mathbf{n}}(\theta_a) \cdot \mathbf{r}^a)}{\partial \mathbf{r}^{\mu}} = \frac{\partial \mathbf{r}^a}{\partial \mathbf{r}^{\mu}} \cdot \hat{\mathbf{n}}(\theta_a) \tag{A8}$$

With these expressions, taking the derivative in Eq. A8 is tedious but straightforward. This becomes

$$\frac{\partial r_j^a}{\partial r_i^\mu} = \frac{1}{6A_a} \left[\delta_{ij} (\mathbf{r}^\lambda \times \mathbf{r}^\mu) \cdot \hat{\mathbf{k}} + \delta_{ij} (\mathbf{r}^\mu \times \mathbf{r}^\gamma) \cdot \hat{\mathbf{k}} \right. \\
+ (r_j^\lambda + r_j^\mu) R(\pi/2)_{ik} r_k^\lambda + (r_j^\mu + r_j^\gamma) R(-\pi/2)_{ik} r_k^\gamma \\
- 3(R(\pi/2)_{ik} r_k^\lambda + R(-\pi/2)_{ik} r_k^\gamma) r_j^a \right], \tag{A9}$$

where

$$R_{ij}(\theta) = \begin{bmatrix} \cos(\theta) & -\sin(\theta) \\ \sin(\theta) & \cos(\theta) \end{bmatrix}$$
 (A10)

is an active vector rotation by an angle θ , i,j index cartesian components in the x-y plane and Einstein summation convention is assumed for repeated indices. While the expression in Eq. A9 is a bit unwieldy, we note that it is translationally invariant and may consider it in a more convenient coordinate system which has its origin at \mathbf{r}_a . Using this, we find the force on vertex μ due to the motility of cell a as

$$\begin{split} \mathbf{f}_{i}^{\mu(a)} &= v_{0}\hat{n}(\theta_{a})_{j} \frac{\partial r_{j}^{a}}{\partial r_{i}^{\mu}} \\ &= \frac{v_{0}\hat{n}(\theta_{a})_{i}}{3A_{a}} [A_{a}^{\lambda\mu} + A_{a}^{\mu\gamma}] \\ &+ \frac{v_{0}}{6A_{a}} \left[R(\pi/2)_{ik} \tilde{r}_{k}^{\lambda} \hat{n}(\theta_{a})_{j} (\tilde{r}_{j}^{\lambda} + \tilde{r}_{j}^{\mu}) \right. \end{split} \tag{A11}$$

$$&+ R(-\pi/2)_{ik} \tilde{r}_{k}^{\gamma} \hat{n}(\theta_{a})_{j} (\tilde{r}_{j}^{\mu} + \tilde{r}_{j}^{\gamma}) \right],$$

where $A_a^{\lambda\mu}$ and $A_a^{\mu\gamma}$ are the area of triangles with vertices $\{{\bf r}^a,{\bf r}^\lambda,{\bf r}^\mu\}$ and $\{{\bf r}^a,{\bf r}^\mu,{\bf r}^\gamma\}$ respectively, as shown in Fig. 7. While this expression does not lend itself to insight, we can see on inspection that the first terms will move the vertex in the direction of $\hat{\bf n}(\theta_a)$. The second terms in the large square brackets will roughly serve to shrink the length of interfaces at the back of the cell, while expanding the length of interfaces at the front.

While these forces determine the continuous timeevolution of the vertices, there can be no large-scale migration of cells through the tissue until topological rearrangements are allowed. We therefore include a protocol for T1 rearrangements (as shown in Fig 8) which are sufficient to explore the space of cellular topological configurations at constant density. In practice, the edge lengths are periodically checked after a time t_{T1} for values lower than a threshold ℓ_c . These edges are then topologically rearranged so that the two cells which initially share the short edge are no longer in contact. The edge is then rotated by $\pi/2$ and the length extended by a factor λ_{T1} . The value of $t_{T1} = 0.05$ used here is chosen for speed of simulation, while the rescaling factor $\lambda_{T1} = 2$ is chosen to avoid "T1-traps" whereby the same transition may repeat itself regardless of energetic favorability. the cutoff length $\ell_c = 0.04$ is chosen small enough to make the transition appear continuous but also large enough so that a vertex may "find" and activate the desired T1. The parameters ℓ_c , λ_{T1} and t_{T1} have been separately varied within reasonable ranges and the impact on the dynamics appears insubstantial.

