Extracellular Matrix Levels Modulate Outgrowths Dynamics in Ovarian Cancer.

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ABSTRACT

Ovarian carcinoma (OC) form outgrowths that extend from the outer surface of an afflicted organ into the peritoneum. OC outgrowth formation is poorly understood because there is a limited availability of OC cell culture models to examine the behavior of cell assemblies that form outgrowths. Prompted by immuno-chemical evaluation of extracellular matrix (ECM) components, laminin γ 1 and collagens, in human tissues representing untreated and chemotherapyrecovered OC, we developed laminin and collagen-rich ECM-reconstituted cell culture models amenable to studies of cell assemblies that can form outgrowths. We demonstrate that ECM promotes outgrowth formation in fallopian tube non-ciliated epithelial cells (FNE) expressing mutant p53^{R175H} and various OC cell lines. Outgrowths were initiated by cell assemblies that had undergone outward translocation and, upon mechanical detachment, could intercalate into mesothelial cell monolayers. Electron microcopy, optical coherence tomography (OCT) and small-amplitude oscillatory shear experiments revealed that elevating ECM levels increased ECM fibrous network thickness and led to high shear elasticity of ECM environment. These physical characteristics were associated with suppression of outgrowths. Culture environment with low ECM content mimicked viscoelasticity and fibrous networks of ascites and supported cell proliferation, cell translocation and outgrowth formation. These results highlight the importance of ECM microenvironments in modulating OC growth and could provide additional explanation of why primary and recurrent ovarian tumors form outgrowths that protrude into the peritoneal cavity containing ascites.

INTRODUCTION

Ovarian carcinoma (OC) can arise from the outer mucosa of the epithelial cell surfaces covering reproductive track organs including endometrium¹, ovary² and the fallopian tubes³. Within malignant tissue, transformed epithelial cells can form cell assemblies that protrude, away from the basement membrane and sub-epithelial stroma, toward the peritoneal space⁴. These cell assemblies can detach, transit into the abdominal cavity and, through superficial attachment and intercalation into mesothelial cell layers, contribute to the spread of the peritoneal disease^{5,6}. After surgery and chemotherapy, surviving tumor cells continue to form outgrowths leading to recurrent disease characterized by the enrichment of detached tumor clusters in the peritoneal cavity often containing ascites, the inflammatory fluids rich in various mitogenic factors including ECM molecules⁷. Thus, investigating the processes associated with OC outgrowth formation would provide important information about the mechanisms of disease progression.

ECM is a molecular scaffold rich in numerous glycosylated proteins including laminin γ1⁸ and collagens⁹ that, through interaction with cell surface receptors¹⁰, provide mechanical¹¹⁻¹⁴ and biochemical^{15,16} cues to regulate tissue development^{17,18} and malignant progression^{19,20}. Recent histologic examination of OC outgrowths protruding from the fallopian tube demonstrated enrichment of laminin γ1 within tumor cells forming outgrowths²¹. Consistent with these findings, we previously reported²² that both detached human fallopian tube non-ciliated epithelial (FNE) cells, expressing various mutant p53 proteins and patient-derived ovarian tumor cell clusters deposited on their cell surfaces, fibronectin, an ECM molecule that supports tumor survival and cell-ECM-cell adhesion. Most recent studies²³ using second harmonic visualization of outgrowths arising in the fallopian tube revealed the presence of yet other ECM components, the collagens. Collagen molecules were found to form fibrillar "wavy" networks beneath basal surfaces of transformed epithelium. Taken together these data provoke questions of whether and how tumor microenvironments, such as ECM, regulate the behavior of cell assemblies that form outgrowths.

Reconstitution of cell cultures with a laminin and collagen-rich basement membrane extract, Matrigel® (MG), has become a standard approach²⁴ to study the role of ECM in cellular growth²⁵, migration²⁶ and death²⁷. Several recent studies²⁸⁻³¹ employed MG to grow OC cells to evaluate their response to treatments in the context of three-dimensional (3D) tumoroid structures that resemble malignant tissue organization more closely than do two-dimensional (2D) cell cultures. While these studies provided a static view of tumor cells with ECM reconstitution, the contribution of ECM microenvironments to the dynamic process of OC growth remains unknown.

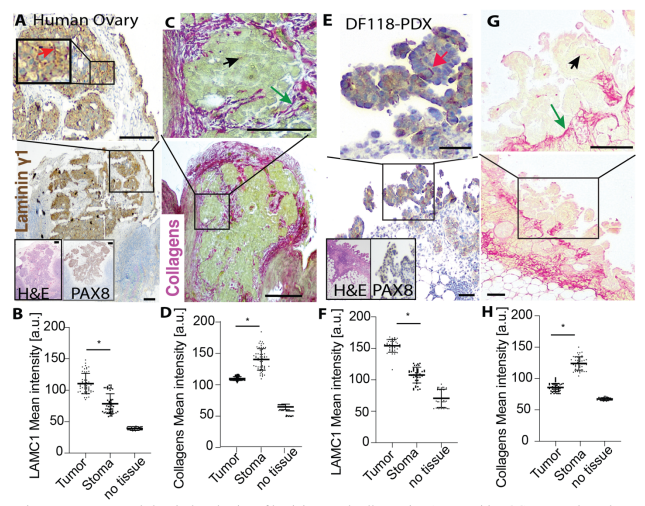
In this report we have performed immuno-chemical evaluations of laminin $\gamma 1$ and collagens in OC representing disease progression before and after chemotherapy. As in of OC tumorigenesis from the fallopian tube²¹, we found that laminin $\gamma 1$ was predominantly associated

with tumor cells, whereas collagens formed fibrillar networks within the surrounding microenvironment. These studies indicated that ECM deposition within tumor cells and adjacent connective microenvironments contributes to OC outgrowth dynamics. Motivated by these results we deployed laminin and collagen-rich (MG) ECM-reconstituted 96-well-based suspended or adhered cell culture models to study dynamics of OC cell assemblies using live-cell imaging and quantitative image processing of cell proliferation and translocation. We found that ECM reconstitution of suspended cell clusters induced outgrowths in FNE cells expressing mutant p53^{R175H}, the cells-of-origin for a proportion of OCs, and genetically distinct and well-established OC cells lines. Live-cell recordings revealed that outgrowths were initiated by cell assemblies that had undergone outward translocation and, upon mechanical detachment, could intercalate into mesothelial cell monolayers, the most frequent metastatic sites for OC. Increased ECM content altered fibrous network thickness and viscoelastic characteristics of the microenvironment, which led to the suppression of cell proliferation, directional cell translocation and outgrowths. Furthermore, ECM-reconstituted cultures that supported outgrowths, cell proliferation and cell translocation mimicked viscoelastic properties of ascitic fluids. Our results are consistent with a model whereby low-elasticity ECM microenvironments support cell proliferation and translocation of cell assemblies that form outgrowths. Our data could partially explain why primary and recurrent OCs form outgrowths that protrude into ascites, as opposed to breaching underlying collagen-dense tissues.

RESULTS

ECM deposition is associated with OC outgrowths.

