Aligned Nanotopography Promotes a Migratory State in Glioblastoma Multiforme Tumor Cells

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Glioblastoma multiforme (GBM) is an aggressive, Grade IV astrocytoma with a poor survival rate, primarily due to the GBM tumor cells migrating away from the primary tumor site along the nanotopography of white matter tracts and blood vessels. It is unclear whether this nanotopography influences the biomechanical properties (i.e. cytoskeletal stiffness) of GBM tumor cells. Although GBM tumor cells have an innate propensity to migrate, we believe this capability is enhanced due to the influence of nanotopography on the tumor cells’ biomechanical properties. In this study, we used an aligned nanofiber film that mimics the nanotopography in the tumor microenvironment to investigate the mechanical properties of GBM tumor cells in vitro. The data demonstrate that the cytoskeletal stiffness, cell traction stress, and focal adhesion area were significantly lower in the GBM tumor cells compared to healthy astrocytes. Moreover, the cytoskeletal stiffness was significantly reduced when cultured on aligned nanofiber films compared to smooth and randomly aligned nanofiber films. Gene expression analysis showed that tumor cells cultured on the aligned nanotopography upregulated key migratory genes and downregulated key proliferative genes. Therefore, our data suggest that the migratory potential is elevated when GBM tumor cells are migrating along aligned nanotopographical substrates.

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Greater Cytoskeletal Stiffness Observed in Astrocytes than in GBM Tumor Cells. As GBM is categorized as a Grade IV astrocytoma, the difference in cytoskeletal stiffness between GBM tumor cells and non-cancerous healthy primary astrocytes was measured using atomic force microscopy (AFM). The cytoskeletal stiffness was tested on two GBM cell lines (U87MG and A172), and primary GBM CICs (BT145). Conventional thought is that primary GBM tumor cells were derived directly from genetically mutated astrocytes or glial precursor cells (i.e. EGFR amplification/mutation, PTEN loss/mutation, etc.)36. Therefore, primary rat post-natal day 2 astrocytes and mouse neural stem cells were used as the non-cancerous, healthy cells. Average stiffness measurements and representative images for each cell type are shown in Fig. 1. Astrocytes were significantly stiffer than each GBM tumor cell type, with an average stiffness of $4184 \pm 102.3$ Pa ($p < 0.0001$). There was no statistical difference between the two GBM cell lines, U87MG and A172 tumor cells, which had an average stiffness of $1315 \pm 39.98$ Pa and $1138 \pm 68.58$ Pa, respectively. Primary GBM CICs, BT145, were statistically less stiff than the primary astrocytes and the GBM tumor cell lines ($p < 0.01$), with an average stiffness of $653.3 \pm 35.37$ Pa (Fig. 1A). Finally, NSCs had a similar stiffness to the CICs when plated on laminin (data not shown). Morphologies of the cells were also noticeably different between the various cell types (Fig. 1D). Astrocyte morphology was more spread on the TCPs compared to the spindled morphology exhibited in the tumor cells.

As a healthy cell undergoes oncogenic mutations, many cellular attributes are altered resulting in the abnormal growth and migratory behavior of cells15. Due to the role of the cytoskeleton on cell migration, previous studies identified biophysical attributes, specifically cytoskeletal stiffness, as a biomarker for invasive potential in a variety of cancers. Tumor studies using AFM to measure cytoskeletal stiffness have shown that more invasive breast, ovarian, and prostate cancer cells are less stiff than their less invasive, benign, or healthy cell counterparts18,20,37. Significant reductions in stiffness, 2–5 fold, were observed between immortalized ovarian surface epithelial cells and different ovarian cancer cell lines20. In addition, Andolfi et al. showed that high and low grade glioma stem cells are less stiff than the non-tumorigenic glioma-associated-stem cells23. Similar results were observed in our experiments with the GBM tumor cells, as the highly invasive CICs were more compliant than the U87MG and A172 tumor cell lines, which together, were significantly softer than non-cancerous astrocytes. During invasion, cancer cells need to deform their bodies in order to conform to the surrounding tissue and migrate. The reduced stiffness indicates a greater capability to deform within the aggressive tumor cells, which facilitates the migration and invasion through the surrounding ECM leading to secondary tumor sites15. In addition, NSCs have also been shown to be highly migratory within the brain. Due to their enhanced migratory capacity toward cancerous/diseased tissue, these cells have been researched as a tool to deliver therapies to tumor cells34,39.
Cellular Traction Stress Greater in Astrocytes than GBM Tumor Cells. Average cell traction stress of the primary astrocytes, U87MG, A172, and BT145 were measured using traction force microscopy (TFM) (Fig. 2). Cells were cultured overnight on polyacrylamide gels before being measured. For the primary astrocytes, a stiffer gel was used to measure the cell traction forces as the softer gels used for the tumor cells were too compliant and were pulled by the extremely contractile astrocytes, thus causing the beads to be in a different focal plane than the rest of the gel. This resulted in inaccurate measurements (data not shown). Therefore, a stiffer gel using 8% acrylamide and 0.1%-bis-acrylamide was used for the astrocytes. As the BT145 tumor cells did not adhere well or generate traction on the aforementioned gel, the BT145 tumor cells were plated on a 5% acrylamide and 0.08%-bis-acrylamide gel. As gel stiffness influences the traction force generated, U87MG and A172 tumor cells were plated on both the 8% and 5% gel, and compared with the traction forces from the astrocytes and BT145 tumor cells, respectively. Cell traction forces were normalized to respective projected cell area to determine average cell traction stress.

