

Measuring the Mechanical Stress Induced by an Expanding Multicellular Tumor System: A Case Study

V. D. Gordon^{1,2}, M. T. Valentine^{1,2}, M. L. Gardel^{1,2},
D. Andor-Ardó¹, S. Dennison², A. A. Bogdanov³,
D. A. Weitz^{1,2,*} and T. S. Deisboeck^{1,4,5}

¹ Division of Engineering and Applied Sciences and ² Department of Physics, Harvard University, Cambridge, MA 02138; ³ Center for Molecular Imaging Research, Massachusetts General Hospital, Harvard Medical School, Charlestown, MA 02129; ⁴ Complex Biosystems Modeling Laboratory, Harvard-MIT (HST) Athinoula A. Martinos Center for Biomedical Imaging, HST-Biomedical Engineering Center, Massachusetts Institute of Technology, Cambridge, MA 02139; ⁵ Molecular Neuro-Oncology Laboratory, Massachusetts General Hospital, Harvard Medical School, Charlestown, MA 02129.

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***Corresponding author (and reprint requests):**

David A. Weitz, Ph.D.
Department of Physics
Harvard University
Cambridge, MA 02138
Tel: 617-496-2842
Fax: 617-496-9564
Email: weitz@deas.harvard.edu

ABSTRACT

Rapid volumetric growth and extensive invasion into brain parenchyma are hallmarks of malignant neuroepithelial tumors *in vivo*. Little is known, however, about the mechanical impact of the growing brain tumor on its microenvironment. To better understand the environmental mechanical response, we used multi-particle tracking and microrheological methods to probe the environment of a dynamically expanding, multicellular brain tumor spheroid that grew for six days in a three-dimensional Matrigel-based *in vitro* assay containing 1.0 μm latex beads. These beads act as reference markers for the gel, allowing us to image the spatial displacement of the tumor environment using high-resolution timelapse video-microscopy. The results show that the volumetrically expanding tumor spheroid *pushes* the gel outward and that this tumor-generated pressure propagates to a distance greater than the initial radius of the tumor spheroid. Intriguingly, beads near the tips of invasive cells are displaced inward, towards the advancing invasive cells. Furthermore, this localized cell *traction* correlates with a marked increase in total invasion area over the observation period. This case study presents evidence that an expanding microscopic tumor system exerts both significant mechanical pressure *and* significant traction on its microenvironment.

Keywords: brain tumor, multicellular tumor spheroid, growth dynamics, invasion, traction, stress, strain, Matrigel, in vitro assay, particle tracking, microrheology

INTRODUCTION

The outcome for patients suffering from highly malignant brain tumors remains dismal in spite of all therapeutic efforts. The most malignant form, glioblastoma, which accounts for 23% of all primary brain tumor cases, has a mean patient age at diagnosis of 62 years and, in the age group of 45-64 years, a 5-year survival rate of only 2.0 % [1]. The reasons for the almost inevitable treatment failure include rapid volumetric growth, early development of treatment resistance and, most importantly, extensive tissue infiltration, which leaves these neoplasms surgically incurable.

A better understanding of the structural remodeling of brain parenchyma by tumor proliferation and invasion is essential to understanding the processes that facilitate diffuse infiltration and tumor recurrence. However, the mechanical relationship between an expanding tumor system and its microenvironment is still poorly understood. By placing multicellular tumor spheroids in increasing concentrations of an agarose gel, Helmlinger *et al.* [2] found that such spheroids can overcome a mechanical stress of up to 45 mmHg (6 kPa) before they become growth-inhibited at stresses between 45 and 120 mmHg (6 and 16 kPa). Furthermore, in addition to withstanding pressure, neoplastic cells much like normal cells may also exert traction. The traction forces generated by skin fibroblasts have been measured by Delvoye *et al.* [3] via strain-gauge measurements. Using distortable sheets of silicone rubber, Harris *et al.* [4] established a method by which traction forces exerted by individual cells can be visualized. In fact, they distinguish “compression wrinkles,” directly beneath the traction-generating cell, from “tension wrinkles,” which radiate outwards. Interestingly, in their follow-up paper [5] the authors compare various cell types and observe that glia cells from dorsal root ganglia exert very strong traction forces. Since then, a variety of methods and several other gels, including polyacrylamide gels, collagen gels, and basement membrane gels, have been used to study mechanical stresses at the cell-substrate interface [e.g., 6, 7]. For instance, reorganization of basement membrane matrices has been studied extensively with endothelial cells and fibroblasts. Using Matrigel, a commercial basement membrane gel, Vernon *et al.* [8] reported that cellular traction leads to extracellular matrix (ECM) alignment, in which matrix filaments form lines or tracks; this alignment promotes cellular elongation and directs migration. This leads ultimately to the formation of multicellular cords and tubelike structures, characteristic of endothelial cells. Moreover, a quantitative correlation of “contact guidance” with collagen fibril orientation has been described for human fibroblasts [9]. These traction forces can have significance for remodeling connective tissue by tumors as well. In fact, contraction of collagen fibrils has been linked to the metastatic potential of melanoma cells [10] and may also be important for cell locomotion in other cancers, including malignant brain tumors.

Tumor tissue invasion itself is a multistep process, however, including cell attachment, degradation of ECM components, and active cell motility. ECM-degradation in turn requires the expression of proteolytic enzymes by the tumor cells, which has been shown to occur in gliomas *in vitro* and *in vivo* [11, 12]. Interestingly, the parental (high-grade glioma) cell line of the one used in our experiment, U87MG, has been shown to release gelatinase A, known also as metalloproteinase 2 (MMP-2) [12, 13]. As Chintala *et al.* [14] have reported, MMP-2 can degrade a variety of extracellular matrix (ECM) components, including collagen IV, which is a major (30%) ECM-component of the bio-gel used in our assay. A rather complex interplay between mechanical forces and proteolytic enzymes is supported by the work of Vernon and Sage [15], who studied the remodeling of collagen type I gels by bovine aortic endothelial (BAE)

cells. Using zymography to detect enzymatic activity, the authors found that BAE cells that had contracted collagen gels also had secreted MMP-2. They conclude that both cellular traction and proteolysis are important for endothelial cell invasion in the process of angiogenesis.

To elucidate the role of and potential interplay between compression and traction in the dynamic expansion of a micro-tumor system, we implanted a growing multicellular (brain) tumor spheroid (MTS) in a gel matrix and investigated the tumor’s impact on its ECM-gel microenvironment with methods from microrheology. Specifically, we used 1.0 μm beads as reference markers for the gel and monitored their spatial displacement with high-resolution timelapse video-microscopy. In addition, videorate imaging and particle tracking were employed to study the thermally-driven Brownian motions of these tracer beads. In the vicinity of invasive tips, such position fluctuations remain constant in size throughout the observation period. The results show not only that the volumetrically expanding spheroid *pushes* the gel outward but also that beads near the tips of advancing invasive cells are displaced inward, towards these tips. This is consistent with cell *traction* producing tension in the gel. In addition, significant localized gel strain towards invasive cell tips correlates well with a marked increase in overall gel invasion throughout the observation period, indicating that cell traction and invasion are linked. To our knowledge this is the first study presenting evidence that a growing microscopic tumor system can exert *both* significant pressure and traction on its microenvironment.

