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Flow induces epithelial-mesenchymal transition, cellular heterogeneity and biomarker modulation in 3D ovarian cancer nodules

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Seventy-five percent of patients with epithelial ovarian cancer present with advanced-stage disease that is extensively disseminated intraperitoneally and prognosticates the poorest outcomes. Primarily metastatic within the abdominal cavity, ovarian carcinomas initially spread to adjacent organs by direct extension and then disseminate via the transcoelomic route to distant sites. Natural fluidic streams of malignant ascites triggered by physiological factors, including gravity and negative subdiaphragmatic pressure, carry metastatic cells throughout the peritoneum. We investigated the role of fluidic forces as modulators of metastatic cancer biology in a customizable microfluidic platform using 3D ovarian cancer nodules. Changes in the morphological, genetic, and protein profiles of biomarkers associated with aggressive disease were evaluated in the 3D cultures grown under controlled and continuous laminar flow. A modulation of biomarker expression and tumor morphology consistent with increased epithelial-mesenchymal transition, a critical step in metastatic progression and an indicator of aggressive disease, is observed because of hydrodynamic forces. The increase in epithelial–mesenchymal transition is driven in part by a posttranslational up-regulation of epidermal growth factor receptor (EGFR) expression and activation, which is associated with the worst prognosis in ovarian cancer. A flow-induced, transcriptionally regulated decrease in E-cadherin protein expression and a simultaneous increase in vimentin is observed, indicating increased metastatic potential. These findings demonstrate that fluidic streams induce a motile and aggressive tumor phenotype. The microfluidic platform developed here potentially provides a flow-informed framework complementary to conventional mechanism-based therapeutic strategies, with broad applicability to other lethal malignancies.

cancer metastases are responsible for 90% of cancer-related deaths, but the biological and physical factors that determine the fate and heterogeneity of metastatic tumors remain poorly understood (1–6). Ovarian cancer is the leading cause of deaths related to gynecological malignancies, and is frequently diagnosed at an advanced stage. Initially, ovarian cancer metastasizes by direct extension to sites that are proximal to the primary tumor through a complex series of events including migration, assembly, and proliferation (7–10). Dissemination to distant sites prognosticates the poorest outcomes for ovarian cancer patients and occurs via transcoelomic, lymphatic, or hematogenous routes (7, 9, 10). Among these routes, transcoelomic metastases are the most frequent and are responsible for the highest morbidity and mortality rates, which in turn are associated with the frequent production of malignant ascites (7, 9, 10). Under normal physiologic conditions, the great majority of peritoneal fluid is resorbed by the vasculature and the lymphatics and is removed from the body (11). The buildup of ascites occurs when the rate of fluid production exceeds the rate of clearance, typically because of an underlying pathology such as cancer. Multiple factors drive the accumulation of this transudate, including the overproduction of peritoneal fluid, increased leakiness of tumor microvasculature, and obstruction of lymphatic vessels (11–15). The abnormal buildup of protein- and cell-rich ascitic fluid in the peritoneal cavity occurs frequently in ovarian cancer and is associated with a dismal quality of life and a grim median survival (7, 11). A major route of dissemination for ovarian tumors is along natural ascitic currents in the peritoneal cavity that are established by the physiologic movement of peritoneal fluid (Fig. 1A, blue arrows) (7, 16–18). Among the physiological factors that drive the formation of ascitic currents are gravity, negative subdiaphragmatic pressure, and organ mobility as well as recesses formed by key anatomical structures (7, 16–18). Detached ovarian cancer cells primarily colonize distant sites under the influence of ascitic flow; however, little is known about the impact of dynamic physical forces on the molecular and architectural heterogeneity of ovarian cancer metastases.

A key molecular marker relevant to ovarian cancer biology, particularly advanced-stage disease, is the epidermal growth factor receptor (EGFR), a cell-surface receptor that regulates proliferation, growth, and survival (19–21). In ovarian cancer, high EGFR expression is associated with increased proliferation and a more aggressive and invasive phenotype and is observed more frequently in metastases than in samples of primary tumors (22–24). The EGFR may also be an important negative prognostic indicator of disease-free survival and overall survival, depending in part on the stage of the disease and the technique used to evaluate expression of this biomarker (20, 25). EGFR activation has been shown to down-regulate E-cadherin (26–28), a transmembrane glycoprotein involved in intercellular adhesions mediated by a protein complex that includes β-catenin (29). Loss of E-cadherin is associated with a poor clinical outcome and increased metastastic potential of ovarian cancer cells mediated in part by the cell ECM receptor αvβ3 integrin, which binds to...
fibronectin (26, 28). Decreased E-cadherin expression also is a hallmark of epithelial-to-mesenchymal transition (EMT) (26–30), a critical step in metastatic progression and a prognostic indicator of aggressive and refractory disease (31–35). Along with loss of E-cadherin, EMT is characterized by a range of molecular and morphological features, including the acquisition of a motile and mesenchymal phenotype with increased spindle-like morphology as well as increased expression of vimentin, an intermediate filament that maintains cell and tissue integrity and helps confer resistance to mechanical stress (30, 32, 35–37). The cell-cycle regulator p27Kip1 modulates proliferation, motility, and apoptosis by inhibiting an array of cyclin-dependent kinases including cell-division control protein kinase 2 (CDC2) and also plays a role in cytoskeletal remodeling (38, 39).

