



## Synthetic tumor networks for screening drug delivery systems



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### ABSTRACT

Tumor drug delivery is a complex phenomenon affected by several elements in addition to drug or delivery vehicle's physico-chemical properties. A key factor is tumor microvasculature with complex effects including convective transport, high interstitial pressure and enhanced vascular permeability due to the presence of "leaky vessels". Current *in vitro* models of the tumor microenvironment for evaluating drug delivery are oversimplified and, as a result, show poor correlation with *in vivo* performance. In this study, we report on the development of a novel microfluidic platform that models the tumor microenvironment more accurately, with physiologically and morphologically realistic microvasculature including endothelial cell lined leaky capillary vessels along with 3D solid tumors. Endothelial cells and 3D spheroids of cervical tumor cells were co-cultured in the networks. Drug vehicle screening was demonstrated using GFP gene delivery by different formulations of nanopolymers. The synthetic tumor network was successful in predicting *in vivo* delivery efficiencies of the drug vehicles. The developed assay will have critical applications both in basic research, where it can be used to develop next generation delivery vehicles, and in drug discovery where it can be used to study drug transport and delivery efficacy in realistic tumor microenvironment, thereby enabling drug compound and/or delivery vehicle screening.

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### 1. Introduction

In recent years, myriad technologies have been employed to deliver novel cancer therapeutics ranging from antibodies, cytokines, gene therapy and traditional chemical drugs to tumors. Furthermore, drug delivery vehicles ranging from viral (e.g., adenovirus, lentivirus) and non-viral vectors (e.g., polymers, liposomes, nanoparticles) have been developed [1–3] to enhance the delivery performance. The efficacy of any new therapeutic in eradicating tumors depends critically on uniform and effective delivery of the drugs [4–6] to all the tumor cells. The possibility of even a single cell to not come in contact with the drug can lead to regeneration of tumors and even worse, one that is drug-resistant [6–9].

High-efficiency drug delivery to tumors is a daunting challenge and rendered difficult primarily due to the complexity of the tumor microenvironment. The tumor microenvironment [9,10] is highly heterogeneous comprising of tumor and stromal cells (e.g., fibroblasts, inflammatory cells) embedded in an extracellular matrix connected to a vascular supply for nutrients. It also has gradients of cell proliferation and differential regions of hypoxia and acidity. In addition, solid tumors which account for more than 85% of the cancers have less than 10% of

blood vessels. One of the unique features of the tumor vasculature is their leakiness as a result of the discontinuity of the endothelium [11,12]. Studies using *in vivo* data have shown that the pore size of the leaky vessels ranges from 100s of nanometer to a few microns in a mouse mammary carcinoma [13]. In comparison, the vascular permeability in normal tissues is typically less than 6 nm [14] with the largest size of 150 nm in spleen endothelium [15].

Several *in vivo* techniques have been developed to study tumor drug delivery. A commonly used model employs windowed chambers in dorsal skin [16–18] or brain models [19,20] to study drug distribution. A relatively new method is the use of systems like the IVIS® optical imaging system (PerkinElmer, Waltham, MA) that can detect non-invasively fluorescent tags in live animals. However, such *in vivo* studies are expensive and require skilled personnel due to the use of live animals.

In contrast, *in vitro* models are a cost-effective means to study and screen drug delivery vehicles. In classical studies, the delivery vehicle containing the therapeutic of interest (drug/fluorescent tags) is incubated with the tumor cells in culture. At regular time points, the cells are analyzed either for uptake of the fluorescent tags or reduction in cell proliferation as a measure of delivery efficacy. Improvements to monolayer experiments in tissue culture have led to the development of *in vitro* methods which use multicellular tumor spheroids [21–23]. However, these static methods [24] do not account for transport across the vascular endothelium and the complex microvascular network structure observed *in vivo*. Furthermore, depending of the model, they

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rely exclusively on diffusion for the drugs to permeate the tumors and do not allow real-time visualization to study the diffusion of the delivery vehicle and/or drugs due to the use of semi-permeable membranes. Recent research has focused on the development of microfluidic devices to study cellular behavior under fluidic conditions [25–28]. Studies incorporating angiogenesis, tumor growth, invasion and tumor–endothelial cell interactions have also been reported [29–35]. However, all of these devices are not well-suited for the study of tumor drug delivery vehicles in conditions representing *in vivo* scenarios.