MATERIALS and METHODS

1. Multicellular Tumor Spheroid and Morphometric Measurements:

The human U87MGmEGFR glioblastoma cell line¹ was used to generate the multicellular tumor spheroids (MTS) [16]. We cultured this cell line in DMEM medium (GIBCO BRL, Life Technologies™, Grand Island, NY) supplemented with 10% heat inactivated cosmic bovine serum (HyClone®, Logan, UT) and 400 $\mu\text{g}/\text{ml}$ G418 (Life Technologies™) in a humidified atmosphere (at 37°C and 5% CO₂). These cells co-express both wild-type epidermal growth factor receptor and its mutant variant, mEGF-R [17]. U87MGmEGFR cells tend to form spheroids in culture after reaching monolayer confluence. Once detached, these spheroids can be collected from the medium with Pasteur pipettes. A variation of the assay used in here is described in Deisboeck *et al.* [18]. In the present case study, we used a rather small spheroid, less than 200 μm in diameter, which we placed in a novel 1×1×1 cm plexiglass cube filled with a 3:1 mixture of growth factor reduced (GFR) matrix, Matrigel® (BIOCAT®, Becton Dickinson, Franklin Lakes, NJ) and (non-serum supplemented) OPTI-MEM® medium at 4 °C, at which temperature the mixture remains fluid. When raised to room temperature, this mixture gels to form a reconstituted basement membrane. Latex beads were dispersed in this mixture at 4 °C while it was still liquid, so that they are incorporated into the gel matrix surrounding the MTS. Maintaining the assay at 37°C and 5% CO₂, tumor growth and invasion as well as the bead-containing microenvironment were imaged at both timelapse (1 frame per 30 seconds) and videorates (30 frames per second), allowing particle tracking and analysis of bead motions. The total observation period was six days. **Figure 1** shows part of the tumor approximately 48 hours

¹ Kindly provided by Dr. W. K. Cavenee (Ludwig Institute for Cancer Research, San Diego, CA).

after implantation. The dark gray area is part of the central MTS; lighter invasive branches, consisting of mobile tumor cells following each other in chainlike migration, are also depicted.

The maximum orthogonal diameters of the MTS were measured at an original magnification of 10× and the average diameter, D_{MTS} , determined with an uncertainty of less than 5 μm , was used to calculate the volume, V_{MTS} , of the MTS core:

$$V_{MTS} = \frac{\rho D_{MTS}^3}{6} \quad (1)$$

The average diameter of the entire tumor system, D_{SYS} , which includes the proliferative MTS core and invasive branches, was determined from measurements of orthogonal diameters of the tips of the invasive chains with an uncertainty of less than 10 μm . From this, the cross-sectional area A_{SYS} was calculated using:

$$A_{SYS} = \rho \left(\frac{D_{SYS}}{2} \right)^2 \quad (2)$$

The cross-sectional area of the invasive region only, A_{INV} , can then be calculated with:

$$A_{INV} = A_{SYS} - A_{MTS} \quad (3)$$

2. Multiparticle Tracking:

Particle tracking techniques, already well-developed for studying soft materials, have the advantage in the present case of allowing nondestructive *in situ* measurements over several days to examine the effects of a growing, invading tumor on its local and extended environments. To act as reference markers, latex beads, 1 μm in diameter, were mixed with the liquid Matrigel solution at 4 °C so that the beads were evenly dispersed before the tumor spheroid was added and gelation induced by bringing the mixture up to room temperature. Carboxylated beads were purchased from Interfacial Dynamics Corporation (Portland, OR) and their surfaces coated with amine-terminated PEG chains, MW = 750 Da, which were covalently coupled to the surface carboxyl groups. The empirical observation that PEG-coated materials resist nonspecific protein adsorption has led to the widespread use of PEG in biomedical applications [19], and we have observed that beads coated with PEG using this protocol are much more resistant to protein adsorption than untreated beads [20]. PEG-coating was found to prevent the beads from aggregating together in the gel, allowing more even dispersion and better tracking accuracy. Resistance to protein adsorption also allows the beads to diffuse thermally within the confines imposed by the Matrigel microenvironments.

During tumor growth, the strain field induced in the gel matrix was mapped, via these probe beads, using **timelapsed images** acquired using a Hamamatsu CCD camera controlled by Metamorph imaging software (Universal Imaging Corporation, Downingtown, PA). In-house IDL particle tracking routines [21] were used to analyze the images and Adobe Photoshop 6.0 was used for further image processing. In **Figures 4-7**, the color superimposed at a position

along a bead track indicates the time elapsed from the beginning of the timelapse sequence until the bead was at that position: for each timelapse sequence, each frame has been assigned an unique color value based on its order within the sequence, starting at indigo-blue for early times and shading along the spectrum to red at late times. The increment in color value per frame is adjusted according to the number of frames in the sequence, so that the scaling of color with elapsed time is different for each timelapse sequence.

Furthermore, **videorate images** of the same field of view were taken using a Sony SVO-9500MD videocassette recorder immediately before beginning and immediately after completing most timelapse sequences; this allowed analysis of the beads’ thermal motions. We were able, over the course of the study, to examine Brownian bead position fluctuations near and far from the growing tumor and at the tips of invasive branches. The Brownian position fluctuations of embedded beads have been used to study the microrheology [22] as well as the local microenvironments [23] of inhomogeneous soft materials.

We measured the bulk elastic modulus, G' , of the Matrigel-based assay, using a temperature-controlled **strain-controlled rheometer** (C-VOR rheometer, Bohlin Instruments, East Brunswick, NJ), to be 20-40 Pa for frequencies from 0.05 to 100 rad/s. The viscous modulus was an order of magnitude smaller than the elastic modulus for most of the frequency range examined, indicating that the gel is predominantly a *solid*.

RESULTS

1. Morphometric Measurements:

Over the six-day observation period the MTS **volume** grows from 0.004 mm^3 to 0.009 mm^3 (**Figure 2**) in three phases: initial growth, phase **I** (Day 1-3); plateaued volume, phase **II** (Day 3-4); rapid secondary growth, phase **III** (Day 4-6). The **invasion** cross-section area increases from 0 to 0.10 mm^2 (**Figure 3**) over these six days, showing its most rapid expansion between Days 3 and 4. This marked increase in invasion area precedes the secondary volumetric growth (**III**); similarly, the invasion area plateau (Day 2-3) precedes the MTS volume plateau (**II**).