We note again that this form of the Active Vertex Model is chosen to minimize the differences with the Self-Propelled Voronoi model. The differences are limited to the following: (1) the Active Vertex Model has

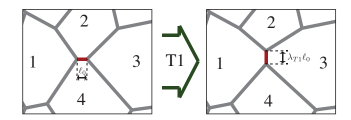


FIG. 8. A T1 topological rearrangement in our AVM simulations identifies an edge with length $l_0 < l_c$, rewires the network connections appropriately, rotates by $\pi/2$, and extends the edge length to a factor λ times its original length.

more degrees of freedom, avoiding the shape constraints of Voronoi tesselations, (2) T1 rearrangements in the AVM must be done by hand, while in the SPV they come about naturally and (3) motility in the AVM is designed to propel the centroid of the cell, whereas the SPV this is replaced by the Voronoi center.

Simulations are developed and run using the "cell-GPU" codebase developed by one of us previously [44].

2. Subdiffusive Exponents and Exotic Dynamical States in the Active Vertex Model

While the main text provides a coherent description of the glassy transition present in the AVM, Vertex Models governed by an energy of the type in Eq. 1 may also be subject to anomalous energy landscapes and nontrivial inherent stresses [28, 29]. As a reflection of these features, the dynamical signatures such as the mean-square displacement (msd) and the Overlap (as defined in Eq. 1) may appear anomalous in the vicinity of this glassy phase. Here, we explore these anomalies in greater detail.

The mean-square displacement (msd) in thermal particulate systems approaching a glass transition will display the signatures of "caging". This manifests as a flat plateau in the msd as particles reach this cage size (radius) and are temporarily trapped before diffusing away at a later timescale [45, 46]. In contrast, the AVM does not appear to completely flatten at any discernible cagesize. Instead, we observe an extended subdiffusive regime which emerges as the system approaches the glassy state, implying that there is not a well defined cage-size. This is a suggestion that the cells are exploring a distribution of local metastable states as they attempt to realize certain shapes. It may therefore not be useful to think of the travel between these metastable states as "cagebreaking" which come paired with intermittent jumps in displacement on the order of the particle size. Instead, it appears that a distribution of effective cages is continuously traversed by the cells.

Our narrative thus far points to the unconventional ge-

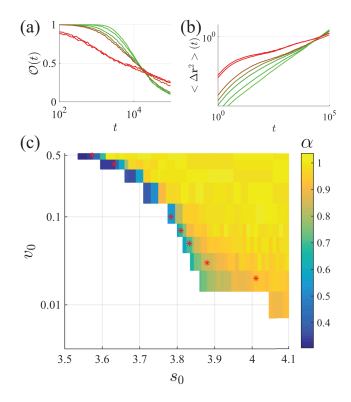


FIG. 9. Revealing the persistent subdiffusion in the Active Vertex Model. In **c**, the long-time exponent of the MSD is plotted on a colormap for the same phase data as in main text Fig. 1c. Phase points where the uncaging time $\tau_{\alpha 0}$ is beyond the simulation time are left white. Along the interface, a region where this exponent α is distinctly lower than unity despite the finite uncaging time indicates the anomalous dynamics. In **a**, and **b**, we display the varied character of the overlap (**a**) and the MSD (**b**) curves for points with the same uncaging time $\tau_{\alpha 0} \sim 2 \cdot 10^4$ and corresponding, from green to red, to the points at $(s_0, v_0) = (4.01, 0.02)$, (3.88, 0.03), (3.833, 0.05), (3.81, 0.07), (3.783, 0.1), (3.632, 0.4), (3.573, 0.5) as indicated by the red stars in the phase diagram. All data for $D_r = 1.0$ and $N_{cells} = 300$.

ometric mechanics of the Vertex Model as the cause of this anomalous behavior, yet there is another potential culprit. Our model also differs from that of traditional glasses and colloids in the persistence of the propulsive forces. As we have employed a non-zero persistence time of these forces as $\tau_p = 1/D_r = 1$, we also perform simulations with the motility replaced by thermal forces on the vertices at effective temperature T. More precisely, these simulations are run in the AVM with v_0 set to zero, and the equation of motion of a vertex instead obeying

$$\partial_t \mathbf{r}^{\mu} = -\frac{1}{\gamma} \frac{\partial}{\partial \mathbf{r}^{\mu}} E + \eta^{\mu} \tag{A12}$$

where

$$\langle \eta^{\mu}(t) \rangle = 0 \tag{A13}$$

and

$$\langle \eta_i(t_1)\eta_j(t_2)\rangle = \frac{2T}{\gamma}\delta_{ij}\delta(t_1 - t_2)$$
 (A14)