Transcriptional and posttranslational regulation of molecular markers provides a cell with the plasticity to respond to environmental changes, particularly in response to stress. However, changes in gene expression do not always correlate with altered protein levels (40–47). Modulation of molecular pathways and architectural features relevant to ovarian cancer progression by dynamic physical forces may elucidate critical metastatic mechanisms.

Microfluidic systems provide unique opportunities to investigate the influence of physical and biological parameters on the growth of metastatic tumors (48–50). The current study integrates these observations with 3D tumor growth under sustained flow to study the effects of dynamic forces on molecular alterations critical to the metastatic process. 3D cancer models restore important architectural and stromal cues (51–65) but fail to evaluate the impact of fluid hydrodynamics on tumor adhesion and micronodule growth. A system that integrates both physical and stromal modulators of metastatic biology could provide unique insights into the events that drive cancer progression and tumor heterogeneity. The microfluidic platform described here produces 3D tumor micronodules that exhibit morphological features and express a panel of molecular markers associated with an aggressive and highly metastatic phenotype. The system reveals a flow-induced post-translational up-regulation of EGFR expression and activation, a transcriptionally regulated decrease in E-cadherin expression, increased vimentin expression, and increased EMT-like characteristics. Ovarian micronodules grown under flow also show decreased volume and viability compared with nonflow 3D cultures. The data collectively highlight the importance of investigating dynamic physical influences to reflect the complex determinants of metastatic fate that cancer cells encounter under physiological conditions.

**Results**

**Modeling Fluidic Determinants of Ovarian Cancer Dissemination and Growth.** A microfluidic platform was developed to evaluate the effects of flow (Fig. 1A) on the growth of 3D ovarian micronodules as shown in the schematic in Fig. 1B and the corresponding image in Fig. 1C. Additional details of the chip dimensions are provided in Fig. 1. Modeling fluidic determinants of ovarian cancer dissemination and growth. Tumor cell dissemination and colonization of distant sites is influenced by a complex array of factors including the physical stresses that tumor cells encounter as they interact with stromal beds. (A) Ovarian cancer disseminates predominantly via movement of intraperitoneal fluid leading to a distinctive distribution pattern of tumor nodules (orange) involving four common abdomino-pelvic sites: (1) cul-de-sac (peritoneal fold between the rectum and the posterior wall of the uterus); (2) right infracolic space (the apex formed by the termination of the small intestine of the small bowel mesentry at ileocecal junction); (3) left infracolic space (superior site of sigmoid colon); (4) Right paracolic gutter (communication between the upper and lower abdomen defined by the ascending colon and peritoneal wall). This characteristic distribution is influenced by topographical pathways of ascitic flow (blue arrows) that are established by the hydrodynamics of intraperitoneal fluid motion in the abdomino-pelvic cavity. The direction and strength of these fluidic pathways are determined by physical influences including negative subdiaphragmatic pressure, gravity, and organ mobility as well as by recesses formed by key anatomical structures: (i) cul-de-sac, (ii) termination of small intestine mesentry, (iii) sigmoid colon, (iv) faliform ligament, and (v) phrenicocelel ligament. In contrast to this flow-based dissemination, the absence of ascites leads to metastatic spread that is largely proximal to the primary tumor. (B) Schematic of a microfluidic chip used to study the effect of sustained flow on the growth and molecular features of 3D ovarian cancer nodules. (C) Photograph of a microfluidic chip used in the experiments. (D) A side view of a microfluidic chip designed to study the impact of flow on the attachment and growth of ovarian cancer cells to a stromal bed. (E) Ovarian cancer cells were cultured successfully under continuous flow for 7 d in the microfluidic chip and formed 3D micronodules as shown in a 3D rendering (Left) and an optical slice (Right) of a two-photon autofluorescence image.
Fig. S1 A and B. Tumor cells entered the channels through gas-permeable silicone tubing and flowed over stromal beds of growth-factor reduced (GFR) Matrigel. A portion of the cells was effluxed from the chip via the outlet tubing; those that adhered to the Matrigel beds (Fig. 1D, Fig. S2, and Movie S1) were cultured under continuous flow for 7 d. Fig. 1E shows a representative 3D ovarian micronodule grown under continuous flow (Fig. 1E, Left, a 3D rendering of two-photon autofluorescence high-resolution z-stacks; Fig. 1E, Right, a corresponding optical slice from the 3D rendering showing cellularity throughout the nodule).

Computational Modeling of Flow in the Microfluidic Platform. To characterize the cross-sectional profile of the stromal bed inside the microchannel (Fig. 2A), 1.02-μm yellow-green fluorescent–labeled microspheres were incorporated into cooled Matrigel, and the chips were fabricated as described in Microfluidic Chip Design and Fabrication. High-resolution z-stacks were obtained by confocal microscopy, which demonstrated uniform thickness and spreading of the Matrigel in the middle of the channels. To demonstrate that flow velocity, shear rate, and vorticity are uniform in the midchannel region, where biological characterization of the ovarian cancer cells was performed, we numerically simulated the flow conditions in microchannels based on the cross-sectional profile of the Matrigel coating from Fig. 2A. Results showed that the capillary curvatures of the Matrigel bed at the two edges of the channel did not have a significant effect on the vorticity and shear-rate profiles in the midchannel region, where shear stress reached up to 0.3 Pa at the walls. To validate temperature and gaseous (CO₂) equilibration of culture medium under constant flow, we performed computational modeling of culture medium flow in the porous tubing used in this study (Fig. 2B). Results showed that at the flow rates used the CO₂ level (Fig. 2C) and temperature (Fig. 2D) of the culture medium reached equilibrium with the cell-culture conditions in the incubator [5% (vol/vol) CO₂, 37 °C] before reaching the microchannel (Fig. S1 C and D).