2. Timelapse Multiparticle Tracking:

As the tumor system grows volumetrically and invades the surrounding gel matrix, timelapse image sequences show the displacements of beads, embedded in the gel, following local gel motion. These reference markers allow the time-dependent strain field induced by the tumor to be mapped, at a variety of locations, over the period of observation.

Four hours after implantation (Day 1), bead tracks in **Figure 4** depict gel movement radially inward toward the tumor. Tracks show movement with a circumferential component, along a line tangential to the tumor surface, as well as a radial component (upper center of image) at a location where an early invasive cell is later seen to emerge. Significant tumor growth begins about 6 hours after implantation. The increase in MTS volume (see **Figure 2**) displaces the gel radially *outward* (**Figure 5**). Tracks show more circumferential movement (middle right of image) at a location where a cluster of invasion pathways has appeared by the end of this 21-hour timelapse sequence. Twenty-four hours after implantation, MTS growth continues to displace gel radially outward (**Figure 6**). However, an emergent invasion pathway

(center of image) is associated with a local gel strain with a significant component radially inward, toward the tip of the invading cell and opposite to the bulk gel movement. In **Figure 7**, three days after implantation of the MTS (Day 4) and approximately 65 μm from the spheroid edge, gel near the tip of an invasive branch moves a significant distance *in, toward* the leading invasive cell. Meanwhile, the bulk surrounding gel is only slightly displaced outward. The magnitude and spatial extent of inward strain associated with the invasion is highly directional and not isotropic about the cell tip. Beads more than 30 μm from the leading tip of the invasive cell along the line of invasion exhibit significant local inward strain.

Gel strain inward toward invasive cell tips is consistently observed throughout this study. Established invasion pathways close to the MTS seem to induce gel strain outward, along the direction of invasion.

3. Mean Square Displacement (MSD) of Beads :

Throughout the period of observation, the bead-averaged mean square displacement (MSD) for groups of Brownian beads near invasive tips, at a lag time of 0.3 seconds averaged over 3 to 6 minutes, remained within the range 0.012-0.022 μm^2 showing displacements much less than the 1 μm bead diameter.

DISCUSSION

The combination of this 3D-ECM gel assay with analytical methods from microrheology and soft condensed matter science yields novel and important insights into the dynamic interaction between tumor and microenvironment. The timelapsed imaging of tracer beads embedded in the gel matrix allows direct mapping of the displacement field induced by the tumor within its microenvironment and clearly shows that the volumetrically expanding tumor spheroid displaces the bulk matrix radially outward while gel near the tips of invasive cells is pulled inward toward those tips. Since such gel consists of interconnected sheets of proteins [24], these phenomena should be attributed to *stress propagation*, i.e. movement of distant beads results from transmission of the local mechanical effect exerted by the expanding tumor or by the tip of an invasive branch. In the following, we will discuss observations of the microtumor system expanding within the 3D environment, on both *single-cell* and *multicellular* scales, as well as implications of the material properties found in our investigation of the tumor-free Matrigel-based gel mixture.

Qualitative observations of gel displacement and cell traction are not all that can be obtained from these timelapse sequences. Equipped with knowledge of the gel’s bulk viscoelastic properties, we can *quantitatively* estimate the traction exerted by invasive cells. **Figure 7**, depicting the result of a timelapse sequence acquired for 165 minutes during a period of rapid invasion and negligible volumetric growth, shows that beads 7-14 μm from the leading invasive cell tip (tracks 2 and 3) are displaced about 8 μm in toward the tip. One bead about 30 μm from the tip (track 1) is displaced about 5 μm . Gel 46 μm from the cell tip is effectively motionless, probably as a result of the superposition of the invasive tip’s inward pull and the growing spheroid’s outward displacement, as shown in **Figure 7** by the beads most distant from the invasive cell. We therefore treat the matrix at a distance of 46 μm from the cell tip as fixed and can approximate the unstrained length of gel at a distance r from the cell tip as $L = 46 \mu\text{m} -$

r . From this, we find the strain $\Delta L/L$ induced in the gel by the cell tip is approximately $8\text{mm}/(46\text{mm} - 14\text{mm}) \approx 5\text{mm}/(46\text{mm} - 30\text{mm}) \approx 0.3$. Using the bulk elastic modulus (G) of about 30 Pa, and neglecting the viscous modulus since it is about an order of magnitude smaller, we find that this invasive cell tip has exerted traction of $G^*(\Delta L/L) \approx 9$ Pa, plus an additional term correcting for the outward pressure resulting from MTS expansion.

This corrective term should be quite small, however, as we may see if we approximate that the cell tip applies a force at a point in an infinite elastic medium. At about 46 μm from the cell tip, the cell tip traction should be equal in magnitude and opposite in sign to the outward pressure caused by MTS volumetric expansion, since we observe negligible displacement of the gel at this distance. The stress resulting from the application of such a force is inversely proportional to the square of the distance from the point of application [25]. We therefore approximate that the traction caused by the cell tip is much greater than the outward pressure at this location. To put it in perspective, the traction applied by the glioma cell tip shown here is three orders of magnitude less than the maximum traction measured during fibroblast locomotion on an elastic substrate [6] and approximately 1-2 orders of magnitude less than the maximum tractions measured at the front of migrating fibroblasts using a novel micromachined force transducer [26]. Interestingly, stronger traction does not always correlate with greater cell motility, however, and in fact Harris and coworkers [5] found in their studies of various differentiated cell types that traction strength was weakest in the most mobile and invasive cells studied.

If we treat the traction-induced bead displacements as the result of a point force, applied at the invasive cell tip in a direction determined by averaging displacement vectors, we can estimate the magnitude of this force using the equation of equilibrium for a three-dimensional elastic solid. A displacement \mathbf{u} will have the form

$$\bar{u} = \frac{1 + \mathbf{n}}{8pE(1 - \mathbf{n})} \frac{(3 - 4\mathbf{n})\bar{F} + \bar{n}(\bar{n} \cdot \bar{F})}{r} \quad (4)$$

where $r = (x^2 + z^2)^{1/2}$ is the distance from the origin, where the force is applied, E is the elastic modulus, and \mathbf{n} is Poisson’s ratio [27]. For a true point force, equation (4) should be valid at all locations where r is large compared to the dimension of the region where the force is applied – in this case, at distances more than a few microns from the invasive tip. If we choose coordinate axes such that $\mathbf{F} = F\mathbf{z}$, the orthogonal components of $\mathbf{u} = w\mathbf{z} + v\mathbf{x}$ can be simplified to yield

$$w = \frac{F(1 + \mathbf{n})}{8pE(1 - \mathbf{n})r} \left(3 - 4\mathbf{n} + \frac{z^2}{r^2}\right) \quad (5)$$

and

$$v = \frac{F(1 + \mathbf{n})}{8pE(1 - \mathbf{n})} \frac{xz}{r^3} \quad (6)$$

Setting \mathbf{n} to $1/2$, Poisson’s ratio for an incompressible medium, the magnitude of the point force F applied over the timelapse period can be approximated from observed displacements. From w and v components of the four bead tracks (1-4) indicated in **Figure 7**, we estimate that, over the 165 minutes shown in this timelapse, this invading cell tip pulls nearby gel inward with a force in

the range 10-100 nN. The forces thus estimated are significantly less than those reported for fibroblasts as measured on elastic substrates, $\sim 2 \mu\text{N}$ [6, 28], but only about 2-10 times less than those measured at the front of a migrating fibroblast [26] and comparable to the force, up to 30 nN, applied at a single focal adhesion by human foreskin and cardiac fibroblasts [29].