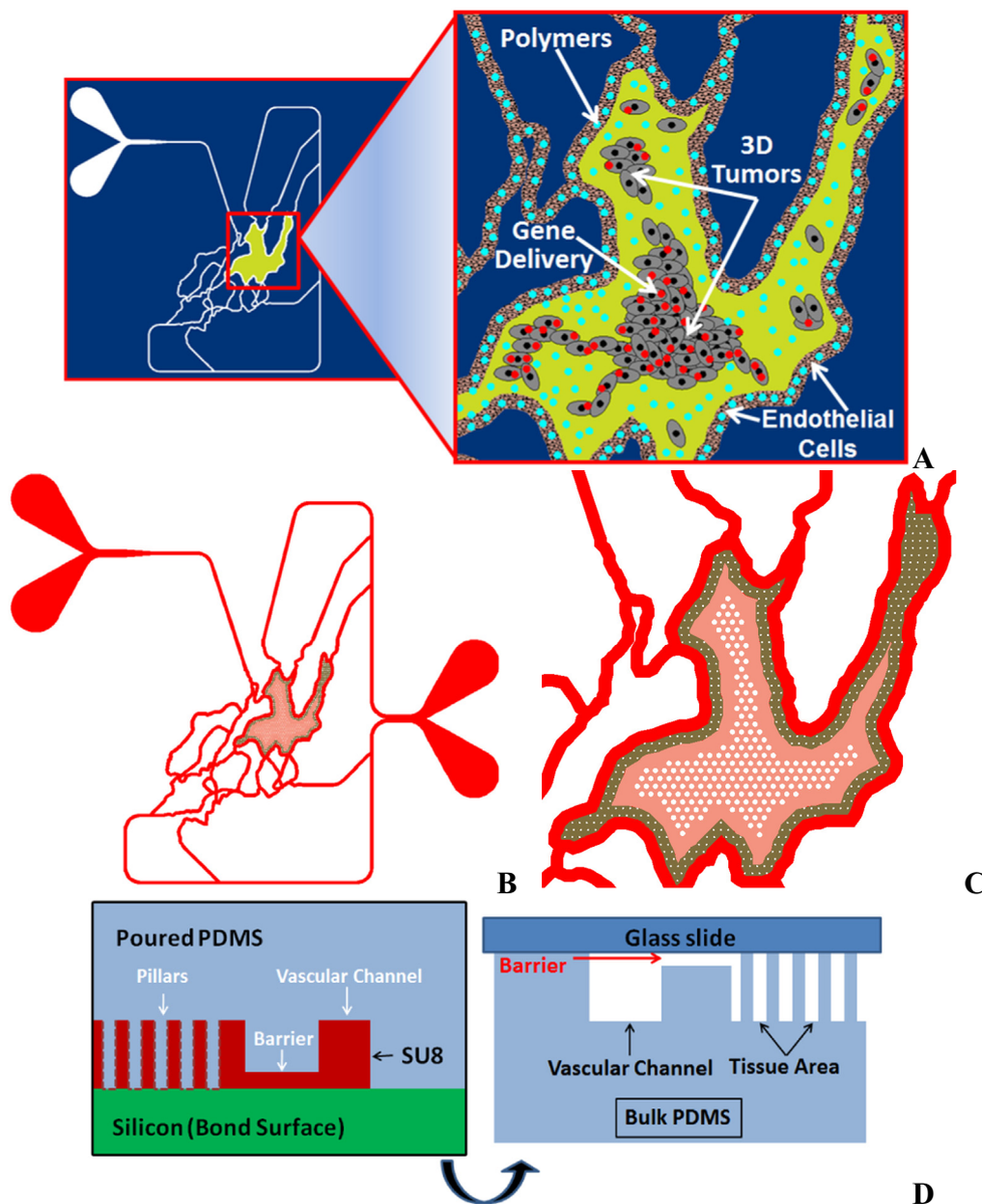
In this study, we report on the development of a microfluidics based synthetic vasculature assay that models the tumor microenvironment observed *in vivo*. This synthetic tumor network builds upon our previous work where we developed a novel methodology for reproducing microvascular networks digitized from *in vivo* images of rodent vasculature onto a microfluidic device [36–38]. The microfluidic device recreates the *in vivo* tumor microenvironment encompassing (a) circulatory flow in the vessels derived from *in vivo* morphology, (b) transport across the leaky vessel walls based on engineered barriers between

the vascular and the tumor cells, and (c) delivery to 3D culture of tumor cells across the interstitial space. The combination of these features distinguishes the present synthetic tumor network model from other *in vitro* models discussed above. Two nanopolymeric based gene delivery systems were tested and the results were compared with *in vivo* rodent data highlighting the predictive ability of the microfluidic device and assay.