Detached OC cell clusters deposit ECM on cell surfaces to support survival and cell-ECM-cell adhesion²². Pathologic examination of early tumors indicated that the sources of detached OC cell



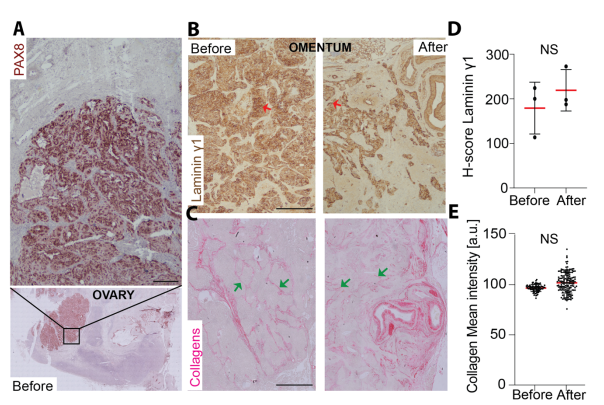
assemblies are malignant outgrowths that protrude into the peritoneal space from the outer mucosal

Figure 1. Immuno- and chemical evaluation of laminin γ 1 and collagens in PAX8 positive OC outgrowths and PAX8 negative surrounding connective tissue. (A, E) Representative images of laminin γ 1, and (C, G) collagen expression in OC outgrowths and extra-tumoral connective tissue protruding from the surface of the human ovary (A,C), or in xenograft experiments, from the surface of the omentum (E, G). Red arrows point to the deposition of laminin γ 1. Green arrows indicate fibrillar collagen structures within the tumor's connective tissue, whereas black arrows represent collagen deposition within tumors. (B, F) Quantification of laminin γ 1 and (D,H) collagen expression in PAX8 positive tumor outgrowths and PAX8 negative surrounding tissue. Each data point represents one region of interest within tumor outgrowth or surrounding stroma or area without tissue, and the line reflects the median of the data point distribution. An unpaired, two-tailed, non-parametric student t-test was used to examine statistical differences between data point distribution representing tumor and extra-tumoral stroma. * denotes p<0.05. Bars are 100 µm. No tissue values represent background level after RGB image conversion to 8-bit gray scale (0-255 pixels range)⁵³.

surface of the fallopian tube⁴. To examine whether OC cell assemblies are associated with ECM deposition, we used a combination of immunohistochemistry (IHC) for laminin γ 1 and picrosirius red stain that decorates dense fibrillar collagens (I, II and IV)³² contained within connective tissue and the basement membrane. These approaches revealed that PAX8-positive (marker of OC) cell assemblies formed outgrowths protruding from the surface of the human ovary (**Figure 1 A**), or in a patient-derived xenograft model (PDX), from the omentum (**Figure 1 E**). OC outgrowths contained laminin γ 1 (**Figure 1A, E red arrows**) and, to a much lesser degree, collagens (**Figure 1C, G, black arrows**). In the same tissue sections, we observed that adjacent extra tumoral connective space was low in laminin γ 1 and contained dense fibrillar "wavy" collagen networks (**Figure 1C, G, green arrows**). Image analysis of laminin γ 1 and collagens expression revealed significant difference in localization of these ECM molecules. Laminin γ 1 was enriched within tumors (**Figure 1B, F**), whereas collagens were mostly enriched within extra-tumoral space (**Figure 1D, H**). Taken together, these results indicate that OC outgrowths are associated with at least two ECM microenvironment factors, laminin γ 1 and collagens.

Laminin y1 and collagens are associated with OC before and after chemotherapy.

Some patients develop recurrent disease characterized by the presence of detached tumor clusters within ascitic fluids accumulating in the peritoneal cavity⁷. These clinical observations indicate the that similarly to primary tumors, tumors that have recovered from chemotherapy are associated with laminin γ 1 and collagens -rich ECM microenvironments. Thus, we wanted to evaluate ECM deposition in tumors recovered from chemotherapy. Pathologic examination of PAX8-positive primary tumors in the ovary (**Figure 2A**) and matched omental metastases representing tumors before and after neo-adjuvant chemotherapy revealed similar laminin γ 1 and collagen deposition (**Figure 2B-E**). These data indicate that OC recovery from chemotherapy is



associated with the presence of laminin $\gamma 1$ and collagens and support the

Figure 2. Metastatic OC before and after chemotherapy is associated with spatially distinct laminin $\gamma 1$ and collagen ECM microenvironments. (A) Hematoxylin and PAX8 stain of high-grade serous ovarian cancer (HGSOC) growing within the ovary before chemotherapy; bar is 100µm. (B) Laminin $\gamma 1$ (red arrows) (C) collagens (green arrows) deposition in matched omental tissue of HGSOC before and after taxane-platin based treatment; bar is 500 µm. (D) H-score based quantification of laminin $\gamma 1$ positivity in matched tissue samples representing HGSOC before and after taxane-platin based therapy. Tissues were analyzed from three patients. Wilcoxon sum rank test was used to compare the difference between the groups, NS= Not-Statistically different. (E) Quantification of collagens expression in omental metastases before and after taxane-platin based chemotherapy. Each data point represent one region of interets of RGB image converted to 8-bit gray scale (0-255 pixels range)⁵³. An unpaired, two-tailed, non-parametric student t-test was used to examine statistical differences between data points.

idea that laminin- and collagen-rich ECM can contribute to OC outgrowth dynamics before and after chemotherapy.

ECM reconstitution of suspended OC cell clusters promotes outgrowths.

Laminins (including laminin γ 1) and collagens (including collagen IV) represent scaffold components of the basement membrane^{33,34} and of the reconstituted basement membrane extract, MG³⁵. To partially mimic ECM microenvironments associated with OC, we tested whether MG supports formation of OC outgrowths in suspended cell cultures. We initially chose to examine MG effects on outgrowth formation in FNE cells expressing mutant p53^{R175H} (FNE-m-p53). We chose this cell line, because (1) fallopian tube non-ciliated cells, with mutations in the TP53 gene, are thought to be precursors of OC and an initial source of outgrowths that can disseminate into the peritoneum, and (2) we have recently demonstrated²², using this cell line, that expression of m-p53 in FNE cells induces acquisition of transformed phenotypes associated with OC progression including survival of detached cellular clusters and mesothelial intercalation²². Here, we examined outgrowth forming capabilities of FNE-m-p53 or FNE cells expressing empty plasmid. First, we stimulated cell assembly formation by clustering cells in suspension for 24 hours in single wells of low-adhesion 96-well plates (Figure 3A). Second, to mimic laminin- and collagen-rich ECM microenvironments, we reconstituted cell assemblies with media containing 2% MG (Figure 3A). We chose this ECM reconstitution condition because we have previously reported that 2% MG supported growth²⁷ of suspended breast cancer cell cultures. Live-cell microscopy revealed that, FNE-m-p53, but not FNE expressing empty plasmid significantly expanded and formed outgrowths initiating from the cell assemblies that, over time, began to extend beyond the cell cluster (Figure 3B-C, and MOVIE 1). We observed that FNE-m-p53 cell clusters formed outgrowths with an average frequency of 30%, whereas no outgrowths were observed in cell clusters composed of FNE cells expressing empty plasmid or FNE-m-p53 cultured without MG (Figure 3D). These data are consistent with the crucial role of ECM deposition in supporting

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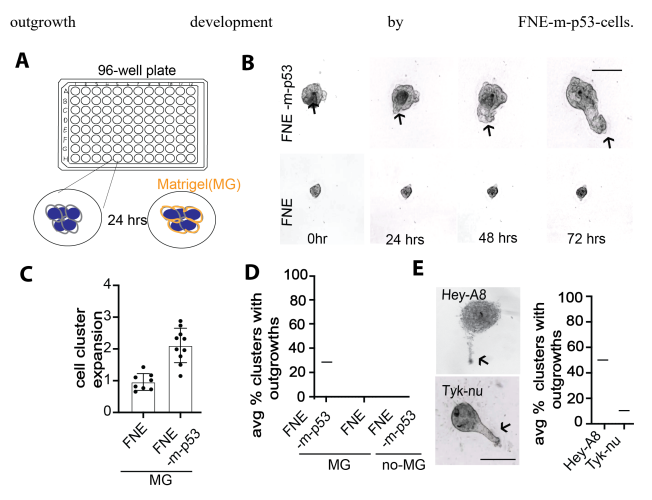


Figure 3. Laminin and collagen-rich ECM reconstitution stimulates outgrowths in suspended cultures of various OC cells. (A) Graphical representation of the assay design to study outgrowth dynamics. (B) Representative bright-field images of outgrowth formation in FNE cells expressing plasmid containing m-p53^{R175H} (FNE-m-p53), or control plasmid, and cultured as suspended clusters reconstituted with 2% MG. Cell structures were cultured for 7 days before imaging and subsequently floowed for additional 72 hrs. (C) Quantification of 3D structure expansion, each dot represents fold changer in area, over 72 hrs of filming time, in one structure. * denotes p<0.05, for an unpaired, non-parametric student t-test. (D) Quantification of cell clusters with outgrowths in FNE-m-p53 or FNE control cells. Bars represents an average percentage of outgrowth formation with at least 30 suspended cell clusters analyzed. (E) Representative bright-field images of outgrowths in Hey-A8 and Tyk-nu ovarian cancer cell lines, and quantification of outgrowths in Hey-A8 or Tyk-nu. Bars represent an average % of outgrowth formation in three independent experiments with 20-30 structures scored per experiment ; scale bars are 200 μm.