However, we note that a cell likely consumes some of the surrounding gel as it invades, so that gel volume is not conserved and the system is not truly Hookean. Furthermore, the way in which proteolytic enzymes may change the gel’s elastic properties has not been examined here. Better characterizations of the effects, if any, of enzymatic proteolysis on gel viscoelastic properties and of the rate and spatial distribution of gel consumption by invasive cells will allow refinement of these traction and force estimations, as will measurement of the gel’s true Poisson ratio. Moreover, as the cell invades the gel, the point at which the inward force is applied, which we have taken as the origin of our coordinate system above, should move outward with the cell as well. Indeed, this cell-ward force does not seem to be, in fact, a true point force. A closer examination of **Figure 7** reveals that not all beads shown to the right of the invasive cell are displaced in toward the cell, nor are the displacements of those beads that do move inward all consistent with the displacement field described by equations (5) and (6). A more sophisticated treatment, incorporating multiple, moving force origins as well as a model of cell adhesion to and exertion of traction on gel filaments, might significantly improve interpretation of these observations.

On a larger length and longer time scale, observation of the tumor system shows that both spheroid volume and invasion area increase over the observation period. However, these increases are neither monotonic nor synchronized; rather, a marked increase in invasion area (Day 3-4) trails a rapid gain in volume (Day 1-3) and precedes a second rapid volumetric expansion (Day 4-6), as shown in **Figures 2** and **3**. A similar pattern has been described previously by Deisboeck *et al.* [18], and present observations are in agreement with the notion that a *feedback* mechanism may link volumetric growth and invasive expansion. As indicated by timelapsed observation of bead displacement far from the MTS, the mechanical impact of the microscopic tumor on its environment reaches well beyond a distance, measured from the MTS edge, greater than twice the initial radius of the MTS. Since the clinically relevant, macroscopic situation presents tumor radii of several centimeters [30] and since tumor cells have been found *in vivo* at a distance greater than 4 cm from the gross tumor [31], a cautious extrapolation of these findings to the clinical situation suggests that the tumor’s mechanical impact may be exerted throughout, and perhaps even beyond, the entire ipsilateral brain hemisphere. Interestingly, this notion is already supported by preliminary results from a brain tumor patient, found using specific diffusion-tensor MR-imaging, showing tumor-related changes in the diffusion anisotropy of water throughout the entire brain [32].

Furthermore, we find that marked ECM-gel *invasion* appears to be correlated with tumor cell *traction*. This corresponds well with the findings of Klein *et al.* [10], who report that two highly aggressive melanoma cell lines efficiently contract 3D collagen type I gels and that their synthesis of $\alpha_2\beta_1$ integrins is upregulated. The upregulation of such cell adhesion receptors is important in this context since integrins mediate also in malignant brain tumor cells both interaction with ECM components and invasiveness [33]. However, the identity of specific mechanism(s) used by tumor cell-generated traction to facilitate directed movement is still unclear. Vernon and Sage [15], for example, propose that traction-mediated alignment of an ECM may produce specific pathways that other cells follow. Davis and Camarillo [34] suggest a similar concept drawn from their studies of endothelial cells, which caused linear distortions of

Matrigel; these distortions correspond to the migration pathways of endothelial cell processes. These authors call this phenomenon “matrix guidance pathways” and say it may result from the generation of tension between endothelial cells. Tranquillo [35] presented a similar concept for fibroblasts, proposing that “cells align, exert traction and migrate preferentially in the direction in which surrounding fibrils are aligned.” Deisboeck *et al.* [18] presented a “least resistance, most permission, highest attraction” concept for the emergence of invasive branching patterns: brain tumor cells would follow each other because of increased chemical attraction, enhanced haptotactic permission as well as reduced mechanical resistance within a preformed path. The aforementioned ‘matrix guidance’ would thus be in accordance with this concept as it also facilitates at least the haptotactic element. Our timelapse observations indicate that the traction-induced tension associated with invasion is highly localized at the tips of invasive branches and not alongside established invasive branches closer to the MTS. The fairly constant average thermal MSD of beads within the area near such invasive cell tips may argue for a relatively minor role of proteolytic enzymes in the tip area, since substantial matrix degradation would be expected to result in an increase in thermally-driven bead fluctuations. However, since the parental cell line (U87MG) has been shown to secrete MMP-2, which degrades collagen IV, a component of GFR-Matrigel, and since the release of MMP-2 has been linked to collagen contraction by endothelial cells [13, 14, 15], the possibility that proteolytic enzymes may contribute to tumor invasion also in the present study cannot be eliminated. This is further supported by the findings of Vaithilingam *et al.* [36], who report an (primarily extracellular) increase of general proteolytic activity in C6 astrocytoma spheroids with increasing spheroid diameter.

The 3D-assay system used in the current study does have potential shortcomings. For example, the presence of some tissue culture medium around the spheroid when the MTS is implanted cannot be avoided. The early inward gel movement depicted in **Figure 4** probably results from the rapid metabolism of traces of incubation medium by highly proliferative MTS surface cells. Generally, a higher incubation medium concentration might produce an immediate environment that is more favorable for volumetric tumor growth and hence result in the first marked increase of MTS volume **(1)** in **Figure 2**. A loss of fluid from the gel towards the end of the observation period is another concern and might indicate a rather serious limitation of the assay, since the loss of fluid and the resulting increased rigidity of the gel may render forward cell movement more difficult. To address this issue, it is helpful to look carefully at the invasive behavior, depicted in **Figure 3**. The invasion area clearly still continues to increase towards the end of the observation period, which suggests a relatively minor influence of drying on cell motility.