## 2. Materials and methods

### 2.1. Design of synthetic tumor network

The microvascular network digitized previously [37] was modified to include regions for growth of tumors and the leaky gaps between the vessel lumen and the tumor growth region. The largest tissue area from the network was selected and the vessel wall adjacent to the tumor growth region was modified in AutoCAD to include 2  $\mu\text{m}$  size leaky gaps, typical pore size found in MCA-IV mouse mammary



**Fig. 1.** Design of synthetic tumor network. A. Concept. B. Schematic showing the vascular channels for culturing endothelial cells and the tissue compartment for culturing tumor cells. C. Magnified view of the tissue chamber showing the scaffolds for the 3D tumor. D. Concept with side view showing the 2  $\mu\text{m}$  leaky gaps.

carcinomas vessel walls [13]. A cylindrical micro-pillar array with prescribed dimensions of 50  $\mu\text{m}$  diameter, 100  $\mu\text{m}$  height and 50  $\mu\text{m}$  spacing was designed to create a scaffold for 3D tumor growth in the tissue area. Fig. 1A shows a schematic of the synthetic tumor network and Fig. 1B–C shows the image highlighting the microfabricated pillars for 3D culture. Fig. 1D shows the side view schematic of the 2  $\mu\text{m}$  leaky gaps and the microfabricated scaffolds.

## 2.2. Microfabrication of synthetic tumor network

The designed devices were fabricated using PDMS based soft-lithography. The tumor area was separated from the vascular channels using the barrier method shown in Fig. 1D. The barrier is structured on SU-8 by patterning an extra layer in addition to the fluidic layer, which contained the pillars, channels and access port holes, to form a thin slab between the tissue area and the vascular channels. The two step fabrication process for the soft lithography masters was as follows: (a) 500  $\mu\text{m}$  thick, 4" diameter p-type Si wafers were organically cleaned and dehydrated @ 200 °C for 5 min, (b) SU8 spin deposition to obtain 2  $\mu\text{m}$  film, (c) hot plate @ 65 °C for 1 min  $\rightarrow$  95 °C for 2 min, (d) exposure at  $\sim$ 250  $\text{mJ}/\text{cm}^2$ , (e) hot plate @ 65 °C for 1 min  $\rightarrow$  95 °C for 1 min (allow to cool for 10 min), (f) develop in PGMEA (SU8 developer) until field clears (<1 min), (g) spin coat fluidic SU8 layer @ 100  $\mu\text{m}$  over existing features, (h) hot plate @ 95 °C for 30 min, (i) exposure @  $\sim$ 250  $\text{mJ}/\text{cm}^2$ , (j) hot plate @ 65 °C for 1 min  $\rightarrow$  95 °C for 5 min (allow to cool for 10 min), (k) develop in PGMEA (SU8 developer) until field clears (5 min) and finally rinse with IPA. SEM images of the SU-8 masters were acquired using a Hitachi S-2600 N (Hitachi High Technologies America, Inc., Pleasanton, CA) scanning electron microscope. Samples were coated with 50 nm of Au using a Hummer 6.2 sputtering system (Anatech Ltd., Union City, CA). An acceleration voltage of 15 kV was used for subsequent imaging.

Sylgard 184 PDMS (Dow Corning) was poured over the developed master to generate devices in PDMS and cured at 60 °C overnight in an oven, following which the PDMS was peeled off from the master. Through holes, defining the inlets and outlets, were punched using a 1.5 mm biopsy punch. For injection of tumor cells, a 30 gauge blunt and sharpened needle was used to punch holes in the tumor area using a stereo microscope for proper alignment of the access port. The surfaces of the PDMS and a pre-cleaned glass slide were cleaned using oxygen plasma treatment prior to bonding. Tygon Microbore tubing with an outside diameter of 0.06 in and inner diameter of 0.02 in. served as the connecting ports for fluidic interface.