The mode of intraperitoneal dissemination is a common feature among different types of OCs such as serous and non-serous types ⁶. Therefore, we wanted to evaluate whether OC cell lines, representing other tumor types, form outgrowths with 2% MG reconstitution. We examined additional OC cell lines, Hey-A8 and Tyk-nu representing non-serous and likely serous OC models as defined by Domcke *et al.*,³⁶. As shown in **Figure 3 E**, reconstitution of Hey-A8 (**MOVIE2**) or Tyk-nu (**MOVIE3**) cellular clusters with 2% MG promoted outgrowth formation. The average percentage of outgrowth formation among Hey-A8 and Tyk-nu cell clusters varied from 50% to 10%, respectively (**Figure 3 E**). Our data suggest that ECM deposition contributes to the formation of outgrowths in some suspended cell clusters representing fallopian tube cells expressing cancer driving mutant p53 protein, serous and non-serous OC cells.

Detached outgrowths clear mesothelial monolayers.

OC outgrowths can ultimately detach and intercalate into mesothelial surfaces of abdominal organs, such as the ovary¹. So, we wanted to know whether OC outgrowths, that develop in ECM-reconstituted cultures, can intercalate into mesothelial cell layers. To do this, we examined Hey-A8 MG-reconstituted cell cultures, as these formed the longest protruding outgrowths that could be most easily dissociated from cell clusters under a dissecting microscope. We mechanically separated Hey-A8 outgrowths from cell clusters and co-cultured both entities on the surface of mesothelial cell monolayers that express green fluorescent protein (GFP) (**Figure 4A**). Of note, we observed that, upon detachment, outgrowths changed their shapes from "rod-like," to a spherical structure resembling free-floating OC spheroids commonly found in malignant ascites. To examine the outgrowths 'and cell clusters' abilities to invade mesothelial monolayers, we used a mesothelial cell monolayer³⁸. Hole formation is visualized by the time-dependent

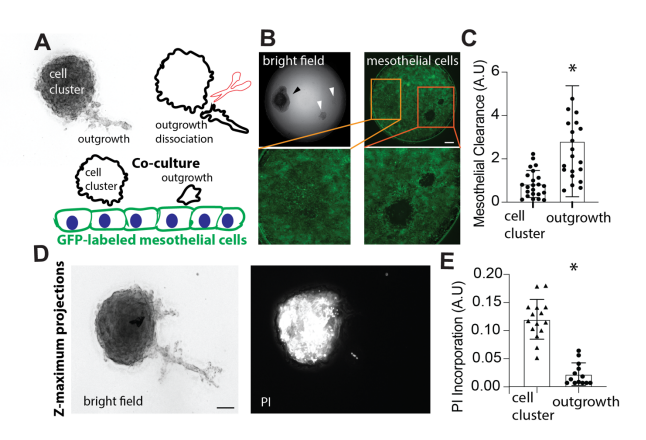


Figure 4. Detached outgrowths clear the mesothelial monolayers. (A) Schematic representation of Hey-A8 outgrowth detachment and co-culture with mesothelial cells expressing GFP. (B) Representative bright-field and corresponding fluorescent images of mesothelial clearance by Hey-A8 cell cluster (black arrow) or detached outgrowths (white arrows); bar is 200 μ m. (C) Quantification of mesothelial clearance by Hey-A8 cell clusters and detached outgrowths. Each data point represents a cell cluster or outgrowth. Twenty outgrowths and cell clusters representing two independent experiments were analyzed. * p<0.05, for an unpaired, non-parametric student t-test. (D) Representative bright-field and corresponding fluorescent maximum projection images of Hey-A8 cell clusters forming outgrowth and treated with propidium iodide (PI) to spatially visualize cell death within the structure. (E) Quantification of PI incorporation by cell clusters or outgrowths. Sixteen Hey-A8 cell structures were analyzed in two independent experiments.

exclusion of GFP fluorescence reporting on mesothelial cell migration beneath an OC cell assembly. By using this assay, we observed that detached outgrowths induced mesothelial clearance, while cell clusters, from which the outgrowths were separated, showed less clearance (**Figure 4B-C**). The observed results could be explained by the possible difference in viability

between cells contained within outgrowths and cell clusters. Spatial examination of cell viability, within cellular structures, using propidium iodide (PI), a fluorescent dye permeating only dead cells, revealed that many dead cells were localized to the cell clusters whereas outgrowths showed minimal PI incorporation (**Figure 4D-E**), which was indicative of a good cell viability. These data are consistent with the hypothesis that OC cell assemblies forming outgrowths are viable and upon detachment can intercalate into mesothelial monolayers.

Increase of ECM deposition inhibits outgrowths.

Up to this point, our data provide evidence that ECM deposition induces OC outgrowths that can intercalate into mesothelial cell layers. Next, we wanted to gain more information about the role of ECM in the regulation of OC outgrowths. Hence, we examined whether modulation of ECM content influences outgrowth formation. We compared outgrowth formation capability of suspended FNE-m-p53 and Hey-A8 cell clusters reconstituted with 2% or 25% MG. As expected, reconstitution of both cell types with 2% MG led to outgrowth formation; however, 25% MG culture conditions significantly attenuated outgrowth formation (Figure 5A-B), indicating that increasing ECM content suppresses outgrowths. We noticed that under 25% MG conditions, both FNE-m-p53 and Hey-A8 cells clusters appeared to be moderately, but significantly, smaller when compared to 2% MG reconstitution conditions (S. Figure 1A, B). Of note, as opposed to FNE-mp53 cells, Hey-A8 cells cultured in 25% MG were less compact and more disorganized (Figure 5B, inset), suggesting that elevating ECM levels may disrupt Hey-A8 cell-cell adhesion. Nevertheless, these data indicated the possibility that increasing ECM deposition affects cell proliferation. To explore this possibility, we took advantage of cell culture assay in which cell monolayers are overlayed with MG³⁹ (S. Figure 1C), partially mimicking conditions of ECM encapsulation. We integrated this assay with live-cell imaging of FNE-m-p53 and Hey-A8 cells