In summary, we have demonstrated that an expanding, microscopic tumor system exerts significant mechanical forces upon its environment, in this case a particular extracellular matrix gel-composition placed inside a novel plexiglass cube. More specifically: **(1)** The multicellular brain tumor spheroid system exerts both compressive pressure *and* tension on its microenvironment. **(2)** These mechanical forces can be linked to the tumor’s proliferative and invasive growth dynamics, which seem to induce each other. **(3)** Spatially, the mechanical impact of the volumetrically growing tumor propagates to at least twice its initial radius. **(4)** The growing MTS exerts this outward pressure, but inward traction is generated by invasive tips only. **(5)** Furthermore, this cell traction does not relax while volumetric growth of the MTS ceases. Also, the invasive tip maintains this tension for five consecutive days during which the

overall invasive area increases substantially. Cell-imposed traction and tumor invasion therefore seem related.

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REFERENCES

1. CBTRUS, Central Brain Tumor Registry of the United States, Statistical Report, 1992-1997.
2. Helmlinger, G., Netti, P. A., Lichtenfeld, H. C., Melder, R. J., and Jain, R. K. (1997). Solid stress inhibits the growth of multicellular tumor spheroids. *Nat. Biotechnol.* **15**, 778.
3. Delvoye, P., Wiliquet, P., Leveque, J.-L., Nusgens, B. V., and Lapiere, C. M. (1991). Measurement of mechanical forces generated by skin fibroblasts embedded in a three-dimensional collagen gel. *J. Invest. Dermatol.* **97**, 898.
4. Harris, A. K., Wild, P., and Stopak, D. (1980). Silicon rubber substrata: a new wrinkle in the study of cell locomotion. *Science* **208**, 177.
5. Harris, A. K., Stopak, D., and Wild, P. (1981). Fibroblast traction as a mechanism for collagen morphogenesis. *Nature* **290**, 249.
6. Dembo, M., and Wang, Y.-L. (1999). Stresses at the cell-to-substrate interface during locomotion of fibroblasts. *Biophys. J.* **76**, 2307.
7. Jenkins, G., Redwood, K. L., Meadows, L., and Green, M. R. (1999). Effect of gel re-organization and tensional forces on $\alpha 2\beta 1$ integrin levels in dermal fibroblasts. *Eur. J. Biochem.* **263**, 93.
8. Vernon, R. B., Angello, J. C., Iruela-Arispe, M. L., Lane, T. F., and Sage, E. H. (1992). Reorganization of basement membrane matrices by cellular traction promotes the formation of cellular networks in vitro. *Lab. Invest.* **66**, 536.
9. Guido, S., and Tranquillo, R. T. (1993). A methodology for the systematic and quantitative study of cell contact guidance in oriented collagen gels. Correlation of fibroblast orientation and gel birefringence. *J. Cell Sci.* **105**, 317.

10. Klein, C. E., Dressel, D., Steinmayer, T., Mauch, C., Eckes, B., Krieg, T., Bankert, R. B., and Weber, L. (1991). Integrin $\alpha 2\beta 1$ is upregulated in fibroblasts and highly aggressive melanoma cells in three-dimensional collagen lattices and mediates the reorganization of collagen I fibrils. *J. Cell Biol.* **115**, 1427.
11. Apodaca, G., Rutka, J. T., Bouhana, K., Berens, M. E., Giblin, J. R., Rosenblum, M. L., McKerrow, J. H., and Banda, M. J. (1990). Expression of metalloproteinases and metalloproteinase inhibitors by fetal astrocytes and glioma cells. *Cancer Res.* **50**, 2322.
12. Nakano, A., Tani, E., Miyazaki, K., Yamamoto, Y., and Furuyama, J. I. (1995). Matrix metalloproteinases and tissue inhibitors of metalloproteinases in human gliomas. *J. Neurosurg.* **83**, 298.
13. Uhm, J. H., Dooley, N. P., Villemure, J. G., and Yong, V. W. (1996). Glioma invasion in vitro: regulation by matrix metalloproteinase-2 and protein kinase C. *Clin. Exp. Metastasis* **14**, 421.
14. Chintala, S. K., Tonn, J. C., Rao, J. S. (1999). Matrix metalloproteinases and their biological function in human gliomas. *Int. J. Devl. Neurosciences* **17**, 495.
15. Vernon, R. B., and Sage, E. H. (1996). Contraction of fibrillar type I collagen by endothelial cells: a study in vitro. *J. Cell Biochem.* **60**, 185.
16. Sutherland, R. M. (1988). Cell and environment interactions in tumor microregions: the multicell spheroid model. *Science* **240**, 177.
17. Nishikawa, R., Ji, X. D., Harmon, R. C., Lazar, C. S., Gill, G. N., Cavenee, W. K., and Huang, H. J. (1994). A mutant epidermal growth factor receptor common in human glioma confers enhanced tumorigenicity. *Cancer Res.* **91**, 7727.
18. Deisboeck, T. S., Berens, M. E., Kansal, A. R., Torquato, S., Stemmer-Rachamimov, A. O., and Chiocca, E. A. (2001). Pattern of self-organization in tumour systems: complex growth dynamics in a novel brain tumour spheroid model. *Cell Proliferation* **34**, 115.
19. Harris, J. M., and Zalipsky, S. (1997). “Poly(ethylene glycol): Chemistry and Biological Applications,” American Chemical Society, Washington, D. C.
20. Valentine, M. T., Perlman, Z., Gardel, M. L., Shin, J., Matsudaira, P. T., Mahadevan, L., Mitchison, T., and Weitz, D. A. (in preparation). Effect of surface chemistry on microrheology experiments: Comparison of BSA-blocks, PEG-coated, and Carboxylated beads.
21. Crocker, J. C., and Grier, D. G. (1996). Methods of digital video microscopy for colloidal studies. *J. Colloid Interface Sci.* **179**, 298.
22. Mason, T. G., and Weitz, D. A. (1995). Optical Measurements of Frequency-Dependent Linear Viscoelastic Moduli of Complex Fluids. *Phys. Rev. Lett.* **74**, 1250.
23. Valentine, M. T., Kaplan, P. D., Thota, D., Crocker, J. C., Gisler, T., Prud’homme, R. K., Beck, M., and Weitz, D. A. (2001). Investigating the microenvironments of inhomogeneous soft materials with multiple particle tracking. *Phys. Rev. E* **64**, 061506.
24. Kleinman, H. K., McGarvey, M. L., Hassell, J. R., Star, V. L., Cannon, F. B., Laurie, G. W., and Martin, G. R. (1986). Basement membrane complexes with biological activity. *Biochemistry* **25**, 312.