## 2.3. Fluidic testing

A fluorescent marker (FITC) was used to visualize the leakiness of the fabricated synthetic tumor network. FITC at a concentration of 10  $\mu\text{g}/\text{ml}$  was injected into the network using a syringe pump (PHD 2000, Harvard Apparatus, MA) at a flow rate of 1  $\mu\text{l}/\text{min}$ . An image of the entire device was acquired using an automated stage (LEP Ltd) mounted on an inverted fluorescence microscopy system (NIKON, Melville, NY). Images were visualized using NIKON Elements software. In order to test the leakiness of the 2  $\mu\text{m}$  barriers, 1  $\mu\text{m}$  and 5  $\mu\text{m}$  fluorescent particles (Fisher Scientific, Pittsburgh, PA) were injected into the vascular channel and their penetration into the tumor chamber was visualized.

## 2.4. Co-culture of endothelial and tumor cells in synthetic tumor network

We utilized an immortalized endothelial cell line, RBE4 (courtesy of Dr. Michael Aschner, Vanderbilt University Medical Center, Nashville, TN) to represent the vascular cells while the commonly used tumor cell line (HeLa-cervical cancer) was chosen to represent the tumor cells in the synthetic tumor network. RBE4 cells were cultured in Eagle's Minimum Essential Medium and Ham's F-10 media (1:1)

supplemented with 10% FBS, 1% Pen/Strep, 2 mM L-glutamine and G418 (300  $\mu\text{g}/\text{mL}$ ). Cells were incubated at 37 °C, 95% humidity and 5%  $\text{CO}_2$  until confluent. HeLa cells were obtained from ATCC (#CCL-2™) and maintained in DMEM media with 10% serum supplemented with, 4 mM L-glutamine and 100 U/ml penicillin/streptomycin on T25 tissue culture flask at 37 °C in 5%  $\text{CO}_2$ . Confluent cells for both types were trypsinized and sub-cultured at a ratio of 1:3 until ready for experiments.

HeLa cells ( $\sim$ 10<sup>7</sup>/ml) were harvested and mixed in a ratio of 1:3 with cold Matrigel™ for a total volume of 50  $\mu\text{l}$ . The solution was mixed uniformly and 10  $\mu\text{l}$  of the solution injected slowly into the tumor area access port of the device. The device was kept on an ice bath until this process was completed. Sterile cell culture media without serum was continuously perfused in the vascular channel side at a flow rate of 10  $\mu\text{l}/\text{min}$  to flush out any HeLa cells entering the vessel lumen. The device was then incubated overnight at 37 °C, 95% humidity and 5%  $\text{CO}_2$  until confluent. The next morning, fibronectin at a concentration of 50  $\mu\text{g}/\text{ml}$  and flow rate of 1  $\mu\text{l}/\text{min}$  was injected into the vascular channels for 30 min followed by incubation for another 30 min. Endothelial cells were trypsinized and injected into the vascular channels at concentration of  $5 \times 10^6$  cells/ml. Flow was stopped for 30 min by clamping the inlet and outlet for 2 h. At the end of 2 h, fresh media (RBE4 media mixed with HeLa media at 1:1) was injected into the channels overnight and allowed to perfuse overnight at a flow rate of 0.1  $\mu\text{l}/\text{min}$ . RBE4 cells and HeLa cells were allowed to grow together for additional 24 h prior to initiation of the delivery system screening experiments. Co-cultured RBE4 cells and HeLa cells were assayed using Calcein AM (Life Technologies, Carlsbad, CA), a cell-permeant dye used to determine cell viability.