Tumor Volume and Viability Are Significantly Lower in 3D Ovarian Nodules Grown Under Continuous Flow than in Equivalent Nonflow Cultures. Intrachannel tumor cell adherence densities were quantified immediately postflow in three regions of the channels: proximal to the inlet (1.6–6.5 mm), middle (11.4–14.7 mm), and proximal to the outlet (19.4–24.3 mm) (Fig. 3A and B). Malignant ascites contains a mix of tumor cells and other cell types as well as proteinaceous secretions (11–15). The concentration of ovarian cancer cells in this milieu likely changes as the disease progresses and the volume of ascites increases. For the present study (Fig. 3B), we chose cell concentrations that span three orders of magnitude (10⁶, 10⁷, and 10⁸ cells/mL) to account for a broad range of possibly relevant seeding densities (12, 66). Quantitative analysis (Fig. S3A) of intrachannel tumor cell adherence densities for the three initial cell concentrations shows a concentration-dependent increase in the number of adherent cells (Fig. 3B). A significant increase in intrachannel tumor cell adherence density is observed at an initial cell concentration of 10⁶ cells/mL compared with the other cell concentrations (P < 0.05). There was a slight trend toward decreasing tumor cell adherence density along the channel, particulary in the section distal to the inlet, but these differences were not statistically significant. A more detailed characterization of the number of adherent cells per linear mm [λ (y)] immediately postflow (Fig. 3C) shows a similar trend as a function of distance along the channel at an initial cell concentration of 10⁶ cells/mL. The number of cells adhered per linear mm across the channel [λ (y)] (Fig. 3D) indicated higher concentrations toward the channel center, possibly because of increased vorticity at the edges of the channel (Fig. 2A). After 7 d of growth under continuous flow, 3D micronodules were distributed along the full length of the channel (x axis) (Fig. 3E) and across the width of the channel (y axis) (Fig. 3F). The middle region (5–15 mm along the channel) produced the highest number of viable 3D tumors, possibly because of the stabilization of flow in this section of the channel. To evaluate the impact of flow on 3D nodule volume and viability (Fig. 3 G–I and Fig. S3B), nonflow 3D cultures were plated at densities equivalent to the intrachannel tumor cell adherence densities quantified in Fig. 3B (120, 40, and 8 cells/mm²) and were grown for 7 d (Fig. S4). Growth under continuous flow resulted in a significant decrease in mean tumor volume relative to nonflow cultures, at equivalent plating densities (Fig. 3G). At an equivalent plating density of 120 cells/mm², mean tumor volume after 7 d of growth in nonflow cultures was 5.2 × 10⁸ μm³.
Characterization of Tumor Cell Distribution in Microfluidic Channels Immediately After Flow (A-D)

Distribution of 3D Tumor Nodules in Microfluidic Channels After 7 Days of Growth (E-F) and Comparison of 3D Nodule Volume and Viability with Equivalent Non-flow Cultures (G-I)

Fig. 3. Characterization of tumor cell distribution in microfluidic channels immediately after flow (A–D), distribution of 3D tumor nodules in microfluidic channels after 7 d of growth (E and F), and comparison of 3D nodule volume and viability with equivalent nonflow cultures (G–I). (A) Intrachannel distribution of adherent ovarian cancer cells immediately after the introduction of three initial cell concentrations (10^5, 10^6, and 10^7 cells/mL) into the channels was quantified in three regions: proximal to the inlet (1.6–6.5 mm), middle (11.4–14.7 mm), and proximal to the outlet (19.4–24.3 mm). (B) A concentration-dependent increase in the intrachannel tumor cell adherence density was observed in all three regions (P < 0.05). Within each initial cell concentration, results indicated no statistically significant difference in adherent cell densities across the channel in the three regions analyzed (P > 0.05). There was a trend toward decreased cell adherence density distal to the outlet at the initial cell concentration of 10^7 cells/mL. (C) Characterization of the number of adherent cells per linear mm [i(x)] showed similar trends as a function of distance along the channel at the initial concentration of 10^6 cells/mL. (D) The number of cells adhered per linear distance across the channel [i(y)] indicated higher concentration toward the channel center. (E and F) At an initial concentration of 10^6 cells/mL, adherent ovarian cancer cells that grew into 3D micromodules under the influence of continuous flow for 7 d were distributed along the full length of the channel (x axis) and across the width of the channel (y axis). (G) Compared with nonflow 3D cultures at all equivalent plating densities (black bars), growth under continuous flow (blue striped bars) resulted in a significant decrease in mean tumor volume. ***P < 0.001, *P = 0.01 to 0.05. (H) A shift toward smaller tumors was observed in 3D nodules cultured under continuous flow (blue bar) compared with equivalent nonflow 3D cultures (white bar). (I) At all equivalent plating densities, 3D nodules grown under continuous flow (blue striped bars) had a significantly lower fractional viability than corresponding nonflow cultures (black bars). ***P < 0.001.