Figure 2A illustrates that non-cancerous astrocytes exerted significantly greater cell traction stress on the 8% acrylamide gel compared to the malignant GBM tumor cells. The average cell traction stress exerted by the astrocytes was $530 \pm 56.5$ Pa compared to $91.2 \pm 14.7$ Pa exerted by the malignant GBM tumor cells ($p < 0.0001$). Further, highly invasive CICs were significantly less stiff than less invasive GBM cell lines ($p < 0.05$). (B) After differentiation CICs with FBS, stiffness of cells significantly increased 2-fold ($p < 0.01$). (C) Representative brightfield images of astrocytes and tumor cells. N = 3, mean ± SEM.
Cellular traction forces have also been investigated in other cancers to demonstrate how differing traction stresses can correlate to the tumor cells' invasive potential; however, conflicting results have been reported. Similar to what was observed in this study, an inverse relationship between traction stress and cell metastatic potential in breast cancer cells was observed\(^4\). Similarly, MacKay et al. treated GBM tumor cell lines with constitutively active (CA) RhoA and found that the increased expression in RhoA led to increased cell traction forces and decreased migration, which was probably due to the tumor cells adhering too strongly to the substratum\(^2\). However, other studies that investigated transformed fibroblasts, breast, prostate, and lung cancer cells, showed the opposite trend, as these cell types exerted significantly more total cell traction force than their non-metastatic counterparts\(^4\). One potential reason for these conflicting results is the differing native microenvironments between the different cell types. The brain has a highly specialized ECM. In addition to being considerably softer than other tissues, the brain has a lower percentage of fibrous proteins and an abundance of hyaluronic acid compared to other tissues\(^3\). Cell types will likely respond to varying substrates differently. It has been previously reported that substrate stiffness affects both migration speed and traction forces in a biphasic manner, with an optimal stiffness that promotes the migration speed varying between different cell types, suggesting that migration and traction forces may be linked due to the cells' innate environment\(^5\).

**Actin and Focal Adhesion Analysis of Astrocytes and GBM Tumor Cells Demonstrate that Tumor Cells Exhibit Reduced Focal Adhesion Presence.** To determine if there were differences in cytoskeletal organization between astrocytes and malignant tumor cells, cells were stained for vinculin (red), a focal adhesion protein, as well as F-actin (green) (Fig. 3). The fluorescent images of the vinculin and F-actin co-stain can be seen...
Figure 3. Cytoskeletal organization of astrocytes and tumor cells by staining for vinculin, a focal adhesion protein, and F-actin. (A) Representative images of astrocytes and tumor cells of vinculin (red), F-actin (green), and counterstained with Hoechst (blue). Densely organized actin networks were observed in astrocyte cultures; however disorganized actin filaments were observed in U87MG and A172 tumor cell cultures. Further, vinculin was largely localized throughout the entire cell body of astrocytes, however primarily localized to the periphery of tumor cells. (B) Quantification of focal adhesions demonstrated that tumor cells resulted in significant reduction (p < 0.05) in vinculin area/cell area compared to healthy astrocytes. N = 3, mean ± SEM.
in Fig. 3A. Astrocytes exhibited aligned, densely-packed stress fiber networks distributed throughout the cell body, while actin distribution in GBM tumor cells appeared less dense. As marked differences were noted in focal adhesion formation, the percentage of positive vinculin area within a cell was quantified for the astrocytes and the GBM cell lines, U87MG, and A172. As astrocytes displayed a larger spreading area than the tumor cells, the ratio of vinculin area to cell area provides a reliable normalized measurement. As seen in Fig. 3B, astrocytes exhibited significantly larger focal adhesion areas (22.5 ± 2.52%) than the U87MG (43.6 ± 1.12%) and A172 (12.5 ± 1.66%) tumor cells (p < 0.05). A statistical difference in the percentage of positive vinculin area between the two GBM cell lines, U87MG and A172, was not observed. This reduction in focal adhesion area also provides reason as to why the GBM tumor cells had weaker cell traction forces. The primary BT145 CICs were stained for vinculin and actin; however, an observable amount of vinculin was not positively stained, which may be due to the CICs being a non-adherent tumor sub-population and vinculin not being localized to the focal adhesions.