## 2.5. Delivery system screening in synthetic tumor network

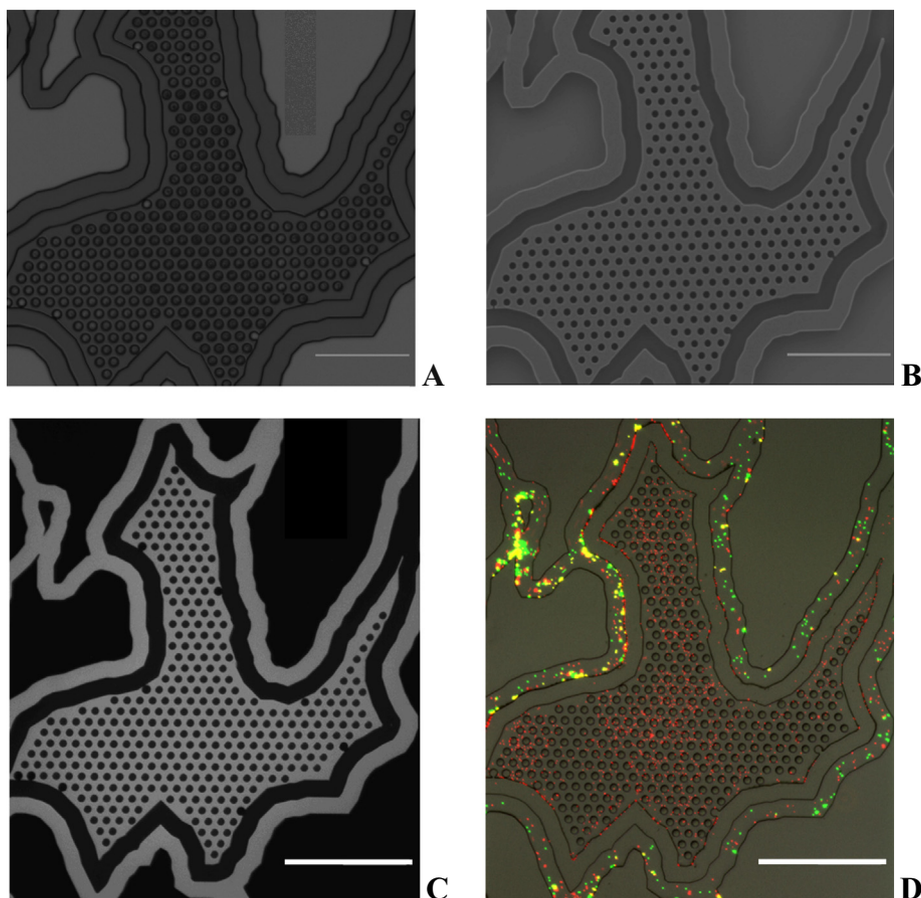
We compared two nanopolymer based gene delivery systems: (1) PPC, and (2) Express-In; in the synthetic tumor network. Express-In is a commercially available polymer based transfection reagent that has been shown to produce very high transfection activity in a variety of cell types in vitro [39]. PPC is a polymeric delivery system that has been shown to efficiently deliver plasmid in vivo [40] and has been tested in clinically for the delivery of IL-12 plasmid in ovarian cancer patients with recurrent platinum resistant ovarian cancer [41,42]. When tested in vivo (intraperitoneal delivery), Express-In is associated with relatively high levels of toxicity in contrast to PPC which has shown to be well tolerated in both pre-clinical and clinical studies [40–42]. The two polymers were labeled with Rhodamine (fluorescent tag) at a ratio of 3.6:1 wt/wt and were then complexed with GFP encoding DNA for a total concentration of 10  $\mu\text{g}/\text{ml}$ . This concentration is the optimal concentration utilized for the transfection studies.

Assays were conducted in two ways. In the first assay, HeLa cell transfection was monitored following polymer injection from the vascular channel. In the second assay, HeLa cell transfection was monitored by injecting the complexed polymers directly at the tumor site. To ensure that at least 3 $\times$  volume of the polymer/GFP complex was circulated in the networks, the complexes were injected into the network for 30 min at a flow rate of 0.5  $\mu\text{l}/\text{min}$ . At the end of 30 min, flow was immediately switched to cell culture medium comprising of 1:1 of RBE4 and HeLa culture media. A circulating flow was maintained for 24 h before GFP expression was measured in the tumor area.