25. Timoshenko, S. P., and Goodier, J. N. (1987). “Theory of Elasticity,” 3^d ed., McGraw-Hill, New York.
26. Galbraith, C. G., and Sheetz, M. P. (1997). A micromachined device provides a new bend on fibroblast traction forces. *Proc. Natl. Acad. Sci. USA*, **94**, 9114.
27. Landau, L. D., and Lifshitz, E. M. (1986). “Theory of Elasticity,” 3^d edition. Pergamon Books Ltd., New York.
28. Wrobel, L. K., Fray, T. R., Molloy, J. E., Adams, J. J., Armitage, M. P., Sparrow, J. C. (2002). Contractility of single human dermal myofibroblasts and fibroblasts. *Cell Motil. Cytoskeleton* **52**, 82.
29. Balaban, N. Q., Schwarz, U. S., Riveline, D., Goichberg, P., Tzur, G., Sabanay, I., Mahalu, D., Safran, S., Bershadsky, A., Addadi, L. and Geiger, B. (2001). Force and focal adhesion assembly: a close relationship studied using elastic micropatterned substrates. *Nature Cell Biol.* **3**, 466.
30. Kansal, A. R., Torquato, S., Harsh IV, G. R., Chiocca, E. A., and Deisboeck, T. S. (2000). Simulated brain tumor growth dynamics using a three-dimensional cellular automaton. *J. Theor. Biol.* **203**, 367.
31. Silbergeld, D. L., and Chicoine, M. R., (1997). Isolation and characterization of human malignant glioma cells from histologically normal brain. *J. Neurosurg.* **86**, 525.
32. Zhang, S., Laidlaw, D. H., Bastin, M. E., Sinha, S., and Deisboeck, T. S. (submitted). Computational visualization and analysis of structural heterogeneity in a diffusion tensor MRI-data set from a brain tumor patient.
33. Paulus, W., and Tonn, J. C. (1994). Basement membrane invasion of glioma cells is mediated by integrin receptors. *J. Neurosurg.* **80**, 515.
34. Davis, G. E., and Camarillo, C. W. (1995). Regulation of endothelial cell morphogenesis by integrins, mechanical forces, and matrix guidance pathways. *Exp. Cell Res.* **216**, 113.
35. Tranquillo, R. T. (1999). Self-organization of tissue equivalents: the nature and role of contact guidance. *Biochem. Soc. Symp.* **65**, 27.
36. Vaithilingam, I. S., Stroude, E. C., McDonald, W., and Del Maestro, R. F. (1991). General protease and collagenase (IV) activity in C6 astrocytoma cells, C6 spheroids and implanted C6 spheroids. *J. Neuro-Onc.* **10**, 203.

FIGURE CAPTIONS

Figure 1: Multicellular Tumor Spheroid (MTS) with invasive branches. Taken on **Day 3**, the image displays a part of the main tumor (dark gray) and shows invasive branches originating from the surface of the MTS. Forming these branches, invasive cells advance in a chain-like pattern. (Original Magnification: 10×).

Figure 2: MTS Volume. Volume of the tumor spheroid over 6 days of observation is shown. The gray shading and Roman numerals indicate the three growth phases described in the text.

Figure 3: MTS Invasion Area. Total invasion cross-section area, exclusive of the central MTS core, over the 6 days of observation is shown.

Figure 4: Timelapse Multiparticle Tracking (Day 1). Tracks are drawn to trace the paths of moving beads as color is used to timestamp bead positions: early times are indigo-blue and colors shade along the spectrum to red at late times. The tracks have been superimposed on the first frame in the timelapse sequence. Note darkened multicellular tumor spheroid top, right. Ticks indicate acquired image dimensions, ten pixels per tick mark. (Bar = 10 μm ; Original Magnification: 40×; Timelapse duration: 110 min).

Figure 5: Timelapse Multiparticle Tracking (Day 1). For orientation, note darkened MTS bottom, right. The MTS radius, determined from the colored arcs tracing the outgrowing spheroid edge, increased by more than 10 μm while this sequence was acquired. (Bar = 10 μm ; Original Magnification: 40×; Timelapse duration: 832 min).

Figure 6: Timelapse Multiparticle Tracking (Day 2). Note a single invasive branch emerging from the darkened MTS. Beads in the bulk gel, not near the invasive cell, show 3-5 μm displacement radially outward. The displacements of beads near the invasive cell, however, show a significant component radially inward, as much as 2 μm in some cases, as bead paths arc over the course of this acquisition. (Bar = 10 μm ; Original Magnification: 40×; Timelapse duration: 224 min).

Figure 7: Timelapse Multiparticle Tracking (Day 4). Note the elongated single cell at the tip of the cell branch invading from top, right, indicated by the *arrow*. Beads 7-14 μm from the cell tip (tracks 2 and 3) are displaced about 8 μm in toward the tip over this timelapse observation. A bead (center, right) about 30 μm from the cell tip (track 1) is displaced about 5 μm , while beads 46 μm and more distant from the tip do not show significant displacement inward. A bead located along a line almost orthogonal to the line of invasion (track 4) shows displacement toward the tip much greater than would be expected to result from a true point force producing the displacements observed for beads located along the line of invasion (tracks 1-3). Beads very far from the invasive tip show a slight displacement outward. (Bar = 10 μm ; Original Magnification: 40×; Timelapse duration: 165 min).