The differing biomechanical properties between the cancer and non-cancer cells are largely due to the changes in the cytoskeleton organization. Similar to what was observed in this study, it has been shown that non-malignant breast and ovarian cells formed a dense network of parallel actin filaments distributed throughout the cell body, however, their cancerous counterparts formed fewer filaments, most of which were shorter and largely disorganized, leading to the reduction in cytoskeletal stiffness\(^4\). Focal adhesion complexes play a significant role in cell migration, as well as adhesion to the substrate. When focal adhesions attach and pull the surrounding ECM along the leading edge, traction forces are generated within the cell, moving the cell forward. In conjunction with focal adhesion formation, there must be a turnover of focal adhesion attachment at the trailing edge to allow for continuous directional cell migration\(^5\). Due to its role in migration, overexpression of vinculin has been shown to reduce cancer cell migration and invasion. Conversely, a lack of vinculin expression has been associated with the development of many cancers, and potentially plays a role in epithelial-mesenchymal transition (EMT), thus suggesting a potential reason for the reduction of vinculin in the GBM cells\(^4\).

**Analysis of Biomechanical Properties Correlate CIC Differentiation with Decreased Invasive Capabilities.** CICs, which are highly migratory and mainly responsible for tumor recurrence, are a key target for cancer therapies. As these cells are resistant to traditional chemotherapeutic strategies, differentiation therapy is being developed as an approach to treat GBM tumors to inhibit their stem-like phenotype thereby reducing tumorigenicity. By inducing differentiation in stem-like primary glioma cells, Campos \textit{et al.} showed a decrease in cellular invasion\(^6\). While the majority of other studies focus on the biochemical mechanism of CIC differentiation, we investigated the effect of differentiation on the biophysical properties of these cells.

To determine if the difference in biophysical properties between the GBM cell lines and the GBM CICs was due to differentiation, the BT145 CICs were differentiated using media without EGF or FGF, but containing 10\% FBS for 3 weeks. Following differentiation, the cytoskeletal stiffness, average traction stress, and percent of positive vinculin area were measured. The cytoskeletal stiffness of the differentiated BT145 (dBT145) tumor cells significantly increased to 1310 ± 90.0 Pa (p < 0.01), similar to that of the U87MG and A172 tumor cell lines (Fig. 1B). In addition, the cell traction forces of the dBT145s (29.2 ± 1.65 nN) were measured and the traction forces significantly increased (p < 0.01) compared to the primary BT145 tumor cells (Fig. 2). Finally, positive vinculin staining was quantified, and contrary to their non-differentiated counterparts, the dBT145 tumor cells adhered well onto the surface with observed stress fiber formation and had a similar percentage of positive vinculin area (7.93 ± 0.92\%) as the U87MG and A172 tumor cells (Fig. 3). Moreover, cell morphologies between dBT145 tumor cells and their non-differentiated counterparts were observably distinct. After differentiation of the BT145 tumor cells, the cytoskeletal stiffness, average cell traction stress, and percent of positive vinculin area significantly increased compared to their undifferentiated CIC counterparts. In addition, these properties were similar to that of the less invasive cancer cell lines, U87MG and A172, supporting that differentiation of the cancer initiating cells leads to a less invasive phenotype. Therefore, this analysis of the biomechanical properties of the tumor cells supports previous reports that state CICs have a greater invasive potential than differentiated tumor cells\(^45–47\).