Flow Induces EMT and an Aggressive Phenotype in 3D Ovarian Micronodules. To characterize the molecular and biological features of 3D nodules cultured under flow and nonflow conditions, we performed on-chip immunofluorescence staining of the EGFR (Fig. 4A), on-chip mRNA isolation and quantitative real-time PCR analysis (Fig. 4C), and on-chip protein isolation and quantification of select biomarkers relevant to ovarian cancer progression and resistance (Fig. 4D). Compared with nonflow cultures, 3D ovarian cancer nodules grown under flow exhibited morphological features indicative of increased EMT and a more motile phenotype (Fig. 4B). To quantify this contrast in morphology (Fig. 4B), the fractal dimension, d_f, was calculated by box counting in sets of brightfield images of cultures grown under flow and nonflow conditions. The d_f was lower under flow (1.78 ± 0.01; n = 8 fields) than in nonflow cultures (1.87 ± 0.01; n = 6 fields) (P < 0.05). These results were consistent with the expectation that fields containing cells with pronounced linear extensions characteristic of mesenchymal morphology have a lower d_f than the nonflow fields, which contain relatively densely packed nodules that fill the 2D image space more completely. A significant down-regulation of E-cadherin and CDC2 gene expression (Fig. 4C) was induced by flow (1.0 ± 0.10 and 1.0 ± 0.14, respectively) compared with nonflow cultures (1.5 ± 0.16 and 2.0 ± 0.25, respectively) (n = 6) (P < 0.05). EGFR gene expression levels were not significantly different between nonflow (1.1 ± 0.17) and flow (1.2 ± 0.12) (n = 6), whereas a significant increase in p27Kip1 gene levels was observed under flow (5.3 ± 0.71) relative to nonflow cultures (1.0 ± 0.17) (n = 6) (P < 0.05). In the presence of flow, a significant increase in EGFR protein expression and phosphorylation was observed (Fig. 4D) (0.70 ± 0.09 and 0.86 ± 0.10, n = 6 and 4, respectively) compared with nonflow cultures (0.28 ± 0.06, and 0.51 ± 0.09, n = 6 and 4, respectively) (P < 0.05). E-cadherin protein expression decreased significantly as a result of flow (0.19 ± 0.04) compared with nonflow 3D cultures (0.45 ± 0.10) (n = 9) (P < 0.05), and vimentin expression increased significantly under flow (1.14 ± 0.19, n = 9) compared with nonflow (0.11 ± 0.04, n = 8) (P < 0.05) (Fig. 4D). A flow-induced down-regulation of β-catenin expression was observed (mean normalized protein expression under flow was 0.67 ± 0.10 compared with 1.3 ± 0.10 in nonflow cultures) (n = 10) (P < 0.05). There was no change in the protein levels of integrin α5 between flow (0.57 ± 0.10, n = 8) and nonflow (0.72 ± 0.09, n = 8) cultures.

Relative to the gene expression profiles for CDC2 and p27Kip1, there was a trend toward a reversal of the protein expression (and
activation for CDC2), as a consequence of flow. Recent studies have provided increasing evidence that changes in gene expression do not always correlate with protein levels, highlighting the complexities of analyzing genetic and molecular responses in biological systems (40–47). Posttranscriptional regulation allows a more rapid response to stress and may play a more important role than previously assumed. These observations may help explain the discrepancy between RNA and protein expression of CDC2 and p27Kip1.

Discussion

The metastatic cascade is characterized by a high attrition rate and significant modulation of molecular biology caused in part by the dynamic physical forces encountered by tumor cells as they interact with distant stromal microenvironments and develop into micro nodules (1–6). The present study describes a unique microfluidic platform that addresses the significant challenge of 3D cell growth under sustained flow to explore the impact of shear stress on tumor cell attachment and micromodule formation on a stromal matrix. Our findings indicate that fluidic streams significantly modulate the architectural and biological features of ovarian micromodule growth and induce physiologically relevant characteristics that cannot be revealed with existing systems. The present study describes a flow-induced modulation of tumor morphology, biomarker expression, and proliferative rate indicative of EMT. This transition is driven in part by a posttranslational up-regulation of EGFR expression and activation, which induces a transcriptionally regulated decrease in E-cadherin expression and a concomitant increase in vimentin protein expression with no change in the expression of integrin α5 (Fig. 5). These morphological features and molecular characteristics are associated with abnormal progression for ovarian cancer patients (19, 20, 26, 31–33, 35) and collectively suggest the induction of a highly aggressive tumor phenotype.