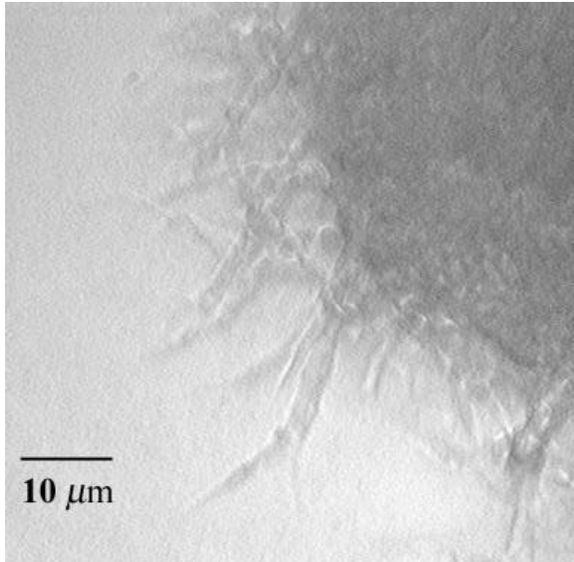


Figure 1

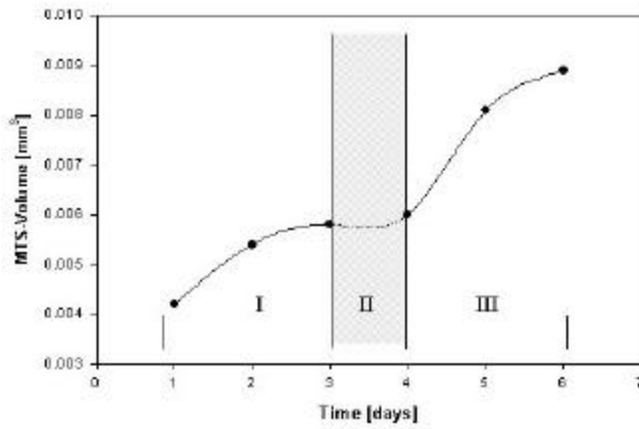


Figure 2

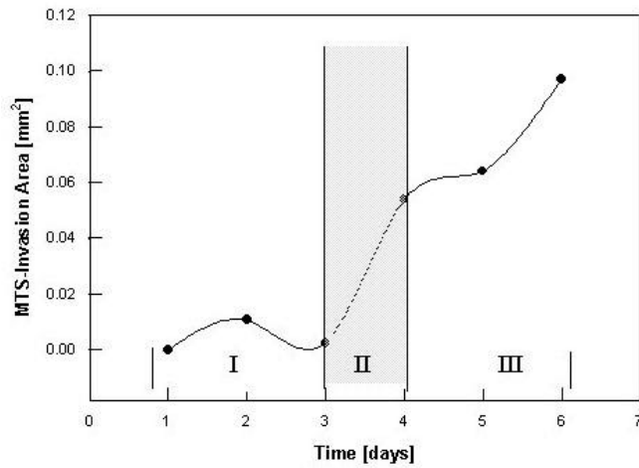


Figure 3

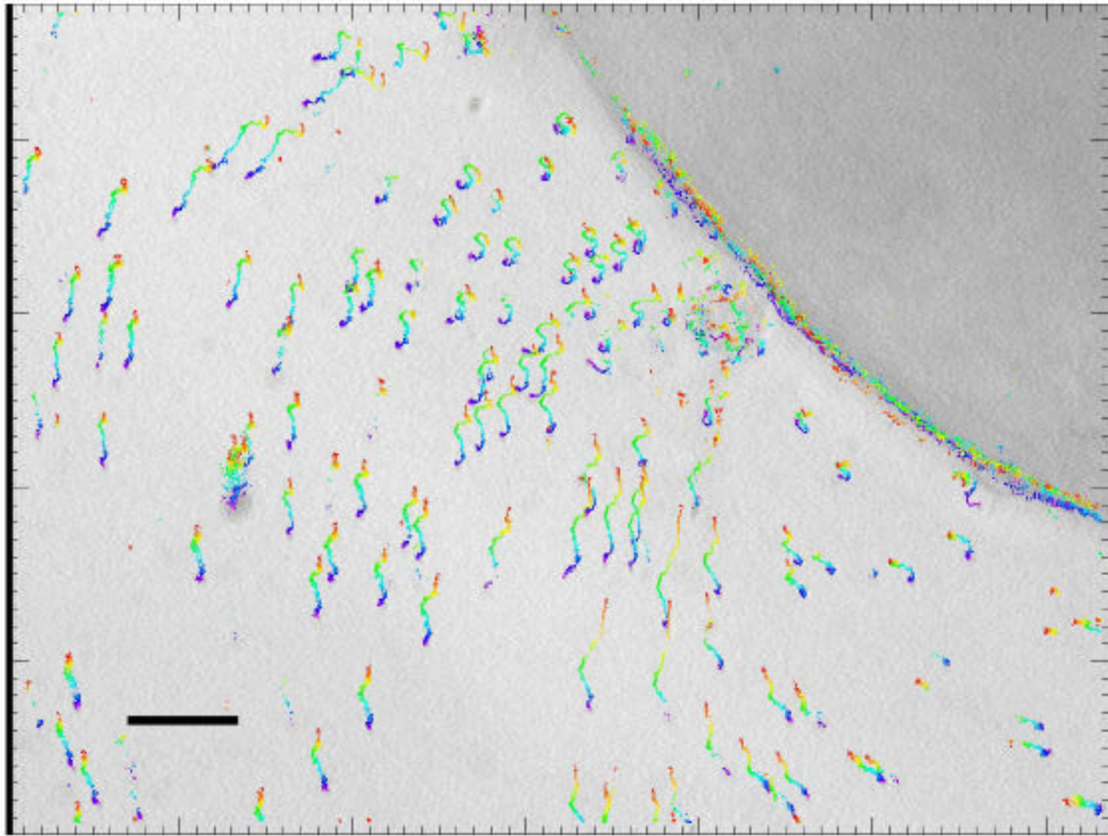