**Cytoskeletal Stiffness Analysis of Cells on Aligned Nanotopography Corroborates that Stiffness is Correlative of Invasive Potential.** The measurement of cytoskeletal stiffness on a rigid 2D substrate provides important baseline characteristics of the cells; however, it is important to measure these values on a scaffold that mimics the mechanical properties of the \textit{in vivo} tumor microenvironment to obtain more biologically relevant results. Therefore, we used polycaprolactone (PCL) substrates (smooth film, randomly aligned nanofibers, and aligned nanofiber films) to investigate whether cell biomechanical properties would change. The nanofibers mimic the topographical cues provided by the white matter tracts and blood vessels. Electrospun nanofibers have been previously used as a model for glioblastoma and breast cancer migration\(^24,30,34\). By using a rotating mandrel, aligned nanofibers with an average diameter of 668 ± 98.6 nm were fabricated, which are within the range of diameters of white matter tracts observed in the human brain. While the diameters of these topographic substrates range from person to person (from 0.3 µm to 10 µm\(^43\)), on average, median and average white matter tract diameters have been below 1 µm\(^46\). Randomly aligned nanofibers had an average diameter of 621 ± 173 nm. SEM images of the different substrates can be seen in Fig. 4A. The difference in alignment can be observed between the aligned and the randomly aligned nanofibers films. By using aligned nanofibers, randomly aligned nanofibers and a smooth, non-topographic film, we were able to investigate how topography and fiber orientation affected cell behavior.

In order to determine the effect of nanotopography on cytoskeletal stiffness, AFM was used to measure the cytoskeletal stiffness of U87MG, A172, and BT145 tumor cells seeded on collagen coated TCPS, smooth film, randomly aligned nanofibers, and aligned nanofibers (Fig. 4B). For both cell lines and CICs, the cytoskeletal stiffness was significantly lower on the aligned nanofibers (p < 0.05) compared to both randomly aligned nanofibers.
and smooth film with stiffness measurements of 995 ± 74.2 Pa, 984 ± 107 Pa, and 661 ± 18.5 Pa, for the U87MG cells on the smooth film, randomly aligned nanofibers, and aligned nanofibers, respectively. Stiffness for U87MG tumor cells cultured on the aligned nanofibers decreased 2-fold compared to TCPS and about 1.5-fold compared to the other substrates. Similar trends in stiffness were also observed for A172 and BT145 tumor cells.

It has been shown previously that cells cultured on aligned nanofibers migrate further and at a faster rate than on randomly aligned nanofibers or substrates without topography, similar to tumor cell migration speeds in vivo. Combined with the data demonstrating that cytoskeletal stiffness is a biomarker for GBM invasive potential, we observe that the cells are more compliant when invasive potential is increased. The actin cytoskeleton is undergoing rapid depolymerization and polymerization during migration, which likely leads to the reduced stiffness. In addition, Roca-Cusachs et al. investigated how the elongated cell shape affects the actin cytoskeleton and reduced cytoskeletal stiffness in endothelial cells. They suggest that the spatial organization of the actin cytoskeleton of elongated cells contributes to the reduced cytoskeletal stiffness. The elongated cells lack the crosslinking between parallel actin networks that occurs in cells that are uniformly spread.

**Morphometric Analysis of Focal Adhesions Correlate Larger, Elongated Adhesions with a Migratory State.** The cytoskeletal organization of cells on glass, randomly aligned nanofibers, and aligned nanofibers was analyzed to further describe the effect of topography on the cytoskeleton. As seen in Fig. 5, cells on the aligned nanofibers formed parallel actin filaments along the direction of the fibers. In addition, clusters of focal adhesions can be observed along the poles of these cells. Cells on randomly aligned fibers observed a more spread morphology, forming extensions in different directions along the fibers. Individual vinculin adhesions were characterized by area, length, and shape factor (Fig. 5B) as previously reported by Kim et al. Focal adhesions on the aligned nanofibers were significantly larger in area and length than the adhesions on glass or randomly aligned nanofibers. Focal adhesion areas on aligned nanofibers, randomly aligned nanofibers, and glass were 5.74 ± 0.441 μm², 3.91 ± 0.268 μm², and 2.96 ± 0.209 μm² respectively. In addition, adhesions also had a significantly smaller shape factor (4π(area)/(perimeter)²) on the aligned nanofibers (0.63 ± 0.020), indicating that the adhesions were more elongated and elliptically shaped than those cultured on glass (0.76 ± 0.17) and randomly aligned nanofibers (0.81 ± 0.012).