Colonization of distant sites by metastatic nodules is a highly inefficient process because of the numerous physiological and biological hurdles as well as the physical stresses encountered by cancer cells throughout the metastatic cascade. The great majority of cancer cells that manage to colonize distant sites either persist as small colonies or suffers attrition over time (1, 6, 67, 68). Recent elegant studies have evaluated the influence of chemo-mechanical gradients on tumor migration and the impact of flow on cell proliferation using fluidic platforms (48–50). However, these studies have not investigated the impact of flow on the growth and molecular expression profiles of biologically relevant 3D tumor nodules. Shields et al. (48) have shown that interstitial flow enhances the migratory potential of metastatic cancer cells. Subsequently, Polacheck et al. (49) demonstrated that the directionality of tumor cell migration depends on the flow strength and cell density in collagen I scaffolds. Chang et al. (50) have shown that flow-induced shear stress causes cell-cycle arrest in monolayer cultures of multiple cancer cell lines. Microfluidic systems also have been used to mimic invasation and extravasation to study cancer metastasis (69, 70). Cells were embedded in Matrigel; however, no flow was present during culture. In addition, gas equilibrium and nutrient transport were not considered during long-term culture in the absence of flow. These studies have provided valuable insights into early steps in the metastatic cascade but have limitations in terms of investigating the effect of flow on cell adhesion and 3D microtumor growth. The present findings provide an understanding of how fluidic streams significantly modulate the morphologic features, biomarker expression, and the volume and viability of 3D tumor nodules. Compared with nonflow cultures plated at equivalent densities for the same growth period, 3D tumors cultured under sustained flow showed increased EMT and expressed biomarkers associated with an aggressive and metastatic phenotype (19, 20, 26, 31–33, 35). A corresponding significant decrease in 3D tumor volume and viability was observed in 3D nodules grown under flow. These findings are consistent with the decreased proliferation that is a characteristic of EMT (30, 32, 35–37) and the high attrition rates that metastatic nodules experience as they cope with the physical and biological stresses associated with colonizing distant sites (1, 6, 67, 68). The results reported here highlight the value of the microfluidic
Fig. 5. Proposed model for hydrodynamic stress-induced modulation of ovarian cancer biology. (0) Baseline levels of EGFR activity and protein expression are maintained by a complex array of factors including recycling and degradation of the activated receptor complex. (1) Flow-induced stress causes a post-translational up-regulation of EGFR expression and activation, which may result from decreased EGFR degradation and increased receptor recycling. (2) The resultant increased EGFR signaling modulates molecular pathways (26, 27) that induce a transcriptionally regulated decrease in (3) E-cadherin protein expression. A concomitant reduction in β-catenin and an increase in vimentin protein expression are observed, indicative of EMT resulting from stress induced by hydrodynamic physical forces.
Meirelles et al. (88) to predict increased colony formation, shorter tumor-free intervals in vivo, and not only resistance but stimulation by doxorubicin. However, these cells were sensitive to Mullerian inhibiting substance (MIS), a fetal testicular protein that targets the MIS receptor type II, which is expressed by a majority of ovarian carcinomas (88). The microfluidic platform provides a potentially valuable system to test chemotherapeutic agents and targeted biologics in unseparated and stem cell-enriched tumors. This system could be leveraged to gain insights into the evolution and reversibility of the morphological and molecular changes that occur under flow and to investigate the role of flow in the selection vs. induction of particular malignant characteristics such as invasiveness and genetic instability. The effectiveness of targeted agents and chemotherapeutics can be enhanced by the rational deployment of mechanistically distinct treatment modalities such as photodynamic therapy (PDT) (89). A biophysical cytotoxic therapy that involves the light-based activation of a photosensitive molecule, PDT has demonstrated clinical promise for the treatment of peritoneal cancers, particularly ovarian carcinoma (90, 91). PDT synergizes with chemotherapeutics and biological agents (19, 60, 89, 92), and these combinations are in early clinical trials.

This study attempts to address the significant challenge of studying metastatic tumor cell growth in physiologically relevant conditions and suggests the need to develop complementary interventional strategies that account for the impact of dynamic physical forces. Microfluidic systems, such as the one described here, may enable the development of therapeutic strategies informed by locoregional differences in tumor characteristics caused in part by the influence of hydrodynamic stress. This advance will provide improved biological insights and therapeutic strategies for an array of lethal malignancies and may enable the study of a variety of physiologic and pathological conditions and drug combinations.

**Materials and Methods**

**Microfluidic Chip Design and Fabrication.** The microfluidic chip (Fig. S1 and S2) was fabricated using 3.175-mm-thick polymethyl methacrylate (PMMA) (McMaster Carr). Microfluidic chip components were cut to 24 × 40 mm dimensions (VersaLASER; Universal Laser Systems Inc.). Three channels with a width of 4 mm were cut into a 254-μm-thick medical-grade double-sided adhesive (DSA) film (ArCaRE 90485; Adhesives Research) to provide the channel height with a 3-mm space in between. The channel inlet and outlet regions were fabricated with a 127° angle to facilitate fluid entrance to and exit from the channels (93, 94). The inlet and outlet ports of microchannels were 2.2 mm in diameter and were positioned 5 mm from the edge of the chip. The channel length is defined as the distance between inlet and outlet. All components were sterilized using isopropyl alcohol followed by air drying to remove excess moisture. Pressure distribution along the tubing was determined based on the following equation: ΔP = 8μLQ/πr², where μ = 0.0006965 Pa(37 °C), L = 100 cm, Q = 0.1 μL/min, and r = 0.38 mm. Note that here ΔP is the difference in pressure between the inlet and the outlet of the tube. The outlet of the tube was set to atmospheric pressure. There was a linear pressure gradient along the tube. The axisymmetric Navier–Stokes equations were solved for modeling fluidic flow inside the gas-permeable silicone tube. To model thermal and gas equilibration, axisymmetric convection and diffusion equations were solved.

**Flow Modeling.** Flow inside the microfluidic channel with Matrigel coating (Fig. 2A) is modeled by Cartesian Navier–Stokes equations. The cells moving closer to the edges of channel were constantly exposed to a small rotational field (see vorticity in z-direction; Fig. 2A). This rotational field might force cells to move closer to channel edges, generating an additional shear of the cell monolayer. Interestingly, the larger number of cells adhering in the midsection of the microchannels can be attributed to this phenomenon.