These results are summarized in Fig. 10 and indicate qualitatively the same behavior as that seen for cell motile forces. We also note that previous studies in the Self-Propelled Voronoi and Thermal Voronoi models have not thus far displayed this behavior. Altogether, this indicates that this strange extended subdiffusion is associated with the Vertex Model geometric energy along with the vertex degrees of freedom that it acts on.

Finally, we investigate temporal fluctuations in the mechanical stress as defined instantaneously by

$$\sigma_{ij} = \frac{1}{A_T} \sum_{cells-a} A_a \left[\frac{\partial E_{shape}}{\partial A_a} \delta_{ij} + \frac{1}{2A_a} \sum_{\mu\nu \in a} \frac{\partial E_{shape}}{\partial l_i^{\mu\nu}} \right]$$
(A15)

as in Ref. [33]. From this global tissue stress, we compute the stress autocorrelation function via

$$C_{\sigma}(t) = \frac{\langle \sigma_{xy}(t_0 + t)\sigma_{xy}(t_0) \rangle_{t_0}}{\langle \sigma_{xy}^2(t_0) \rangle_{t_0}}.$$
 (A16)

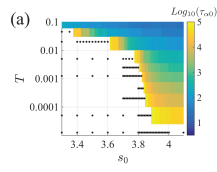
Where the angular brackets denote the average over both noise and over the time t_0 . In practice, the computation of this correlation is accomplished using the algorithm described by Ramírez et. al. in Ref. [47]. This stress auto-correlation function, shown in Fig. 10 for a range of s_0 values, indicates additional unusual behavior. Specifically, we do not resolve the traditionally expected "beta" (fast) and "alpha" (slow) decorrelation times. Instead, the stress appears to decorrelate in three or more stages across a range of timescales. This again agrees with our developing perspective that the process of cage breaking occurs across a broad range of different length scales (and therefore different time scales) in cellular tissues.

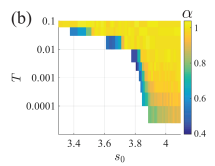
Appendix B: Relating the Uncaging time to the Diffusion Constant for Brownian Particles

This work relies largely on two scalar measures of tissue dynamics, the uncaging time τ_{α} and the diffusion coefficient D. Here we show that, for the case of Brownian motion in two dimensions, these quantities may be related explicitly as

$$\tau_{\alpha} D = c_B \equiv \frac{-b^2}{4ln[1 - \frac{1}{e}]}, \qquad (B1)$$

where b is the radius defining the cage size in the overlap function (Eq. 10). This equation (which is related to the Stokes-Einstein relation) breaks down as a material approaches a glass transition, and therefore provides important context for interpreting dynamical parameters.





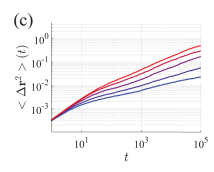


FIG. 10. Summarizing the dynamical signatures in the Thermal Vertex Model. In \mathbf{a} , the uncaging time $\tau_{\alpha 0}$ is displayed in a logarithmic colormap revealing similar behavior to that for the AVM shown in Fig. 1c. Black asterisks again indicate phase points where $\tau_{\alpha 0}$ is out of range. In \mathbf{b} , the long-time MSD exponent α is plotted in a colormap for the same phase space in \mathbf{a} , again with the points where the uncaging time is out of range left blank. A region of phase space with subdiffusive behavior extending sufficiently beyond cage escape is indicated by the dark region near the boundary. In \mathbf{c} , the extended subdiffusion is shown for points (from blue to red) at $s_0 = (3.75, 3.77, 3.79, 3.81, 3.83)$ and T = 0.00045. All data for $N_{cells} = 300$.