Substrate and ECM topography has an effect on stress fiber and focal adhesion formation. Human mesenchymal stem cells cultured on 500 nm gratings of TCPS exhibited increased vinculin protein expression compared to unpatterned controls. In addition, the cells on the topographic substrates exhibited a reduction in cytoskeletal...
Cells on aligned nanofibers had adhesions that were significantly larger in area, longer, and more elliptical (p < 0.001) than on randomly aligned fibers or glass. N = 21, mean ± SEM.

Figure 5. Focal adhesion analysis on glass, randomly aligned, and aligned nanofibers exhibits aligned nanotopography promotes larger and more elongated focal adhesions. (A) Representative images of U87MG tumor cells of vinculin (red), F-actin (green), and counterstained with Hoechst (blue) on aligned and randomly aligned nanofibers. White box highlights a magnified area in the inset for the fluorescent images showing vinculin staining. Aligned nanofibers promoted a spindled morphology, while randomly aligned nanofibers exhibited cells with multiple processes along the fiber directions. (B) Quantification and analysis of vinculin adhesions showed that cells on aligned nanofibers had adhesions that were significantly larger in area, longer, and more elliptical. (go) and proliferative (grow) behavior of the tumor cells. Previous studies have shown that invading cells at the leading edge have been observed in cells on an aligned nanotopography, and may aid in enhancing the migration speed. Together, this suggests the aligned nanotopography provides the guidance cues necessary to reorganize the cytoskeleton to promote a propensity for a migratory state.

**Nanotopography and Alignment Correlates with Enhanced Migratory and Reduced Proliferative Phenotype in Tumor Cells.**

With the effect of topography on the biomechanical attributes of cells investigated, further analysis on gene expression along the migratory signaling pathways was completed using qRT-PCR. Gene expression analysis was completed for pro-migratory gene markers zinc finger protein SNAI1 (SNAI1) and Notch homolog 1 (NOTCH1), anti-migratory gene marker reversion-inducing-cysteine-rich protein with kazal motifs (RECK), and pro-proliferative gene markers cyclin-dependent kinase 20 (CDK20) and cyclin D1 (CCND1). U87MG and A172 tumor cells exhibited greater than 2-fold changes for all of the investigated genes when seeded on the aligned nanofibers compared to TCPS (Fig. 6). The fold increase on the aligned nanofibers compared to TCPS for pro-migratory SNAI1 and NOTCH1 markers was 10.12 and 3.39 for U87MG tumor cells, and 16.75 and 3.09 for A172 tumor cells, respectively. Anti-migratory RECK was downregulated 2.99 and 2.54 fold compared to TCPS for U87MG cells. Similar decreased gene expression was observed in A172 tumor cells, with 2.99 and 2.54 fold respective downregulation. Furthermore, there were statistical differences between the aligned nanofiber films and the smooth film substrate for all of the analyzed genes. When comparing the gene expression of the A172 tumor cells on aligned nanofibers and randomly aligned nanofibers for NOTCH1, a greater than 2-fold difference was observed. However, for the U87MG cells on the aligned nanofibers and randomly aligned nanofibers, a greater than 2-fold expression difference was observed for the expression of SNAI1.

Previous studies showed that culturing GBM tumor cells on aligned chitosan-PCL fibers of 200 and 400 nm significantly upregulated several migratory and EMT gene markers. Our data demonstrated a significant increase in SNAI1 and NOTCH1 expression on the aligned nanofibers. Notch signaling has been shown to promote EMT through regulation of SNAI1. Zhang et al. showed that elevation of NOTCH1 signaling in GBM cells and tumor biopsies led to increased tumor invasion and EMT markers expression. Downregulation of metastasis suppressor RECK, which was observed on the aligned nanofibers, is essential for the invasiveness in GBM cells, as overexpression of RECK in T98G glioblastoma tumor cells altered the cytoskeleton to produce fewer lamellipodia and greater stress fibers, indicating decreased mobility.