**Characterization of Intrachannel Tumor Cell Adhesion Density Immediately Post-Flow.** The microfluidic chips were fabricated as described above. One-milliliter syringes (309628; BD) fitted with 18-G blunt needles (B18-100; SAFI Infusion Technologies Inc.) were loaded with OVCAR5 cells at three stock concentrations (10⁶, 10⁷, and 10⁸ cells/mL) in complete culture medium. The syringes were attached to gas-permeable silicone tubing (20-cm inlet, 20-cm outlet) and were loaded onto a vertically placed syringe pump (Multi-Phaser, NE-1600; New Era Pump Systems) set appropriately for the syringe being used in these experiments. The cell suspensions were pumped into the microchannels at a flow rate of 100 μL/min for 5 min. During flow, cells adhered on the channel surface; those cells that did not adhere were flushed out of the channels and were collected and counted in the outlet reservoir. The cell concentrations in the inlet and outlet reservoirs were quantified using a hemocytometer before and after flow. The adhered cells in the channels were imaged and the intrachannel distribution was quantified as described in *Microscopy and Quantitative Image Analysis*.

**Growth of 3D Tumor Nodules in Microchannels Under Continuous Flow.** All procedures used to prepare the chips and to introduce OVCAR5 cells into the microfluidic channels for 3D cultures were conducted under sterile conditions in a laminar-flow tissue-culture hood. A detailed description of microfluidic chip fabrication is provided in *Microfluidic Chip Design and Fabrication*. The gas-permeable silicone tubing for 3D culture experiments was 100 cm long from the pump to the inlet and 20 cm long at the outlet. Suspensions of OVCAR5 cells were prepared at three concentrations (10⁶, 10⁷, and 10⁸ cells/mL) in complete culture medium and were loaded into 1-mL syringes fitted with 18-G blunt needles. The cell-loaded syringes were attached to the inlet tubing and were loaded onto a vertically placed syringe pump with appropriate settings for the 1-mL BD syringe used during this phase of the experiment. The outlet tubes were inserted through holes made in the caps of 5-mL conical tubes (430291; Corning) to collect the effluent. The cell suspensions were pumped into the silicone tubing at a flow rate of 100 μL/min for 5 min. Upon completion, the tubing was clamped at the inlets and outlets, and the syringes were removed from the pump. Twenty-milliliter syringes

GFR Matrigel was supplied through 100-cm-long tubing (of which 40 cm ran from the incubator lid to the microchannel inlet). The outer and inner diameters of the gas-permeable tube were 2.4 mm and 0.76 mm, respectively. The tube inside the incubator was designed to be long enough so that gaseous and thermal equilibrations were reached before the medium entered the microchannels. The incubator was maintained at standard tissue-culture conditions (5% CO₂ and 37 °C).

In our computational model, we assumed that the medium flowing in the gas-permeable tube was at ambient temperature (25 °C) and CO₂ concentration (0.03% CO₂). Therefore, there was a gradient of CO₂ from 5% CO₂ (incubator) to 0.03% CO₂ (medium flowing inside the tube) across the gas-permeable silicon tube. Similarly, there was a thermal gradient from 37 °C (incubator) to 25 °C (medium flowing inside the tube).

Fig. 2B demonstrates the schematic for the computational model. Axisymmetric Navier–Stokes equations were solved in the fluidic domain to evaluate velocity distribution. Convection–diffusion transport equations of temperature and concentration were solved in both fluidic and permeable tube domains to assess the equilibration of CO₂ and distribution of temperature. We assumed that the diffusion coefficient of CO₂ in medium can be approximated as the diffusion coefficient of CO₂ in water, which was reported as 2 × 10⁻⁹ m²/s (96). The diffusion coefficient of CO₂ across the gas-permeable tube was evaluated by incorporating the permeability coefficient (processystemsaint-gobain.com) and pressure drop across the tube. Pressure drop along the tube was calculated following the equation based on the following equation: ΔP = 8μLQ/πr², where μ = 0.0006965 Pa(37 °C), L = 100 cm, Q = 0.1 μL/min, and r = 0.38 mm. Note that here ΔP is the difference in pressure between the inlet and the outlet of the tube. The outlet of the tube was set to atmospheric pressure. There was a linear pressure gradient along the tube. The axisymmetric Navier–Stokes equations were solved for modeling fluidic flow inside the gas-permeable silicone tube. To model thermal and gas equilibration, axisymmetric convection and diffusion equations were solved.

**Flow Modeling.** Flow inside the microfluidic channel with Matrigel coating (Fig. 2A) is modeled by Cartesian Navier–Stokes equations. The cells moving closer to the edges of channel were constantly exposed to a small rotational field (see vorticity in z-direction; Fig. 2A). This rotational field might force cells to move closer to channel edges, generating an additional shear of the cell monolayer. Interestingly, the larger number of cells adhering in the midsection of the microchannels can be attributed to this phenomenon.

**Characterization of Intrachannel Tumor Cell Adhesion Density Immediately Post-Flow.** The microfluidic chips were fabricated as described above. One-milliliter syringes (309628; BD) fitted with 18-G blunt needles (B18-100; SAFI Infusion Technologies Inc.) were loaded with OVCAR5 cells at three stock concentrations (10⁶, 10⁷, and 10⁸ cells/mL) in complete culture medium. The syringes were attached to gas-permeable silicone tubing (20-cm inlet, 20-cm outlet) and were loaded onto a vertically placed syringe pump (Multi-Phaser, NE-1600; New Era Pump Systems) set appropriately for the syringe being used in these experiments. The cell suspensions were pumped into the microchannels at a flow rate of 100 μL/min for 5 min. During flow, cells adhered on the channel surface; those cells that did not adhere were flushed out of the channels and were collected and counted in the outlet reservoir. The cell concentrations in the inlet and outlet reservoirs were quantified using a hemocytometer before and after flow. The adhered cells in the channels were imaged and the intrachannel distribution was quantified as described in *Microscopy and Quantitative Image Analysis*.