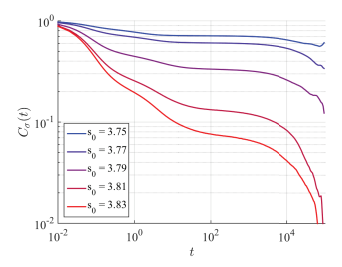


FIG. 11. The shear-stress autocorrelation function $C_{\sigma}(t)$ in the Thermal Vertex Model shows unusual temporal character in the glassy state. This data for T=0.00045 over a series of s_0 values at the same points as in Fig.10c. Three timescales are suggested instead of the traditional two "alpha" and "beta" relaxation times.

To understand Eq. B1, we note that Brownian motion may be described by a smooth probability density function $P(\mathbf{x},t)$ which is governed by Fick's Law

$$\partial_t P(\mathbf{x}, t) = D\nabla^2 P(\mathbf{x}, t).$$
 (B2)

To understand the measured uncaging time, we must first solve for the $P(\mathbf{x},t)$ for a single particle starting at an arbitrary point \mathbf{x}_0 at a time t_0 . This corresponds to solving with an initial condition $P(\mathbf{x}_0,t_0)=\delta(\mathbf{x}-\mathbf{x}_0)$. Preserving the normalization of P, this is solved for $t>t_0$

by

$$P(\mathbf{x},t) = \frac{1}{4\pi D(t-t_0) \exp\frac{-(\mathbf{x}-\mathbf{x}_0)^2}{4D(t-t_0)}}.$$
 (B3)

This probability distribution may be used to take averages and we may therefore rewrite the overlap,

$$\mathcal{O}(t) = \langle \Theta(|\delta \mathbf{r}_i(t)| - b) \rangle$$
, (B4)

as

$$\mathcal{O}(t) = \int_0^b r dr \int_0^{2\pi} d\theta P(r, t), \qquad (B5)$$

where, for convenience, we have set $t_0=0$ and $r=|\mathbf{x}-\mathbf{x}_0|$. The uncaging time is then defined by the condition $\mathcal{O}(\tau_\alpha)=1/e$ and we may insert Eq. B3 into Eq. B5, integrate, and manipulate to find our desired result in Eq. B1. Evaluating this constant for the case b=0.5, we find that $\tau_\alpha D=c_B\sim 0.1363$.

Appendix C: Displacements due to single division or death events

Here we describe our method of converting displacement data for single apoptosis and mitosis events into averaged motions per event $\langle d^2 \rangle_i$ as well as a predicted diffusion coefficient D_δ per Equations 13 and 15.

1. Estimating affine contributions to displacement

To estimate the first and second Diffusion coefficients on the right hand side of Eq. 13, we will use the "single-event" affine motion data $\mathbf{u}_{(m,a)i}(r,\theta)$ obtained in Section IV. On the long timescales which we are interested

in, each event effectively instantaneously moves the cells via the affine displacement $\mathbf{u}_{(m,a)i}(r,\theta)$ and the cells then follow an equation of motion

$$\Delta \mathbf{x}_{(m,a)}(t) = \sum_{i}^{n(t)} \mathbf{u}_{(m,a)}(r_i, \theta_i)$$
 (C1)

in which the n(t) displacements due to previous events are simply summed up. With this EOM, the msd may be computed as

$$\langle \Delta \mathbf{x}_{(m,a)}^2(t) \rangle = \sum_{i}^{n(t)} \langle d^2 \rangle_{(m,a)}^{aff} . \tag{C2}$$

where we have implicitly defined the average affine motion per event $\langle d^2 \rangle_{(m,a)}^{aff}$ and $\langle \rangle$ represents the average over the realizations of these apoptosis and mitosis events. As the stochasticity here comes from the spatial positioning (and orientation) of the mitosis (apoptosis) event with respect to our tracer, finding this average for $\langle \mathbf{u}_{(m,a)i}^2 \rangle$ amounts to integrating over the possible positions and orientations of this mitosis (apoptosis) event. This amounts to a spatial integral

$$\langle d^2 \rangle_{(m,a)}^{aff} = \frac{\rho}{N_{cells}} \int d\mathbf{x} \, \mathbf{u}_{(m,a)i}^2(r,\theta)$$
 (C3)

over the affine displacement data where ρ is the global number density of cells. To directly produce a numeric value of $\langle d^2 \rangle_{(m,a)}^{aff}$ from the data, the integral is turned into a sum over all the bins in r and θ . The Diffusion coefficient produced by these motions is then defined from Eq. C2 as

$$D_{(m,a)}^{aff} = \frac{N_{cells}}{4\tau_{\delta}} \left\langle d^2 \right\rangle_{(m,a)}^{aff} \tag{C4}$$