In addition, we also investigated the effect of substrate topography on genes responsible for proliferation. GBM tumors have exhibited a “go-or-grow” phenomenon in which there is a dichotomy between the migratory (go) and proliferative (grow) behavior of the tumor cells. Previous studies have shown that invading cells at the...
leading edge of the tumor have low proliferative index by inhibiting entry into the cell cycle, while cells located
in the tumor core are highly proliferative, in vitro and in vivo62–66. Using an implantable nanofiber film to direct
tumor cell migration, Jain et al. showed that cells migrating away from the tumor core were in a less proliferative
state on the aligned nanofibers compared to a smooth film control, as suggested with the reduced Ki-67 staining.
F-actin staining analysis showed that cells grown on the aligned nanofiber film exhibited uniform F-actin fila-
ments compared to more punctate F-actin staining of cells on a smooth film, suggesting that the U87MG tumor
cells were less migratory and in a suspended state on the smooth film24. Gene expression of CDK20 and CCND1,
both of which play a role in the transition from the G1/S phase in the cell cycle67,68, were significantly downregu-
lated on the aligned nanofibers, with a greater than 2-fold decrease compared to the TCPS control, further pur-
porting that substrate nanotopography induces a propensity for a migratory GBM tumor cell state

**Conclusion**

This study investigated the biomechanical properties of highly aggressive and migratory GBM tumor cells. We
demonstrate that the cytoskeletal stiffness, traction stresses, and focal adhesion formation is significantly reduced
in the highly invasive tumor cells compared to healthy astrocytes. In addition, by using an aligned nanofiber
film that mimics the white matter tracts and blood vessels, we observed a reduction in cytoskeletal stiffness.
Together, with the upregulation of migratory related genes on the aligned nanofibers, aligned actin cytoskeleton,
and increased presence of larger, elongated adhesions, these data suggest the aligned nanotopography promotes
the biophysical changes in the cells leading to an enhanced migratory state. Testing the biomechanical properties
of GBM is an effective diagnostic methodology to determine the aggressiveness of GBM tumor. This in vitro
model can further be applied to elucidate how the nanotopography of the substrates transduces into altered gene
and protein expression to induce tumor cell invasion and migration, thus providing a mechanism to inhibit the
process to form secondary tumor sites.

**Methods**

**Cell Culture.** Human GBM tumor cell lines, U87MG and A172 (ATCC, Manassas, VA), were maintained
and cultured in DMEM (Corning, Corning, NY), supplemented with 10% fetal bovine serum (FBS, Gemini Bio-
Products, West Sacramento, CA), 1% penicillin-streptomycin, 1% L-glutamine, and 1% non-essential amino
acids (Corning). U87MG cells expressing enhanced green fluorescent protein (eGFP) were generously donated
by Dr. Ravi Bellamkonda (Georgia Institute of Technology). A172 cells were transfected using GFP-Actin fusion
lentiviral particles (GenTarget Inc., San Diego, CA) and positive colonies were selected for expansion. Primary
CICs, BT145 were generously gifted by Dr. Rosalind Segal (Dana-Farber Cancer Institute). The GBM CICs were
cultured in DMEM/F12 (Corning), supplemented with B27 (Gibco, Life Technologies Grand Island, NY), 15 mM
HEPES (Alfa Aesar, Ward Hill, MA), 20 ng/mL EGF (Invitrogen, Carlsbad, CA), and 20 ng/mL FGF (Invitrogen).
Differentiated BT145 tumor cells were grown in DMEM supplemented with 10% FBS. Primary rat astrocytes were
isolated from Neonatal Sprague-Dawley pups (post-natal day 2) and maintained in Neurobasal Medium (Gibco),
supplemented with 10% FBS and 1X GlutaMAX (Gibco). Mouse neural stem cells (Cyagen) were maintained and
cultured in NeuroCult™ NSC Basal Medium (STEMCELL Technologies) supplemented with 1% penicillin-strept-
tomycin, B27, 5 μg/mL FGF, 10 μg/mL EGF, and 0.2% v/v heparin.