**Growth of 3D Tumor Nodules in Microchannels Under Continuous Flow.** All procedures used to prepare the chips and to introduce OVCAR5 cells into the microfluidic channels for 3D cultures were conducted under sterile conditions in a laminar-flow tissue-culture hood. A detailed description of microfluidic chip fabrication is provided in *Microfluidic Chip Design and Fabrication*. The gas-permeable silicone tubing for 3D culture experiments was 100 cm long from the pump to the inlet and 20 cm long at the outlet. Suspensions of OVCAR5 cells were prepared at three concentrations (10⁶, 10⁷, and 10⁸ cells/mL) in complete culture medium and were loaded into 1-mL syringes fitted with 18-G blunt needles. The cell-loaded syringes were attached to the inlet tubing and were loaded onto a vertically placed syringe pump with appropriate settings for the 1-mL BD syringe used during this phase of the experiment. The outlet tubes were inserted through holes made in the caps of 5-mL conical tubes (430291; Corning) to collect the effluent. The cell suspensions were pumped into the silicone tubing at a flow rate of 100 μL/min for 5 min. Upon completion, the tubing was clamped at the inlets and outlets, and the syringes were removed from the pump. Twenty-milliliter syringes
Nonflow 3D Tumor Culture. The protocols for nonflow 3D ovarian cancer cultures were based on previously published methods developed by us and others (51, 53, 55, 57, 59, 60, 75, 76). Briefly, stromal beds were prepared by pipetting 250 μL of chilled GFR Matrigel into each well of a chilled 24-well plate (662892; Greiner Bio-One). After a brief incubation at 37 °C, OVCARS cells were seeded, and the central channels were filled with 2 mL cell suspension containing 4 cells/mm² to 200 cells/mm² (Fig. 54) in 1 mL of 2% GFR Matrigel complete growth medium per well. The 3D cultures were maintained at 37 °C at 5% CO₂ for 7 d with medium changes every 3 d. Based on the range of intra-channel distribution, volume, and viability of the resultant 3D nodules were quantified as described below in Microscopy and Quantitative Image Analysis.

Volume and viability of 3D tumor nodules in nonflow 3D cultures after 7 d growth. To image nodules under nonflow conditions, cultures were incubated with calcein and ethidium bromide reagents for 40 min. Fluorescence images were obtained with the Olympus FV-1000 confocal microscope as 2 x 2 image stacks of DIC images obtained by random sampling throughout respective channels imaged after 7 d of growth in the specified conditions. The overall viability, each channel (calcein, and intercalated ethidium bromide) was binarized as described above, and a mask of nodule regions in the 2D projection was computed as the summation of binary pixel values from each channel. In cases where “live” and “dead” pixels were colocalized, a logical operator was applied to assign the value to 1. Viability was reported as the ratio of total live pixels to total nodule area. This protocol is an adaptation of a methodology previously described to assess cytotoxic response (59, 60), modified here to draw comparisons among multiple untreated arms with differing growth conditions. Using the segmentation methodology above, object size analysis was performed on viable (calcein-positive) cells, and lists of estimated volumes from segmented images were binned into logarithmically spaced bins to create size-distribution histograms as previously described (59). The sensitivity and specificity of the object-identification algorithm in fluorescence images were validated for all initial cell concentrations by manual cell counting of representative fields compared with the same fields with arrows overlaid at each computer-identified object position (Fig. S3B). Data represent averages of multiarea mosaics from 3–6 channels per cell density from three independent experiments.

Measurement of the Matrigel bed profile. To measure the profile of the Matrigel bed in microfluidic channels, 1.02 μm yellow-green fluorescent carboxylate-modified polystyrene microspheres (Polysciences Inc.) were incorporated into the cold GFR Matrigel before gelation to provide fluorescence contrast into beds (otherwise prepared as described above in Microfluidic Chip Design and Fabrication). Confocal fluorescence z-stacks were obtained along the length of the channel, merged, and rotated 90° to visualize the profile.

Quantification of morphology by fractal dimension calculation. To quantify the contrasting morphology of tumor nodules formed under flow and nonflow conditions, we calculated the fractal dimension, (which describes the scaling relation of an object of mass, m, with a radius, r, as m ~ r²) was calculated from sets of DIC images obtained by random sampling throughout respective channels imaged after 7 d of growth in the specified conditions. The applicability of this approach is suggested by previous literature in which d is used to differentiate structural and morphological properties of different tissues (97). Based on the qualitative appearance of nearly circular multicellular objects forming in the absence of flow, we anticipate d > 2, in contrast to the more fibrobast-like morphology with less complete filling of 2D space within the spatial extent of objects formed under flow, corresponding to a lower value of d < 2 between 1 and 2. Calculation of d from image data was based on an open-source box-counting script for MATLAB (mathworks.com). Measurements over six to eight fields from two separate experiments were obtained to report d between each condition. Data represent the minimum of three independently selected fields pooled from two experiments for flow and non-flow conditions.