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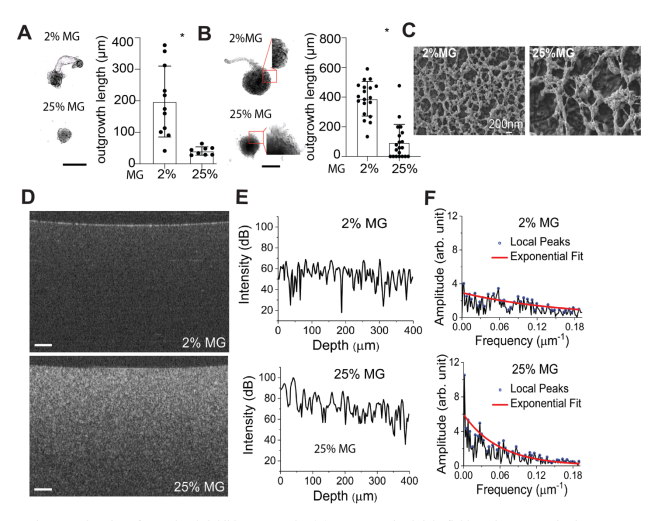
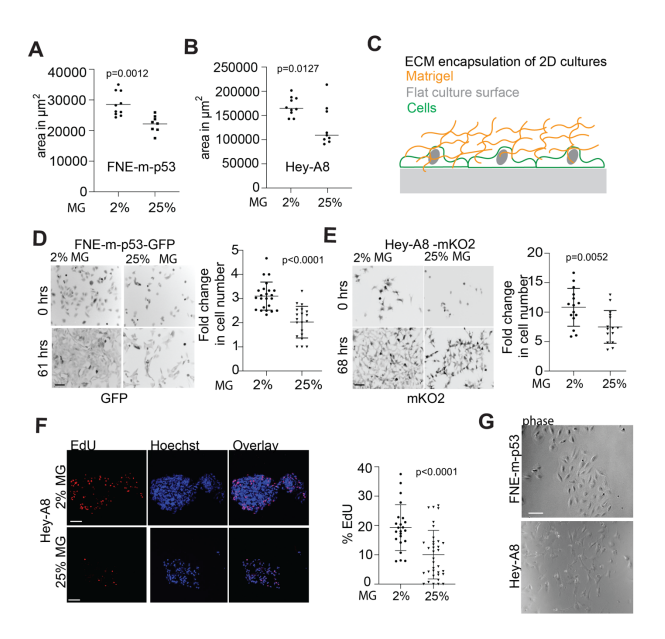


Figure 5. Elevation of ECM levels inhibits outgrowths. (A) Representative bright-field maximum Z-projection images of FNE-m-p53, bar is 250µm, or (B) Hey-8A cells cultured under suspended conditions and reconstituted with 2% or 25% MG, bar is 500 µm. Boxed area shows cell cluster structure edge. FNE-m-p53 and Hey-A8 cell clusters were grown for 9 days and the images show the final time point. Dot plots (A,B) represent distribution of outgrowth lengths from individual cell clusters reconstituted with 2% or 25% MG. * Denotes p<0.05, for Welch's t-test. (C) Representative electron scanning micrography representing 2% and 25% MG. (D) Representative OCT B-scan images of 2% and 25% MG. Scale bars are 100 µm. (E) Representative OCT depth-resolved intensity profiles for spatial frequency analysis. (F) Amplitude spectra of the spatial frequency corresponding to (E) showing relatively stronger low-frequency components from 25% MG, and exponential fit of local peaks indicating a faster decay rate in 25% MG.

expressing GFP or mKO2, respectively. As shown in **S. Figure 1D- E, MOVIE 4 and MOVIE 5** encapsulation of FNE-m-p53-GFP or Hey-A8-mKO2 cells with 25% MG significantly decreased cells proliferation. These results were also supported by the data indicating that 25% MG



S.Figure 1. Elevation of ECM suppresses FNE-m-p53 and HEY-A8 cell proliferation. Quantification of an area in (A) FNE-m-p53 or (B) Hey-8A suspended cell clusters reconstituted with 2% or 25% MG. Dots represent individual cell cluster structure measured 9 days after starting the culture. (C) Cartoon representation of the experimental design for ECM encapsulation of cell monolayers. Representative fluorescent images of (D) FNE-m-p53 or (E) Hey-A8 cells expressing RFP grown under condition of 2% or 25% MG encapsulation. Dot plots represent fold change increase in cell number over time. Each dot corresponds to one field of view. (F) Representative fluorescent images of EdU and Hoechst in suspended Hey-A8 cells and reconstituted with 2% and 25% MG. Cell structures were processed 9 days after starting the culture. Dot plot represent the fraction of EdU positive cells. Each dot corresponds to one structure. Welch's t-test was used to compute statistical difference. (G) Phase contrast images of FNE-m-p53 and Hey-A8 cells grown as 2D adherent cultures. Scale bars are 100µm.

reconstitution of suspended Hey-A8 cell cultures significantly reduced the number of cells undergoing DNA replication, as assessed by 5-ethynyl-2'-deoxyuridine (EdU) incorporation assays (**S. Figure 1 F**). We chose to examine EdU incorporation in Hey-A8 cells because these cells formed significantly larger structures when compared to FNE-m-p53 cell clusters (**S. Figure 1A-B**), making EdU incorporation analysis more robust. Our results are consistent with the idea that ECM levels modulate viscoelastic properties of OC cell environments to control OC cell proliferation and outgrowth formation.

Elevation of ECM content leads to the formation of thick fibrillar networks

Our IHC examination of tumors indicated that OCs do not invade into thick collagen fibrous networks associated with tissue that surrounds tumors (Figure 1). Increasing ECM leads to suppression of outgrowths and cell proliferation (Figure 5 A, B, S. Figure 1). These results motivated us to examine whether elevating ECM increases fibrous network thickness. To do so, we used a combination of electron microscopy and optical coherence tomography (OCT). As shown in Figure 5C scanning electron microscopy (SEM) revealed qualitative differences in ECM networks formed in 2% and 25% MG. We observed that network fibers formed in 25% MG were larger and appeared to be thicker compared to network fibers formed in 2% MG. These results suggest the possibility that elevation of MG concentration leads to the formation of larger ECM fibers. For quantitative evaluation of ECM fibers, we imaged the MG samples with OCT, where the endogenous optical contrast represents the microstructure of the sample. The OCT B-scan images revealed a higher contrast from the ECM contents with larger high-intensity clusters in the 25% MG compared with the 2% MG (Figure. 5D), suggesting a relatively larger ECM fiber size in the 25% MG. Also, the representative OCT depth-resolved intensity profiles show a generally larger spatial period of intensity fluctuations in the 25% MG (Figure. 5E), which can be clearly seen by the relatively more low-frequency components from the corresponding amplitude spectra (**Figure 5F**). This feature in the spatial frequency was quantified with the decay coefficient from an exponential fit of the local peaks in the amplitude spectra (**Figure 5F**), where the higher exponential decay coefficient in 25% MG indicates relatively stronger low-frequency components, suggesting thicker ECM fibers in 25% MG. These results led us to think that increase in ECM fiber thickness results in elevation of OC cell culture elasticity.

ECM levels modulate viscoelasticity of OC culture microenvironments.

Based on the data demonstrating larger size of ECM fiber networks formed in 25% MG cultures, we wondered whether changing concentration of ECM modulates a culture's viscoelasticity *i.e.*, time dependent elasticity and viscous behavior of the MG cell culture⁴⁰.

We characterized the linear viscoelastic material functions of cell culture media containing 2% or 25% MG. This analysis revealed, that under 25% MG culture media, the loss modulus, G", values representing viscous behavior (indicative of the energy dissipated as heat when the culture media is deformed under small-amplitude oscillatory shearing), were within a narrow range of 0.5 to 0.9 Pa in the frequency range of 0.1 to 10 s⁻¹ and thus were not sensitive to the frequency of rotation (**Figure 6A**). Under similar conditions the storage modulus, G', values representing elasticity, *i.e.*, indicative of the energy stored as elastic energy during the deformation of the culture media, were within the range of 3.5 to 4.5 Pa. The storage modulus, G', data were thus also not sensitive to the frequency of rotation (**Figure 6A**).