**Preparation of Nanotopographic Scaffolds.** Aligned and randomly aligned nanofibers were fabricated
using electrospinning as previously described69. Briefly, a 10% polycaprolactone (w/v, PCL, Sigma, St. Louis, MO)
dissolved in 1,1,1,3,3-hexafluoro-2-propanol (HFIP, Sigma) overnight. To fabricate aligned nanofibers with
a diameter of 600–800 nm, a 19 gauge blunt tipped needle was used to eject the polymer jet and was collected on a spinning mandrel (3000 RPM) 10 cm away for 20 min to create a film of 10 μm thickness. A positive voltage (5–8 kV) was applied (Model ES30P, Gamma High Voltage Research). To fabricate randomly aligned nanofibers, the flow rate was adjusted to 0.5 mL/hr, charged with 8 kV, and collected for 10 minutes onto the mandrel spinning at 15 RPM. To fabricate a smooth film substrate, 12% PCL was spread onto a glass coverslip and the HFIP was evaporated under vacuum. Smooth film and TCPS were used as non-topographic controls.

Characterization of Substrates. To characterize the substrates, SEM was performed on the aligned nanofibers, randomly aligned nanofibers, and smooth film substrates as previously described24. Briefly, samples were mounted onto stubs using carbon tape. Smooth film SEM images were taken using a FEI-TeneoLoVac SEM and aligned and randomly aligned nanofiber films were taken using a JSM-7000F SEM. Fiber diameter analysis was completed using ImageJ.

Coating of Topographical Substrates. Substrates were coated with PureCol® Bovine Collagen (100 μg/mL, Advanced BioMatrix, San Diego, CA) prior to culturing the cells to aid in cellular adhesion. The collagen was added onto of the substrates and allowed to incubate at 37 °C for 1–2 hours. The substrates were then washed 3 times with sterile phosphate buffered saline (PBS) prior to seeding the cells. For coating with laminin, substrates were pre-coated with poly-L-lysine (Sigma) for 30 minutes, before being rinsed with sterile water, and coated with 50 μg/mL laminin (50 μg/mL, Invitrogen) overnight at 37 °C.

Cytoskeletal Force Measurements. To measure the cytoskeletal stiffness measurements, tumor cells were seeded on the collagen-coated tissue culture polystyrene (TCPS) or PCL substrates at a density of 20,000 cells per sample. The tumor cells were cultured for 24 hours prior to testing. An MPF3D-Bio AFM (Asylum Research, Santa Barbara, CA) was used to measure the stiffness of cells as previously described70. Briefly, the DNP-D cantilevers were calibrated using the thermal method prior to experiment. A Nikon Ti-S microscope (MVI, Avon, MA), was used to observe and select the desired cells for measurement. The AFM cantilever was placed over the perinuclear region of the cell, and an indentation was performed with a maximum cantilever deflection of 10 nm. After maximum deflection, the z-position of the AFM scanner was oscillated with a frequency of 10 Hz, amplitude of 25 nm, and duration of 1.5 seconds. The AFM tip was removed from the cell, and the z-position of the tip and cantilever deflection was recorded. A total of 3 measurements were obtained at different locations in the perinuclear region of the cell, a minimum of 15 cells was measured for each sample, and each sample was tested in biological triplicates. The Hertz indentation model was used to calculate the stiffness of each cell using a custom MATLAB (MathWorks, Natick, MA) code.

Traction force measurements were determined by culturing the astrocytes or tumor cells on polyacrylamide gels with a thin layer of fluorescent beads embedded within them. Briefly, Fluospheres (Life Technologies) were embedded into a gel comprised of 5% or 8% acrylamide and 0.08% or 0.1% bis-acrylamide, respectively. Prior to plating, the gel was coated and crosslinked with PureCol® Bovine Collagen (100 μg/mL) overnight using Sulfos-SANPAH (Pierce). After coating, 10,000 astrocytes or tumor cells were plated onto the gels. After 24 hours, fluorescent images of the spheres and brightfield images of cell on the stressed gel were taken using an Olympus IX83 inverted microscope. Following this, media was removed and 0.25% trypsin EDTA (Corning) was added to release the cell from the substrate, returning the gel to a relaxed state. Corresponding images of the beads and gel (without the cell) were taken at the same location. A minimum of 7 cells was analyzed per gel, with each gel being repeated in triplicate. Traction forces were calculated using MATLAB and ANSYS software (Ansys Inc., Canonsburg, PA). Traction forces were normalized to respective cell area to determine average cell traction stress.