## 3. Results

### 3.1. Fabrication and testing of synthetic tumor network

Fig. 2A shows the image of the microfabricated synthetic tumor network highlighting the vascular channel, walled barrier and the tumor chamber. Fig. 2B shows the SEM image with a detailed pattern of the

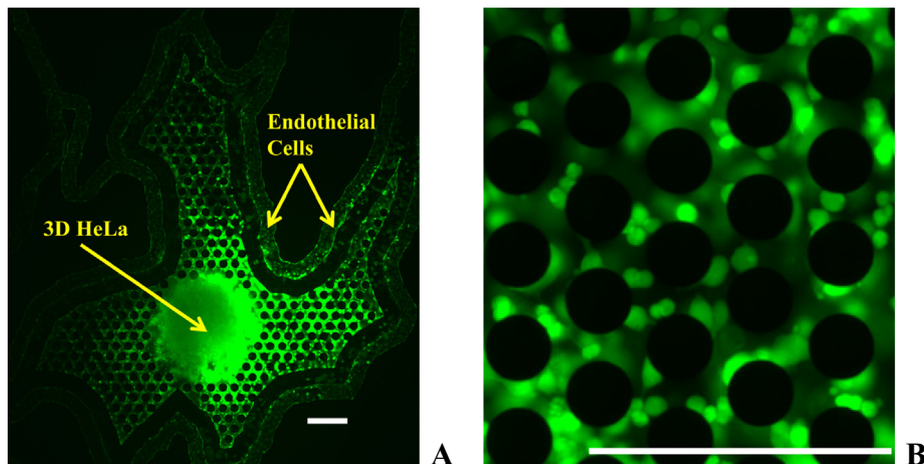


**Fig. 2.** Microfabricated synthetic tumor network. A. Optical image of the SU-8 master with the wall barrier. B. SEM Image showing the tissue area in the network with microfabricated pillars for tumor growth. C. FITC perfused device indicating fluidically connected vascular and tissue chambers. D. Particles (1  $\mu\text{m}$ — red; 5  $\mu\text{m}$ — green) perfused device indicating a leaky barrier. 1  $\mu\text{m}$  particle freely perfuse to the tumor area whereas 5  $\mu\text{m}$  are restricted to vascular channel. Scale bars: 500  $\mu\text{m}$ .

microfabricated pillars, which are used as scaffolds for 3D culture of tumor cells. Fig. 2C shows the network perfused with the fluorescent dye and Fig. 2D shows an image of the network perfused with the particles highlighting the intact barrier between the tumor and vascular channels. In addition, the images demonstrate that the tumor area with microfabricated scaffolds for the 3D culture of tumor cells is fully functional with the 2  $\mu\text{m}$  leaky vasculature.

### 3.2. Co-culture of endothelial and tumor cells in synthetic tumor network

Fig. 3 shows the network with HeLa cells cultured in 3D in the tumor region and endothelial cells in the vascular region stained with calcein AM. As can be seen from the images, the cells (endothelial and tumor cells) were in healthy condition. In addition, several smaller 3D spheroids were observed growing around the micropillars with varying



**Fig. 3.** Co-culture of tumor (HeLa) and vascular (endothelial) cells in synthetic tumor networks. A. Co-culture of 3D HeLa Cells cultured on microfabricated scaffolds and endothelial cells cultured in the vascular lumen separated by the walled barrier. B. Magnified view of the microfabricated scaffolds showing daughter HeLa cell colonies. Scale bars: 250  $\mu\text{m}$ .

number of cellular colonies at each of the location. Uniform calcein AM labeled cells indicate a fully active co-culture system of endothelial cells and tumor cells in the network.

### 3.3. Delivery system screening in synthetic tumor network

Fig. 4 shows images of Rhodamine labeled Express-In (Fig. 4A) and Rhodamine labeled PPC polymers (Fig. 4B) in the synthetic tumor network. As can be observed, Rhodamine signal for Express-In is more intense in the vessel lumen compared to PPC which is more uniform across the vascular and the tumor regions. The increased fluorescence intensity is indicative of particle aggregation, which is not seen in the case of PPC, where minimal aggregation of particles is observed. In addition, Express-In is found to have more aggregation near the tortuous bends and turns of the network compared to linear sections of the network. Finally, PPC is found to be more uniformly dispersed in the tissue chamber compared to Express-In.

Fig. 5A shows GFP expression of the 3D tumor mass using Express-In while Fig. 5B shows GFP expression of the 3D tumor mass using PPC polymers following vascular injection. PPC polymer based GFP transfection is more uniform showing a relatively constant amount of expression across the entire tumor. However, Express-In shows non-uniform GFP expression and the core of the tumor is poorly transfected.