The loss modulus, G", values of samples with 2% MG ranged between 0.05 to 0.1 Pa while the storage modulus, G', values of these 2% MG samples were between 0.09 to 0.3 Pa (**Figure 6B**). The storage moduli of 25% MG reconstitution culture media were significantly higher than the loss moduli (G' > G"), and the ratio of G''/G' (*tan* δ) was below 1 (**Figure 6C**). On the other

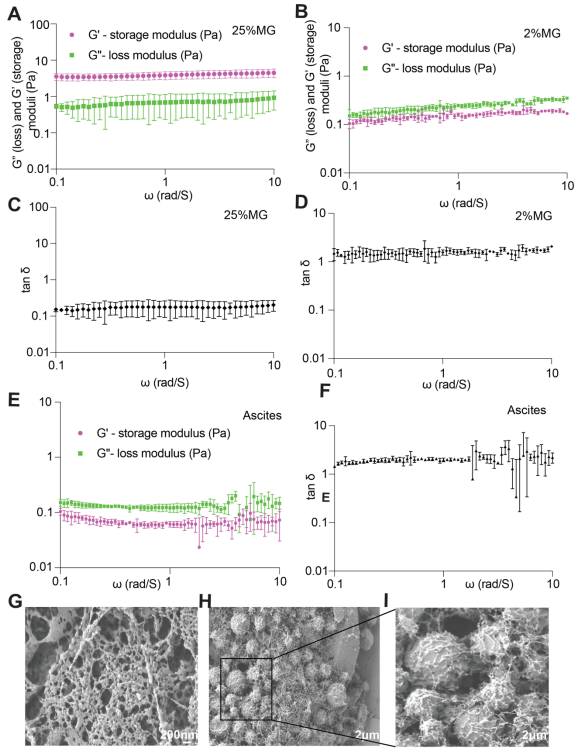


Figure 6. ECM levels modulate viscoelastic properties in the OC culture microenvironment. Measurements of storage (elastic) and loss (viscous) moduli in OC cell culture media reconstituted with 2% or 25% of MG (A, B); or in ascitic fluids isolated from OC patients with relapsed disease (E). (C, D, F) Graphs represent calculated tan δ from values reported in A, B and E. (G, H,I) Scanning electron micrographs representing ascites fibrous ECM network surrounding tumor cells.

hand, 2% MG reconstitution cultures produced tan δ values greater than 1 (Figure 6D) indicating less elastic nature of the 2% versus the 25% MG. These results suggest that elasticity of the 25% MG reconstitution media is significantly higher than that of 2%. Interestingly, measurement of loss and storage moduli of peritoneal ascites isolated from a patient with ovarian cancer, revealed close similarity to G" and G' values and tan δ observed in the 2 % MG cultures (Figure 6E-F), indicating that ascites represent gels with low strength values. These results motivated us to examine whether ascites form extracellular fibrous networks. Scanning electron microscopy evaluation of ascitic fluids revealed the presence of extracellular fibers that were also associated with cells (Figure 6G-I). These results are consistent with the hypothesis that ECM-rich microenvironments, with low shear viscosity and elasticity, such as 2% MG or ascitic fluids, support outgrowth formation.

Increasing ECM deposition suppresses directional translocation.

Based on the data demonstrating that outgrowths were initiated by cell assemblies that translocated outside the cellular structure (**Figure 3**, **MOVIES 1-3**) and that increasing ECM concentration suppressed outgrowths, decreased cell cluster size, and cell proliferation (**Figure 5 and S. Figure 1**), we hypothesized that high ECM levels also suppress cell translocation. To track cells encapsulated by ECM, we overlayed FNE-m-p53-GFP and Hey-A8-mKO2 cells grown on 2D surfaces with 2% or 25% MG (**S. Figure 1C**). Under this condition FNE-m-p53-GFP cells-maintained cell-cell adhesion (**S. Figure 1G**), forming well-defined and coherent colonies. However, Hey-A8 cells remained as single cells with little cell-cell adhesion (**S. Figure 1G**). We analyzed cell translocation of FNE-m-p53-GFP and Hey-A8 cells overlaid with different MG concentrations. To quantify the dynamics of cell translocation over time, we integrated TrackMate

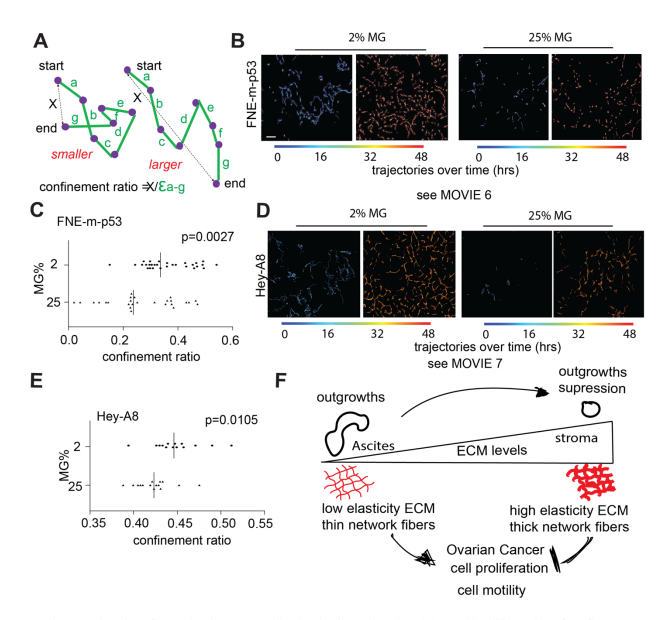


Figure 7. Elevation of ECM levels supresses diretional cell translocation. (A) Graphical illustration of confinement ratio quantification. (B) Representative trajectory images of cell translocation within FNE-m-p53 or (D) Hey-A8 cell monolayers overlayed with 2% or 25% MG. Each line represents single cell trajectory formed during 50 frames/time interval. Time zero correspond to first 50 frames. Distribution of confinment ratio within (C) FNE-m-p53 or (E) Hey-A8 cell monolayers overlayed with various concnetration of MG. Each dot represents one field of view that contained 20-100 cells. Two independent experiments were performed with a total of 12-30 fields of view. p value was derived using unpaired two-tailed t-test. Bar 100µm (E) Graphical representation of key findings in this study.

with Trajectory classifier (both ImageJ plug-ins) to compute confinement ratios, which report on how efficient is track displacement (cell translocation) away from its starting point (**Figure 7A**).

Larger confinement ratio indicates more efficient cell directionality (Figure 7A). We observed that overlaying of FNE-m-p53 or Hey-A8 cells with 2% MG resulted in confinement ratio that was significantly larger than confinement ratio of cells overlaid with 25% MG (Figure 7B-E). We also observed the presence of more sustained (directional) tracks that extended, over time, in cells overlaid with 2% MG as compared to 25% MG culture (Figure 7B, D and MOVIES 6, 7). These results indicated that increase in ECM deposition suppresses directional cell translocation. Our results are consistent with the idea that elevation of ECM levels increases elasticity of OC microenvironments and restricts outgrowth development through the suppression of cell proliferation and directional cell translocation.

DISCUSSION

In tissue samples representing human OC, IHC of laminin $\gamma 1$ and picrosirius based detection of fibrillar- and basement membrane associated collagens revealed that OC outgrowths were associated with two distinct ECM microenvironments. The first ECM microenvironment was observed among carcinoma cells and contained laminin $\gamma 1$, with low levels of collagens. The second showed low laminin $\gamma 1$ extra-tumoral stroma that contained dense collagen fibers. These results prompted us to use a laminin- and collagen-rich reconstituted tissue culture approach to study spatial and temporal evolution of OC outgrowths.