Immunocytochemistry and Analysis. To stain the astrocytes and tumor cell cultures with vinculin and F-actin, approximately 15,000 tumor cells or astrocytes were cultured on collagen-coated substrates overnight (about 24 hours) before being briefly rinsed with cytoskeleton buffer. Following this, cells were fixed with 4% paraformaldehyde for 10 minutes, permeabilized with 0.1% Triton-X-100 (Sigma) for 20 minutes, and blocked in 4% goat serum for 30 minutes. Cells were then stained for focal adhesion protein vinculin (1:500, V9131, Sigma), F-actin with phalloidin (Life Technologies), and counterstained with Hoechst 33342 (Life Technologies). A Leica SP5 Point Scanning Confocal microscope was used to image the cytoskeletal organization on the substrates. Morphometric analysis of the focal adhesions of the GBM tumor cells were conducted on the images using ImageJ software. Briefly, an ellipse outline was drawn on each individual focal adhesion. The length, area, and shape factor were then determined based on the focal adhesion’s shape. An average of each shape descriptor was calculated for each cell. A minimum of 7 cells was analyzed for each substrate sample, and 3 substrate samples were analyzed for each condition.

Analysis of Key Migratory and Proliferative Gene Markers. To analyze key migratory and proliferative gene markers, quantitative reverse transcription polymerase chain reaction (qRT-PCR) was performed. In order to collect the mRNA, 750,000 tumor cells were seeded onto each substrate. Twenty-four hours post-seeding, the media was removed and RLT Buffer (Qiagen, Venlo, Netherlands) was added to the cultures for 20 minutes to lyse cells. The solution was then collected, processed, and purified using the RNeasy kit (Qiagen). The mRNA concentration for each sample was measured at the 260/280 absorbance using NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific). The cDNA was made using the equivalent of 1 μg of RNA and iScript cDNA synthesis kit (Bio-Rad, Hercules, CA). Migratory markers, zinc finger protein SNAI1 (SNAI1), Notch homolog 1 (NOTCH1), and reversion-inducing-cysteine-rich protein with kazal motifs (RECK), and proliferative markers, cyclin-dependent kinase 20 (CDK20), and cyclin D1 (CCND1), were probed using the iTaq SYBR
considered statistically significant for a p value < 0.05. ANOVA with Tukey’s post-hoc analysis was performed to determine differences amongst substrates. Data were reported as mean ± standard error of the mean (SEM), and either a t-test or a one-way ANOVA with Tukey’s post-hoc analysis was performed to determine differences amongst substrates. Data were considered statistically significant for a p value < 0.05.

Table 1. Forward and Reverse Primers.

| Gene    | Forward Primer | Reverse Primer |
|---------|----------------|----------------|
| GAPDH   | 5′-TGTAGTGAGGTCAATGAAGG-3′ | 5′-ACATCGTCTCAGACACCTG-3′ |
| SNAI1   | 5′-GGTTCTCCGGCTGACTGC-3′ | 5′-GCTGGAGGTTAATCTGCGAATT-3′ |
| NOTCH1  | 5′-CAGGCAATCCGGAGACTATG-3′ | 5′-CAACGGTGTGGCTTTCACAG-3′ |
| RECK    | 5′-CAATAGCCATTTCACAGCAG-3′ | 5′-CCAGATTATTGCCCCAGAACA-3′ |
| CDK20   | 5′-CAGTGTCTGCTTCATCTCCG-3′ | 5′-TGTCACACATATCGATGGTCCCTC-3′ |
| CCND1   | 5′-GCCTCGGTGTGCTACTTCTC-3′ | 5′-CTGTTCCTCAGCAGACTCC-3′ |

Table 1. Forward and Reverse Primers.

Green Supermix (Bio-Rad). For analysis of each gene, and the values were normalized to the housekeeping gene GAPDH. Primer sequences are listed in Table 1.

Statistical Analysis. All statistical analysis was performed by using GraphPad Prism (GraphPad Software, San Diego, CA). Data was reported as mean ± standard error of the mean (SEM), and either a t-test or one-way ANOVA with Tukey’s post-hoc analysis was performed to determine differences amongst substrates. Data were considered statistically significant for a p value < 0.05.

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Author Contributions
A.J. conceived and designed the study. A.B. performed the experiments. G.T. helped with performing the measurements using the AFM and J.G. helped obtain and analyze the measurements using TFM. Q.W. provided guidance for the AFM and TFM experiments. The manuscript was written by A.B. and A.J. which was edited and reviewed by all authors.

Additional Information
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