Figure 4

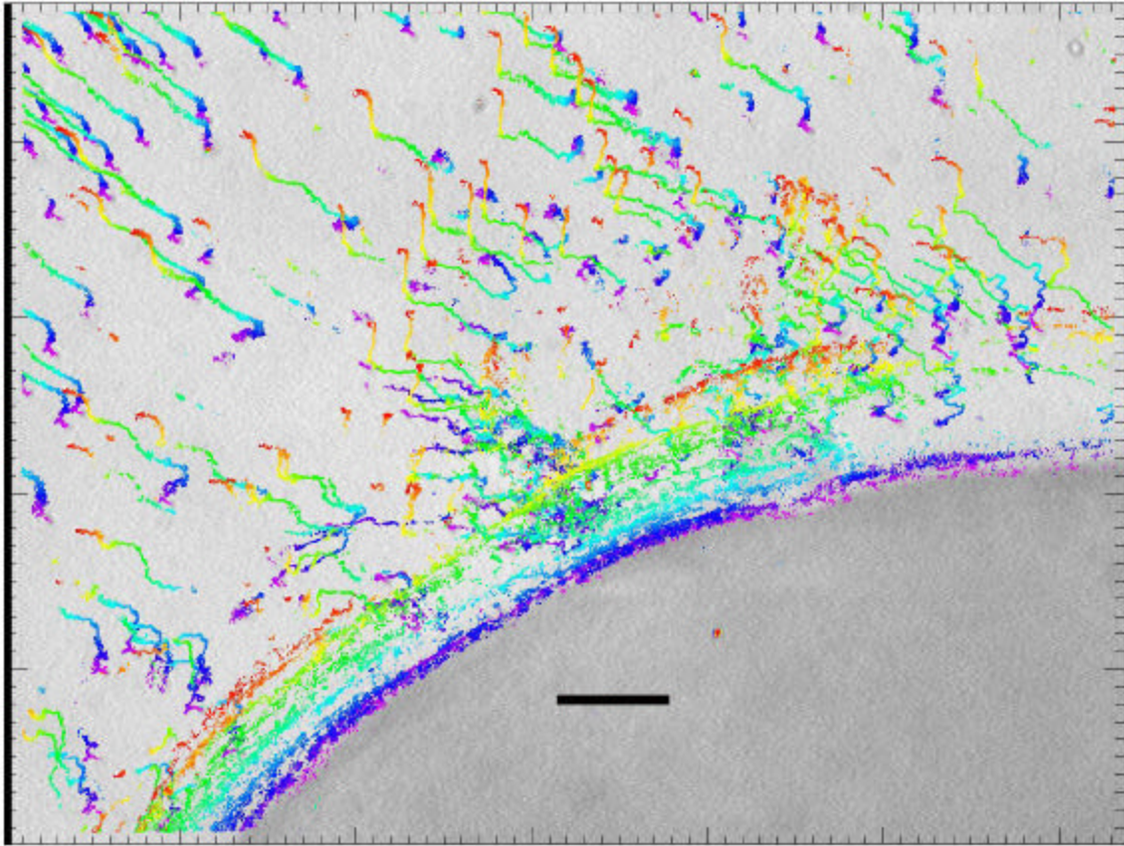


Figure 5

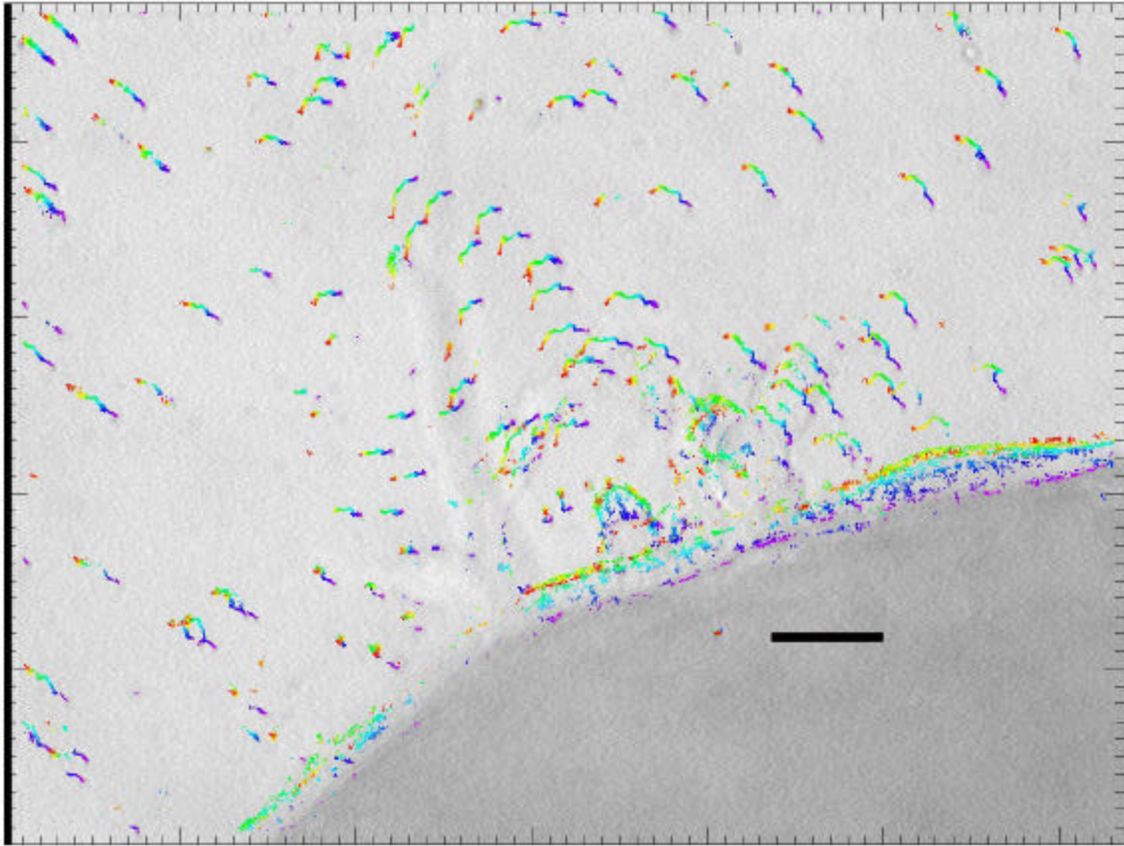


Figure 6

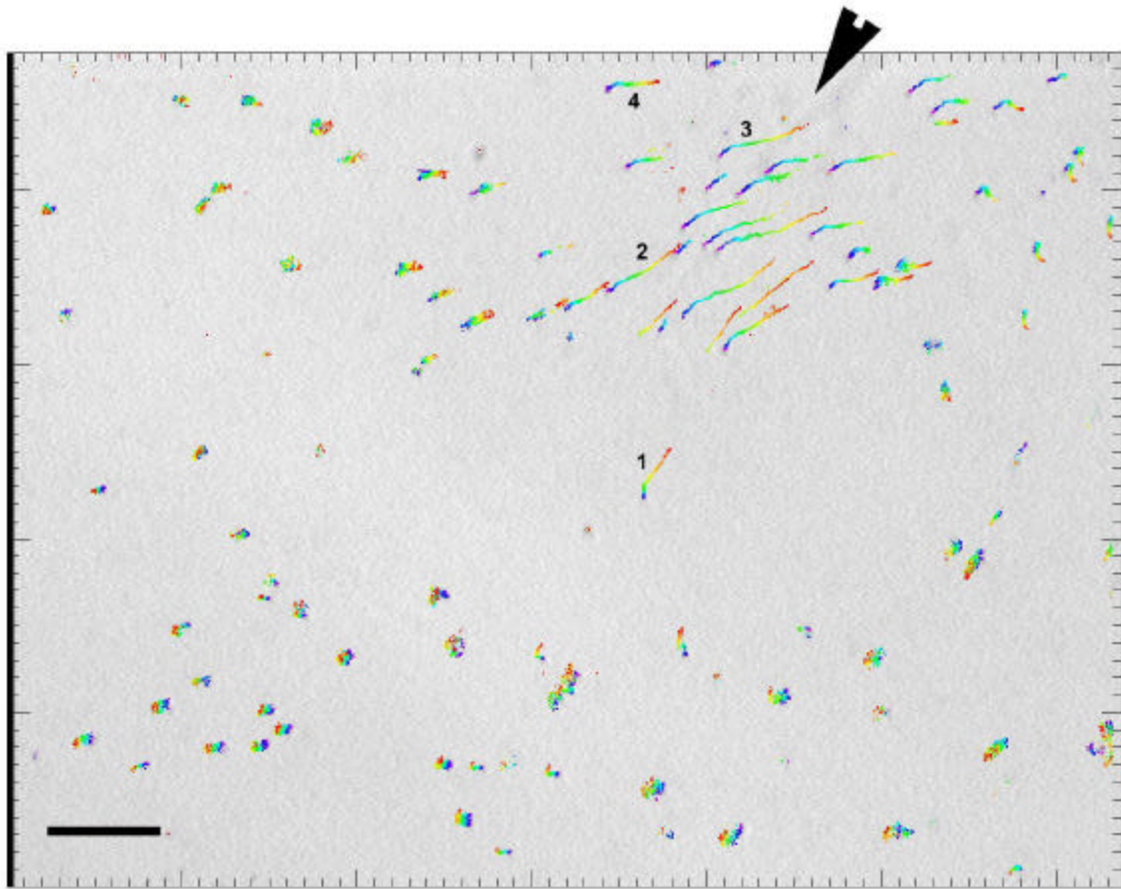


Figure 7