Laminin- and collagen-rich MG³⁵ reconstitution of OC cell cultures was used in combination with live-cell imaging, SEM, OCT and rheology to examine the contribution of ECM to OC cell-assembly dynamics. Live-cell imaging studies demonstrated that ECM reconstitution of suspended cell cultures, representing various OC types, promoted formation of outgrowths that, upon detachment, intercalated into mesothelial cell layers. We observed outgrowth formation in multiple genetically distinct OC cell lines including FNE-m-p53. ECM reconstitution did not promote outgrowths in "normal" non-transformed FNE cultures, indicating that outgrowth formation is associated with transformed phenotypes of the cells-of-origin. OC outgrowths were initiated by cell assemblies that had undergone outward translocation. Increasing ECM deposition led to increased elasticity of OC cell culture media, suppressed proliferation, decreased directional cell translocation and suppressed outgrowths. Based on these results we propose a model in which OC cells favor low elasticity ECM microenvironment to divide, translocate and form outgrowths that extend into the peritoneal cavity (**Figure 7F**). Our data potentially explain why some genetically distinct OC types are superficially invasive and prefer to grow away from ECM dense stroma and form outgrowth protrusions extending into the low elasticity environment of the peritoneal ascites.

Recent IHC evaluation of OC within the fallopian tube revealed enrichment of laminin $\gamma 1$ chain ²¹, a component of the heterotrimeric laminin $\alpha\beta\gamma$ molecule⁴¹. Consistent with these studies, we found that cancer cells representing human OC outgrowths protruding from the ovary and omental metastases were enriched in laminin $\gamma 1$. In contrast to tumor cells, the adjacent stroma displayed low levels of laminin $\gamma 1$ deposition, highlighting a possible involvement of $\gamma 1$ -chain containing laminins in the development of cell-ECM-cell adhesion among tumor cells forming outgrowths. Laminins promote formation of basement membranes that support epithelial cell proliferation and collective cell movements during tissue development ^{42,43}. We speculate that formation of a sufficient ECM adhesion among OC cells could contribute to tumor expansion and cell translocation leading to initiation of malignant outgrowths. To support this speculation, we provide evidence that reconstitution of suspended OC cell clusters with ECM promoted proliferation, directional cell translocation and outgrowths.

We observed a dense and thick collagen fiber network enriched within extra-tumoral spaces. These results were consistent with previous studies demonstrating collagen-fiber localization beneath OC outgrowths originating from the fallopian tube²³ and stroma surrounding metastatic omental implants⁴⁴. Collagen fiber formation occurs when collagen concentration increases⁴⁵, indicating that extra-tumoral fibrils reflect the presence of highly elastic ECM. Metastatic OC superficially invades mesothelial layers of peritoneal organs⁶, and consistently we observed that OC tumors injected into mouse peritoneum colonized surfaces of the omentum and formed superficial outgrowths extending into the peritoneal cavity, with minimal invasion of underlying stroma. Based on these results, we speculated that dynamic evolution of OC outgrowths is supported by low ECM content microenvironments. Thereby, we evaluated the dynamics of cell translocation and outgrowth formation under low- (2%) and high- (25%) MG content conditions. We demonstrated that cell translocation, cell proliferation and outgrowth formation were more robust under 2% MG culture conditions, indicating a possibility that high content of ECM surrounding OC cell clusters provides a physical barrier that suppresses OC progression. To directly test this possibility, we overlayed cell monolayer cultures with media containing 2% or 25% MG and observed that 25% MG significantly suppressed proliferation and restricted directional cell translocation. The correlations between cell proliferation, cell translocation and outgrowth formation are consistent with the ability of recurrent disease to form new metastases and support the idea that OC growth drives peritoneal dissemination.

Recent studies demonstrate that increasing concentration of collagen restricted single cell translocation from the cell clusters, as measured by mean square displacement of the tracked trajectories⁴⁶, and targeting fibroblasts that make collagens results in reduced collagen deposition and increased OC cluster dispersal ⁴⁴. Furthermore, 2D studies of the effects of ECM concentration

on cell motility have shown optimal ECM concentrations that supported directional cell migration. Concentrations below and above these optimal values did not stimulate fibroblast migration to the same level⁴⁷. Our studies support this relationship because OC cells that were reconstituted in a high concentration of ECM showed reduction in directional cell translocation and outgrowth formation. This could be explained by optimal integrin-complex turnover rates under appropriate (likely low) ECM concentrations. It is well established that high concentrations of ECM lead to strong adhesion⁴⁷ and low adhesion turnover⁴⁸. Therefore, we speculate that microenvironments with low ECM content support formation of adhesion complexes that are highly dynamic and, thus, promote directional cell translocation. In contrast, attachment of OC cells to extra-tumoral and dense ECM microenvironments would lead to strong adhesion that could impact directional translocation. This speculation is supported by observation demonstrating larger size of ECM fibrous networks and decreased directional cell translocation under high-ECM reconstitution conditions. This could explain why some OC outgrowths rarely invade collagenous stroma.

Varying ECM content modulates viscoelastic characteristics of the microenvironment and could affect cell behavior⁴⁹, including acquisition of transformed phenotypes⁵⁰. Using electron microscopy, OCT and rheometric analysis of ECM-reconstituted cultures, we provide evidence that 2% MG reconstitution results in the formation of thin (as compared to 25% MG cultures) extracellular fibrous networks and low-elastic properties of cultures, whereas 25% ECM reconstitution led to the development of thick fibrous networks and elastic properties in cell culture media, a characteristic of a more solid environment. Our data demonstrated that ECM concentrations that maintain low elasticity of tissue culture support outgrowths, through cell proliferation and directional cell translocation. Thus, it is conceivable to propose that ECM deposited on the surface of OC tumor cells, creates a low-elasticity microenvironment that provides

enough adhesion and traction force to support, cell division and cancer cell directional translocation. Increasing deposition of ECM within tumor microenvironments would increase adhesiveness, and suppress cell proliferation, cell translocation and outgrowth formation (**Figure 7F**). Our model is consistent with the clinical observations that OC peritoneal dissemination is associated with tumor outgrowths that protrude into the peritoneal cavity^{4,21} where ascites, representing low-elasticity microenvironments (as opposed to solid collagen-rich tissues), are enriched with ECM molecules^{7,51}. To support our model, we provide evidence that shear moduli values of 2% MG cultures resembled ascites enriched with fibrous extracellular networks overlaying carcinoma cells.

OC outgrowths continue to evolve after chemotherapy because detached tumor cell clusters are present within the peritoneal cavity fluids of patients with recurrent disease⁷. We demonstrate that OC cells that have recovered from chemotherapy are in direct contact with ECM deposited within tumors and extra-tumoral space. Thus, the described ECM-reconstitution model of OC outgrowths offers a clinically relevant approach to examine evolution of recurrent phenotypes, such as outgrowths, associated with the disease. The implementation of live-cell imaging, SEM, OCT, and rheology provided important insights into the role of physical characteristics of OC cell microenvironments and their contribution to OC outgrowth dynamics.

METHODS

Cell Culture

Hey-A8 (obtained from Dr. Sumegha Mitra's laboratory, University of Indiana) and Tyk-nu (obtained from Dr.Joan Brugge's laboratory, Harvard Medical School) cells were cultured in a 1:1 ratio of Medium 199 (Gibco) and MCDB105 (Sigma) supplemented with 5% fetal bovine serum (Gibco), 1% Penicillin-Streptomycin (Gibco) and 50 μg/ml plasmocin prophylactic (InvivoGen). FNE cells (obtained from Dr.Tan Ince's laboratory, Weill Cornell Medicine, New York) were

cultured in 1:1 ratio of Media 199 (HiMedia) and DMEM/F12 (HiMedia), 3% HI-FBS (Corning), 1% Penicillin-Streptomycin (Sigma), 0.5 ng/ml of 17 beta-estradiol (US Biologicals), 0.2 pg/ml of triiodothyronine (Sigma), 0.025 µg/ml all-trans retinoic acid (BeanTown Chemicals), 20 mg/ml of insulin (Sigma), 0.5 ng/ml of EGF (Peprotech), 0.5 µg/ml hydrocortisone (Sigma), 25 ng/ml of Cholera Toxin (Calbiochem, USA) and 50µg/ml plasmocin prophylactic (InvivoGen). Cell cultures were tested for the presence of mycoplasma every 3-6 months using the Uphoff and Drexler detection method⁵².