Quantitative Real-Time PCR Analysis. The on-chip total RNA isolation was performed using the Qiagen RNeasy Mini Kit (Qiagen, Inc.). On day 7 of 3D culture under flow, the microfluidic chips were placed immediately on ice, and the inlet and outlet tubes were cut to 2 cm. The shortened inlet tube was connected to the tip of an 18-G 1-inch blunt needle (B18-10G; SAI Inc.) on a 1-mL syringe (309268; BD Biosciences). Each channel was filled with 0.5 mL of RLT lysis buffer (79216; Qiagen Inc.), and the lysate was pushed out of the channel. EDTA (1 mL containing 20 mM Tris·HCl (pH 8.4), 50 mM KCl, 200 μM of each of four dNTPs (dATP, dCTP, dGTP, dTTP), 0.4 mg/mL proteinase K, and 1 x D10 buffer). This sample was then each treatment group and PCR assays in duplicate were used. PCR was carried out with an initial 5-min denaturing step at 95 °C followed by 40 cycles of an annealing step at 58 °C for 30 s, an extension step at 72 °C.
for 20 s, and a denaturing step at 95 °C for 20 s. After completion of the PCR amplification reaction, a melting curve analysis was performed by heating the PCR products and immediately cooling them to 50 °C for 1 min. The samples were then incrementally heated to 95 °C in steps of 1 °C for 10 s per step with the fluorescence measured continuously. GAPDH was used as a reference housekeeping gene for internal control. All real-time PCR reactions were run on the iCycler IQ5 (Bio-Rad). The relative gene expression was quantified by using the ΔΔcycle threshold method. The data represent pooled samples from six channels (flow) or 12 wells (non-flow) from two independent experiments for flow and non-flow conditions, evaluated in triplicate and repeated twice.

**Western Blot Analysis.** Flow 3D samples cultured for 7 d were placed on ice, and the inlet and outlet tubes were cut to 2 cm and clipped to avoid efflux of samples and medium from the channels. A 1-mL syringe fitted with an 18-G blunt needle was used to imbue 70 μL of chilled RIPA lysis buffer (89900; Thermo Scientific) containing a mixture of phosphatase inhibitors (PST26; Sigma) and protease inhibitors (P8340; Sigma) into the channels. Any efflux was collected into Eppendorf tubes and placed on ice. The end of each tube was clipped, and the chip was incubated on ice for 30 min. An additional 30 μL of RIPA lysis buffer was used to flush the lysed cells and Matrigel beds out of channels into Eppendorf tubes at the end of the 30-min incubation period. A blunt needle was used to push air through the tubing of each channel three times to improve removal of the lysate from chip. The samples were then incubated for another 30 min on ice with mixing by pipetting every 10 min during the incubation period. The lysed samples were centrifuged at 16,000 × g at 4 °C for 15 min; then the supernatant containing the protein lysate was removed and aliquoted. Each sample was a pool of three channels.

To prepare protein lysates from 7-d nonflow cultures in 24-well plates, the samples were placed on ice. The culture medium was aspirated carefully, and 200 μL of chilled RIPA buffer containing a mixture of phosphatase and protease inhibitor was added to each well and incubated for 30 min to ensure complete lysis of the cells and Matrigel beds. The lysed samples were centrifuged at 16,000 × g at 4 °C for 15 min; then the supernatant containing the protein lysate was removed and aliquoted. Each sample was a pool of 12 wells.

The Pierce Biinchoninic acid (BCA) protein assay kit (23225; Thermo Scientific) was used to quantify the protein lysate. Protein lysates were separated on 4–15% or 10% (wt/vol) SDS/PAGE gel (Bio-Rad) at 70 V for ~4 h or 1.5 h, respectively. The proteins were transferred onto an immobilob polyvinylidene difluoride (PVDF) membrane (Bio-Rad) at 120 V for 1 h on ice in the cold room. The membranes were probed with the following primary antibodies (obtained from Cell Signaling Technology unless otherwise stated): anti–β-catenin (5982), anti-p27Kip1 (3686), anti–E-cadherin (5296), anti-cdc2 (9112), anti–phospho-cdc2 (9111), anti-integrin alpha 5 (4705), anti–EGFR (4405), and anti–phospho-EGFR Tyr1173 (4407). Rabbit polyclonal vimentin antibody (sc-5655) was obtained from Santa Cruz Biotechnology. All primary antibody dilutions were 1:1,000 except for vimentin, which was used at 1:200 (final antibody concentration was 0.2 μg/μL). Anti-rabbit IgG HRP-conjugated secondary antibody (7074; Cell Signaling Technology) was used to blot for all primary antibodies except for anti-E-cadherin, for which anti-mouse IgG HRP-linked secondary antibody was used. β-Actin rabbit HRP-conjugated antibody (5125; CST) was used for actin-loading control. All proteins were detected using a nonreducing conditions, except for anti-p27Kip1, which was performed under reducing conditions. The HRP signals of protein bands were detected using the Immuno-Star WesternC substrate ECL kit (Bio-Rad) on a Kodak image station 4000R. Protein-of-interest signal intensities were normalized to their respective β-actin loading control. Densitometry was performed using Image J analysis software (National Institutes of Health). Data represent at least two independent experiments each for flow and non-flow conditions, repeated a minimum of five times using two to four replicates of randomly selected pooled samples.

**Statistical Analysis.** The experimental results were analyzed using ANOVA with Tukey’s post hoc test for multiple comparisons and Student’s two-tailed t test for single comparisons with statistical significance threshold set at 0.05 (P < 0.05). Unless otherwise stated, mean values represent three experiments with two or three channels per experiment, and error bars represent standard error of the mean. Statistical analyses were performed with Minitab (Minitab) and GraphPad Prism (GraphPad Software).

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