The estimate produced above leads to useful and accurate predictions of the dynamics as shown in Figures 6 and 4 in the main text. However, this prediction provides little insight into the functional form of these affine diffusions. Some insight can instead be gained by estimating the integral in Eq. C3 using some assumptions based on the data. In Fig. 12(a) we see that averaging these affine displacements over the angular bins reveals a 1/r trend which may be captured by $u_{(m,a)}(r) = u_{(m,a)}/(\rho r)$. This trend persists out to a distance of $\sim 0.3L$ where the event begins to interfere with its periodic image and drops off quickly (here $L = \sqrt{N_{cells}/\rho}$ is the periodic box length). A good approximation of the average displacement is given by

$$\langle d^2 \rangle_{(m,a)}^{aff} = \frac{\rho}{N_{cells}} \int_{r_{congl.}}^{r_{large}} dr r \langle \mathbf{u}_{(m,a)}^2 \rangle_{\theta}(r), \quad (C5)$$

where the small r_{small} and large $r_{large} \sim 0.3L$ cutoffs of integration respectively capture the typical cell neighbor spacing and the extent of the affine field. Inserting

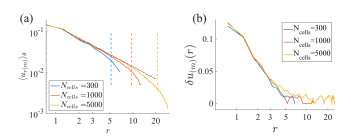


FIG. 12. Scaling of mitosis-based displacement fields with system size for a tissue with $s_0 = 3.8, v_0 = 0.05, D_r = 1.0$. In (a) the affine displacements have consistent form of a 1/r field which fills up to a finite fraction of the box dimension. Dotted vertical lines indicate the cutoff estimate $r_{large} = 0.3\sqrt{N_{cells}/\rho}$. In (b), the non-affine displacements extend over a finite region of fluidization and are unchanged by increasing system size.

the above form of $u_{(m,a)}(r)$ and integrating, we find an expression for the Diffusion

$$D_{(m,a)}^{aff} = \frac{N_{cells}}{4\tau_{\delta}} \langle \mathbf{u}_{(m,a)i}^{2} \rangle$$

$$= \frac{\pi u_{(m,a)}^{2}}{2\tau_{\delta}\rho} ln \left(\frac{r_{large}}{r_{small}}\right).$$
(C6)

Inserting $r_{large} = 0.3 \sqrt{N_{cells}/\rho}$ we will find that this contribution to the Diffusion scales as D $\sim Log(N)$. However, this predicted scaling will only continue as long as the appropriate long-distance cutoff r_{large} is set by the box size. As we consider increasingly large tissues, these affine fields will interfere with that from other events rather than periodic images of the same event. Therefore we expect that the cutoff length (and by proxy, the predicted $D_{(m,a)}^{aff}$) will become independent of the tissue size in large enough tissues.

2. Estimating non-affine contributions to displacement

Similar to the previous section, we now consider the contribution to diffusion from the non-affine (third and fourth) terms in Eq. 13. Similar to Eq. C1, the contribution to the tracer dynamics generated by these non-affine displacements is written as

$$\Delta \mathbf{r}_{(m,a)}^{non-aff}(t) = \sum_{i=0}^{n(t)} \delta \mathbf{u}_{(m,a)i}, \qquad (C7)$$

where $\delta \mathbf{u}_i$ is the non-affine displacement produced by the *i*-th mitosis(apoptosis) event. This displacement is assumed to have a random direction and magnitude determined by the distribution of $\delta u_{(m,a)}(r)$ from Eq. 12. The mean square displacement again relies on the quantity $\langle d^2 \rangle_{(m,a)}^{non-aff}$, which again takes the form of a spatial

average

$$\langle d^2 \rangle_{(m,a)}^{aff} = \frac{\rho}{N_{cells}} \int d\mathbf{x} \, \delta \mathbf{u}_{(m,a)i}^2(r,\theta) \,.$$
 (C8)

Turning this integral into a sum over bins, the non-affine displacement data from Section IV can be used to estimate these contributions to diffusion

$$D_{(m,a)}^{non-aff} = \frac{N_{cells}}{4\tau_{\delta}} \left\langle d^2 \right\rangle_{(m,a)}^{non-aff} . \tag{C9}$$

Similar to the case of the affine displacements, we may estimate the form of the diffusion Eq. C8 based on some more tangible parameters. This is accomplished by approximating the non-affine displacements as a uniform region of fluidization with $\delta u_{(m,a)}(r) = w_{(m,a)}\Theta(R_{\sigma} - r)$.