ECM Reconstitution

100 cells per well were seeded on ultra-low attachment 96-well plates (Corning). Plates were spun at 900 RPM for 3 minutes and incubated until the following day. The next day, on ice and using prechilled pipette tips, 100 μ l of 4% or 50% Matrigel® (MG) (Becton-Dickinson) was added to each well containing 100 μ l of media and cell clusters (final culture volume was ~200 μ L).

Processing of Cell Clusters for Immunohistochemistry (IHC).

Cell clusters were fixed in 4% paraformaldehyde solution (Sigma) then dehydrated in ethanol. Histogel (Thermo Scientific) was liquefied, and cell clusters were mixed with Histogel and left to solidify in a biopsy cassette. Histogel blocks were fixed in 10% neutral buffered formalin overnight then dehydrated in ethanol and cleared in xylene. Processed Histogel blocks were then embedded in paraffin and stored at -20°C until the time of sectioning. Paraffin blocks were sectioned at a thickness of 5 µm using a microtome (Leica).

IHC

Formalin-Fixed Paraffin-Embedded (FFPE) patient-derived xenograft tissue (DF118-PDX) is a kind gift from Dr. Ronny Drapkin, University of Pennsylvania. Sections were deparaffinized in two xylene changes for 10 minutes each, then sections were de-hydrated in a graded series of ethanol

(100%, 90%, 70%, 50%) 5 minutes each, and washed in ultrapure water for 5 minutes. The ABC kit; Vectastain Elite (Vector Laboratories) was used, per the manufacturer's recommendation. The primary antibodies used were anti-PAX8 (1:1000; #10336-1-AP; Protein tech.), anti-LAMC1 (1:500; #HPA 001909; Sigma Aldrich). Peroxidase substrate (ImmPACT DAB; #SK-4105; vector laboratories) was used to develop and visualize staining under the microscope (Nikon ECLIPSE Ci-L using Nikon DS-Fi3 camera). The slides were rinsed in water, dehydrated in a series of ethanol solutions of increasing concentrations, until 100% ethanol was reached, then cleared in xylene, mounted with a non-aqueous mounting medium (#H-5000; Vector Laboratories), sealed using clear nail polish and left to dry before imaging. Collagen staining was performed using a Picrosirius red staining kit (#24901; Polysciences, Inc.) per the manufacturer's recommendation. Deparaffinization and hydration were followed by 1-hour staining in Picrosirius Red, washing in hydrochloric acid, dehydration, and mounting as described above.

IHC Image Acquisition

Stained slides were scanned using an Olympus IX83 microscope equipped with a DP80 color camera and CellSens software. Based on scanned slides, ROIs were identified and recaptured using a Nikon Eclipse equipped with DS-Fi3 color camera and NIS Elements D software. Images were saved as TIFFs for further processing.

IHC Image Processing

As previously described⁵³, RGB images were converted, using a weighted RGB conversion option in FIJI⁵⁴ (Fiji is just ImageJ). software, to an 8-bit grayscale range representing values between white (0) and black (255). To make backgrounds white, images were inverted and multiple ROIs representing tumor, stroma or areas without tissue were selected. Mean gray values (values between 0-255) were calculated for each ROI and plotted.

Live-Cell Imaging of Cell Clusters

Ultra-low adhesion 96-well plates containing ECM reconstituted cell cultures were placed within a BioTek LionHeart FX long-term imaging chamber equipped with enclosed optics, temperature and gas exchange controls. Individual cell clusters were imaged for up to 7 days with indicated time intervals. Multiple XYZ planes of cell clusters were acquired simultaneously.

Quantification of Outgrowth Protrusion Frequency

To quantify the frequency of outgrowth protrusions by cell clusters representing different OC cell lines, different laboratory members set up at least 30 suspended cell cluster cultures reconstituted with 2% MG. Outgrowths were defined as distinct cell populations that appeared outside of the main cell cluster. Scoring was done using a Nikon 2000 tissue culture microscope with a 10X objective.

Pathology Scoring of Laminin γ1 Expression in Human Tumors

Intensity profiles of IHC were qualitatively scored as strong (+3), moderate (+2), weak (+1) and negative (0). IHC results were recorded by a pathologist using H-scores. $H^{score} = (\%cells + 3 \times 3) + (\%cells + 2 \times 2) + (\%cells + 1 \times 1) + (\%cells 0 \times 0)$. Data are represented as dot plots of H-scores for matched biopsy. Since the data falls into a category of a not-normal distribution, a (nonparametric) Wilcoxon rank-sum test was used to calculate significant differences between pre- and post-therapy specimens.

Ascites and solid carcinoma tissue collection

Ascitic fluids were collected from patients with ovarian cancer either by paracentesis under local anesthesia or at the very beginning of a surgery. Ascites fluids were centrifuged at $1,100 \times g$ for 10 min at room temperature to remove cell clusters, aliquoted and stored in -80°C until analysis. All solid tissue collections were performed during surgery. All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and

national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Patients provided a signed informed consent, approved by the Ethics Review Board of Poznań University of Medical Sciences (Consent No 737/17).

Mesothelial clearance assay

Mesothelial cells were plated on glass-bottom dishes (Mat-TEK Corporation), which had been coated with 5µg/ml of fibronectin (Sigma, USA) and/or collagen I (Sigma, USA). Cells were maintained in culture until confluent (48 hours after plating). In co-culture experiments, cell clusters and outgrowths were added to a confluent mesothelial monolayer expressing green fluorescent protein (GFP), allowed to attach, and imaged for up to 120 hours using a LionHeart FX (BioTek) inverted Motorized Widefield Fluorescence Microscope. Only cellular clusters or outgrowths that remained attached during the experiment were used for quantification. Mesothelial clearance was quantified as previously described ⁴⁶.

Determination of Outgrowth Length and Cellular Structure Area

GEN5 image analysis software (BioTek) was used to quantify outgrowth length and ImageJ to analyze cellular cluster size. Before quantification, a series of bright field images, representing multiple XYZ planes of a discreate cellular structure were collapsed to generate maximal projections of a cell cluster and protruding outgrowth. Outgrowth protrusion was identified as a structure that extended beyond the cell cluster, and a line was drawn from the tip across the longer axis of the outgrowth, terminating at the junction between outgrowth protrusion and cell cluster. The size of the cellular structure was calculated using an in-house ImageJ macro. Briefly, binary masks were created to separate a single cellular structure from the non-cellular background.

EdU incorporation assay

MG reconstituted Hey-A8 cells were incubated with 10 mM Click-iTÒ EdU [Thermo Fisher Scientific, catalog number C10340] for 4 hours. Cells were fixed with 4% PFA for 1 hour at room temperature, embedded into HistoGel and sectioned. Sections were permeabilized with 1X PBS containing 0.5% Triton-X. Residual Triton-X was removed by washing twice with 1X PBS supplemented with 3% BSA. To label the incorporated EdU, sections were incubated in the dark for 30 minutes with Click-iT reaction cocktail prepared fresh (<15 minutes before labeling). Click-iT reaction cocktail was then removed, and sections were washed with 1X PBS. To label total cell nuclei, sections were incubated in the dark with 10 µg/mL Hoechst 33342 in 1X PBS for 30 minutes. After washing out the Hoechst dye with 1X PBS, sections were mounted with aqueous mounting media and sealed by coverslips. Sections were imaged on ZEISS LSM -880 confocal microscope using a 10X objective. Images of EdU-labeled nuclei were acquired using far-red laser illumination ($\lambda = 647$ nm) and total nuclei were captured by violet blue laser illumination ($\lambda = 405$ nm). Images were analyzed using open-sourced software FIJI⁵⁴.