The diffusion is then

$$D_{(a,m)}^{non-aff} = \frac{N_{cells}}{4\tau_{\delta}} < \delta \mathbf{u}_{(a,m)i}^{2} >$$

$$= \frac{\rho \pi w_{(m,a)}^{2} R_{w}^{2}}{4\tau_{\delta}}.$$
(C10)

Fig. 5 (b) indicates that all of these parameters above are independent of system size as long as L is sufficiently larger than R_w . Therefore the contribution of these displacements to cell dynamics is also independent of system size as long as $\sqrt{N_{cells}/\rho} > 2R_w$.

These non-affine components may also be checked for scaling with increasing system size. As shown in Fig. 12-b, the mitosis event creates a finite region of fluidization which is consistent across system sizes.

- T. E. Angelini, E. Hannezo, X. Trepat, M. Marquez, J. J. Fredberg, and D. A. Weitz, Proceedings of the National Academy of Sciences of the United States of America 108, 4714 (2011).
- [2] E.-M. Schotz, M. Lanio, J. A. Talbot, and M. L. Manning, Journal of The Royal Society Interface 10, 20130726 (2013), arXiv:arXiv:1307.4454v1.
- [3] J.-A. Park, J. H. Kim, D. Bi, J. A. Mitchel, N. T. Qazvini, K. Tantisira, C. Y. Park, M. McGill, S.-H. Kim, B. Gweon, J. Notbohm, R. Steward Jr, S. Burger, S. H. Randell, A. T. Kho, D. T. Tambe, C. Hardin, S. A. Shore, E. Israel, D. A. Weitz, D. J. Tschumperlin, E. P. Henske, S. T. Weiss, M. L. Manning, J. P. Butler, J. M. Drazen, and J. J. Fredberg, Nature Materials 14, 1040 (2015).
- [4] A. G. Renehan, C. Booth, and C. S. Potten, BMJ (Clinical research ed.) 322, 1536 (2001).
- [5] C. B. Thompson, Science (New York, N.Y.) 267, 1456 (1995).
- [6] D. A. Matoz-Fernandez, K. Martens, R. Sknepnek, J. L. Barrat, and S. Henkes, Soft Matter 13, 3205 (2017).
- [7] D. Matoz-Fernandez, E. Agoritsas, J.-L. Barrat, E. Bertin, and K. Martens, Physical Review Letters 118, 158105 (2017).
- [8] J. Ranft, M. Basan, J. Elgeti, J.-F. Joanny, J. Prost, and F. Julicher, Proceedings of the National Academy of Sciences 107, 20863 (2010), arXiv:78650462567.
- [9] A. N. Malmi-Kakkada, X. Li, H. S. Samanta, S. Sinha, and D. Thirumalai, Physical Review X 8, 021025 (2018).
- [10] A. Puliafito, L. Hufnagel, P. Neveu, S. Streichan, A. Si-gal, D. K. Fygenson, and B. I. Shraiman, Proceedings of the National Academy of Sciences of the United States of America 109, 739 (2012).
- [11] P. Martin, S. M. Parkhurst, and M. Peifer, Development (Cambridge, England) 131, 3021 (2004).
- [12] X. Trepat, M. R. Wasserman, T. E. Angelini, E. Millet, D. A. Weitz, J. P. Butler, and J. J. Fredberg, Nature Physics 5, 426 (2009).
- [13] F. Giavazzi, C. Malinverno, S. Corallino, F. Ginelli, G. Scita, and R. Cerbino, Journal of Physics D: Applied Physics 50, 384003 (2017).
- [14] F. Giavazzi, M. Paoluzzi, M. Macchi, D. Bi, G. Scita, M. L. Manning, R. Cerbino, and M. C. Marchetti, Soft

- matter **14**, 3471 (2018).
- [15] S. Henkes, Y. Fily, and M. C. Marchetti, Physical Review E 84, 040301 (2011).
- [16] L. Berthier and J. Kurchan, Nature Physics 9, 310 (2013).
- [17] L. Berthier, Physical Review Letters 112, 220602 (2014).
- [18] D. Bi, X. Yang, M. C. Marchetti, and M. L. Manning, Physical Review X 6, 021011 (2016).
- [19] R. Farhadifar, J.-C. Röper, B. Aigouy, S. Eaton, and F. Jülicher, Current biology: CB 17, 2095 (2007).
- [20] L. Hufnagel, A. A. Teleman, H. Rouault, S. M. Cohen, and B. I. Shraiman, Proceedings of the National Academy of Sciences of the United States of America 104, 3835 (2007).
- [21] D. B. Staple, R. Farhadifar, J. C. Röper, B. Aigouy, S. Eaton, and F. Jülicher, The European Physical Journal E 33, 117 (2010).
- [22] D. Bi, J. H. Lopez, J. M. Schwarz, and M. L. Manning, Soft Matter 10, 1885 (2014).
- [23] D. Bi, J. H. Lopez, J. M. Schwarz, and M. L. Manning, Nature Physics 11, 1074 (2015).
- [24] M. Merkel, R. Etournay, M. Popović, G. Salbreux, S. Eaton, and F. Jülicher, Physical Review E 95, 032401 (2017).
- [25] D. L. Barton, S. Henkes, C. J. Weijer, and R. Sknepnek, PLOS Computational Biology 13, e1005569 (2017).
- [26] A. J. Kabla, Journal of the Royal Society, Interface / the Royal Society 9, 3268 (2012).
- [27] M. Chiang and D. Marenduzzo, EPL (Europhysics Letters) 116, 28009 (2016).
- [28] D. M. Sussman and M. Merkel, Soft matter 14, 3397 (2018).
- [29] D. M. Sussman, M. Paoluzzi, M. Cristina Marchetti, and M. Lisa Manning, EPL (Europhysics Letters) 121, 36001 (2018).
- [30] C. Metzner, C. Mark, J. Steinwachs, L. Lautscham, F. Stadler, and B. Fabry, Nature Communications 6, 7516 (2015).
- [31] G. Passucci, M. E. Brasch, J. H. Henderson, V. Zabur-daev, and M. L. Manning, PLOS Computational Biology 15, e1006732 (2019).
- [32] R. Sknepnek and S. Henkes, Physical Review E 91,

- 022306 (2015).
- [33] X. Yang, D. Bi, M. Czajkowski, M. Merkel, M. L. Manning, and M. C. Marchetti, Proceedings of the National Academy of Sciences of the United States of America, 201705921 (2017).
- [34] J. Rosenblatt, M. C. Raff, and L. P. Cramer, Current biology: CB 11, 1847 (2001).
- [35] I. Conlon and M. Raff, Journal of Biology 2, 7 (2003).
- [36] H. Dolznig, F. Grebien, T. Sauer, H. Beug, and E. W. Müllner, Nature Cell Biology 6, 899 (2004).
- [37] S. J. Streichan, C. R. Hoerner, T. Schneidt, D. Holzer, and L. Hufnagel, Proceedings of the National Academy of Sciences 111, 5586 (2014).
- [38] J. Baker and D. Garrod, Journal of cell science 104 (Pt 2), 415 (1993).
- [39] F. Puosi, J. Rottler, and J.-L. Barrat, Physical Review E 89, 042302 (2014).

- [40] A. Lemaître and C. Caroli, Physical Review Letters 103, 065501 (2009).
- [41] M. Abercrombie, In Vitro 6, 128 (1970).
- [42] A. Saraswathibhatla and J. Notbohm, (2019), arXiv:1901.10018.
- [43] B. I. Shraiman, Proceedings of the National Academy of Sciences of the United States of America 102, 3318 (2005).
- [44] D. M. Sussman, Computer Physics Communications 219, 400 (2017).
- [45] L. Berthier and G. Biroli, Reviews of Modern Physics 83, 587 (2011).
- [46] E. R. Weeks and D. Weitz, Chemical Physics 284, 361 (2002).
- [47] J. Ramírez, S. K. Sukumaran, B. Vorselaars, and A. E. Likhtman, The Journal of Chemical Physics 133, 154103 (2010).