Cell proliferation, cell motion imaging and quantifications

Monolayers of FNE-m-p53 or Hey-A8 cells expressing GFP or mKO2, respectively were overlayed with MG. Time-lapse imaging was performed on a BioTek Lionheart FX automated microscope using a 10X objective and maximal interval of 22 minutes. After acquisition, time-lapse images were background subtracted (radius 50 pixel), using FIJI⁵⁴ image analysis software. To analyze cell proliferation and movement within MG overlayed cell cultures, we used an ImageJ plugin, TrackMate⁵⁵, to perform single-particle tracking (SPT). TrackMate offers multiple modes of analysis and based on our cell size and the relatively slow movement of cells, we selected and applied the following parameters in TrackMate: LoG (Laplacian of Gaussian) detector, spot diameter of 35 pixels, threshold 10, median filter and LAP tracker with 15-pixel frame linking, 2-frame gap distance

and track-segment splitting. We extracted the "spot" count in the desired time frames (one count every 12 hours or 33 frames) to obtain the number of cells or "spots" in the region of interest at those particular time points. We then divided the number of cells in each time point by the number of cells at the starting time point (time = 0) to normalize and plot it as fold increase. A non-parametric t-test using Welch's correction was used to assess significance in fold increase in cell number between the first and the last time point. Plotting and statistical analysis was computed using GraphPad Prism [version 9.1.0 for Windows, GraphPad Software, San Diego, California USA, www.graphpad.com]. To quantify track displacement, we calibrated the images by converting output in pixels into microns, and frames into hours. For visualization of local tracks, we used 50-frame-depth-, timecolor-coded tracks overlayed on a GFP or RFP-channel time lapse. For trajectory classification we used the TraJClassifier plugin⁵⁶. The plugin uses TrackMate output data and trajectories and computes confinement ratio representing cell directionality over time. After all the trajectories were computed into confinement ration, we plotted the data using Prism GraphPad Prism 9.1.0. Trajectories from multiple fields of view were plotted and directed-motion trajectories were represented as confinement ratio values. Each dot in a dot plot represents one field of view with the horizontal line depicting the mean of all fields of view per condition. Statistical analysis was computed using unpaired two-tailed t-test.

Cryo-SEM Sample Preparation and Imaging

All samples for cryo-SEM were prepared by using a Leica EM HPM100 high pressure freezing (HPF) system. The HPF planchettes were washed in ethanol and exposed to an oxygen plasma for ten minutes prior to use. The frozen hydrated samples were stored in liquid nitrogen. A Leica VCT-100 system was used for subsequent cryotransfer and cryo-imaging. Samples were transferred and coated with sputtered Au (2.5 nm) under cryogenic conditions (T < -135 °C) by using a Leica EM

MED020 system. Prior to SEM imaging, sublimation was used to create topographic contrast by slightly warming frozen-hydrated samples. SEM imaging was done by using a Zeiss Auriga Cross-Beam FIB-SEM equipped with a Schottky field-emission electron gun (FEG) and an Oxford Max-80 ultrathin window (UTW) silicon-drift detector (SDD) interfaced to an Oxford INCA EDS system. Secondary electron imaging (Everhart-Thornley detector) was done by using 2 keV electrons and a 2.5 nm Au coating.

Assessment of ECM network in MG with OCT

A spectral domain OCT system with a central wavelength of ~850 mm was used for MG imaging. The system provides an axial resolution of ~9 μ m in biological samples (1.4 refractive index assumed) and a transverse resolution of ~5 μ m. Parameters for imaging of 2% and 25% MG samples were kept the same. Data processing focused on characterization of the spatial frequency over depth, similar to a previously developed method⁵⁷. Briefly, with the dB intensity A-scan and the identified sample surface, the intensity profile over 0.4 mm starting at ~26 μ m below the sample surface was utilized for analysis. Through a fast Fourier transform, the amplitude spectrum of the spatial frequency was obtained, and an exponential fit of the local peaks was performed. The exponential decay coefficient was extracted as the measure of the spatial frequency over depth and was used for comparison between the 2% and 25% MG samples. A higher decay coefficient represents relatively stronger low-frequency components of the spatial frequency spectrum.

Rotational Rheology

The linear viscoelastic material functions, the storage modulus, G', and the loss modulus, G'', of the 2%, 25% MG samples, as well as ascitic fluids isolated from OC patients were characterized using an Advanced Rheometric Expansion System (ARES) rheometer available from TA Instruments of New Castle, DE. The rotational rheometer was used with stainless steel parallel disks with 25 mm

diameters and had a force rebalance transducer 0.2K-FRTN1. The actuator of the ARES is a DC servomotor with a shaft supported by an air bearing with an angular displacement range of 5x10⁻⁶ to 0.5 rad, and angular frequency range of 1x10⁻⁵ to 100 rad/s. The angular velocity range is 1x10⁻⁶ to 200 rad/s. The sample loading procedure was the same for all the experiments and the gap height between two disks was kept constant at 1 mm. Sufficient volume of samples was used to fill the gap between parallel disks, and the linear viscoelastic properties of the samples were collected as a function of frequency at constant strain of 50% and room temperature. The samples were not presheared. For discussion of dynamic property characterization of complex fluids see, for example, Bird, R.B., Armstrong, R.C. and Hassager, O. "Dynamics of Polymeric Liquids. Wiley, 1987."

Author Contributions: S.A., T.P. and M.I. designed and conceptualized the studies with significant input from S.F., P.B., D.C., and D.K. IHC analysis of human tumors and PDX tissue sections were performed by S.A., and T.P. performed cell tracking experiments and image quantification. S.F. performed mesothelial clearance assays. P.B., L.Q., and D.Kh. designed and performed shear-stress rheometric analyses. P.B. and W.L. designed and performed electron microscopy experiments. M.R., P.J., E.D., E.N-M. and M.P-Z., identified human tumor samples and performed pathologic analysis of laminin γ1 expression in tumors representing disease before and after chemotherapy. S.A., T.P., and M.I. wrote a manuscript with significant input (methodology description) from S.F., P.B., L.Q., D.Kh., S.W., and M.P-Z.

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Competing Interest Statement. The authors declare no competing interests

Supplementary Movies.

MOVIE 1. Outgrowth formation by FNE-m-p53 suspended cellular clusters reconstituted with 2% MG. Cell clusters were imaged for 5 days at 2-hr intervals.

MOVIE 2. Outgrowth formation by Hey-A8 suspended cellular clusters reconstituted with 2% MG. Cell clusters were imaged for 5 days at 2-hr intervals.

MOVIE 3. Outgrowth formation by Tyk-nu suspended cellular clusters reconstituted with 2% MG. Cell clusters were imaged for 5 days at 2-hr intervals.

MOVIE 4. FNE-m-p53-GFP cell proliferation in 2% and 25% MG. Images were obtained every 22 minutes for a duration of 61 hrs.

MOVIE 5. Hey-A8-mKO2 cell proliferation in 2% and 25% MG. Images were obtained every 22 minutes for a duration of 68 hrs.

MOVIE 6. Evolution of trajectories made by FNE-m-p53-GFP monolayer cultures encapsulated with 2% or 25% MG. Images were obtained every 20 minutes for a duration of 61 hours. First frame of the movie corresponds to 13-hour time point.

MOVIE 7. Evolution of cell movement trajectories in Hey-A8-mKO2 monolayer cultures encapsulated with 2% or 25% MG. Images were obtained every 15 minutes for a duration of 68 hours. First frame of the movie corresponds to 20-hour time point.

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