Surprisingly, when injected directly to the tumor site, Express-In based GFP expression was found to be more intense, although both of the polymers exhibited GFP expression (Fig. 5C–D). Fig. 5E shows the quantitative intensity values of the GFP expression using PPC and Express-In polymers for both the direct and vascular injection test conditions. Furthermore, these findings match the in vivo delivery performance of several plasmid based approaches including PPC and Express-In where intra-tumoral injection has shown uniform transfection while intra-peritoneal injection has shown poor transfections [43]. These findings also serve as positive controls indicating that the cells, the polymer and the GFP DNA complex are functional. These results clearly establish the fact that synthetic tumor network is able to predict the drug vehicle characteristics in vivo based on the drug injection route [40–42].

## 4. Discussion

The efficacy of a drug reaching its desired location is dependent upon the attributes of the delivery system. Hence, it's imperative that the delivery system is able to maintain its functional properties in the context of the in vivo environment. Highly complex physical and biological conditions exist in this environment including flow, cell–cell and cell–particle interactions. Unfortunately, standard in vitro tests comprising of static well plate incubation severely misrepresent the in vivo scenario and thus cannot adequately predict or provide a realistic understanding of the properties and behavior of a molecule or particle in vivo.

In this study, a clinical grade polymer PPC and an in vitro grade polymer Express-In were used for transfection of 3D tumors by complexing

with GFP expressing plasmid DNA. In vivo preclinical studies following intra-tumoral injection have shown that these systems behave similarly [43,44] while intra-peritoneal injection and subsequent clinical studies showed that only PPC is fully effective [40–42]. Well plate studies, in contrast, while allowing for direct injection (data not shown) cannot reproduce intraperitoneal or vascular injection scenarios for comparison with in vivo data. Assays in the synthetic tumor network reproduced the exact scenario observed in vivo. Vascular injection of polymers demonstrated higher efficiency for PPC while direct tumor injection showed similar results for both the polymers, although Express-In based GFP expression signal was brighter.

The poor efficacy of Express-In can be attributed due to the fact that serum proteins under flow interact with Express-In, causing it to aggregate to an extent that presents a steric obstacle to uniform transfection. On the other hand, PPC, which remains relatively aggregation-free, is able to flow freely and transfect cells uniformly. The translational diffusion coefficient values can be calculated using the Stokes–Einstein equation and viscosity of cell medium at 0.78 cP [45] and are between 11.7 and 3.69  $\mu\text{m}^2/\text{s}$  for Express-In (diameter range of 50–150 nm) and 11.7 and 3.16  $\mu\text{m}^2$  for PPC (diameter range of 50–175 nm). The diffusivity of these polymers is comparable with those reported in the literature [44,46]. Both of these delivery systems are highly cationic due to the polyethyleneimine (PEI) core structure which allows for the condensation of plasmid DNA into nanoparticles. However, PPC is further modified by the addition of a polyethylene glycol (PEG) which improves serum stability through molecular shielding of the cationic charge [40–42]. The high toxicity of Express-In in vivo (data not shown) may be presumably due to interaction with blood proteins, opsonization and aggregation of the nanoparticle complexes which is significantly attenuated with PPC. In addition, flow and polymer interaction with the cells may express receptors on the cell surface for uptake of GFP which is again not possible to test in static well plate conditions. Detailed studies need to be conducted to understand these cell receptor and delivery system ligand interactions. Although, in the current study, none of the polymers were targeted specifically to the tumor, directed approaches will allow more focused delivery of the drugs or genes to the desired location. In addition, optimization of delivery system receptor type and density to maximize binding strength can be readily tested in the developed assay.

Drug delivery systems come in all shapes and sizes. Recent studies have shown that rod shaped particles have greater binding affinity than spheres for both micro- and nano-sized delivery systems [47–49]. In addition, even simple flow based systems have shown differences in binding affinities compared to static well plate assays [50,51]. The synthetic tumor networks developed in this study can be used to optimize the size and shape of delivery systems in conjugation with targeted receptors.

Drug toxicity is of critical importance in evaluating drugs for efficacy. In this study, we did not focus on the toxicity of the delivery systems to the tumors or the normal cells (endothelial). Studies incorporating toxicity analysis for delivery systems, drugs, etc. will be pursued in the future. An interesting study will be to investigate the difference between a

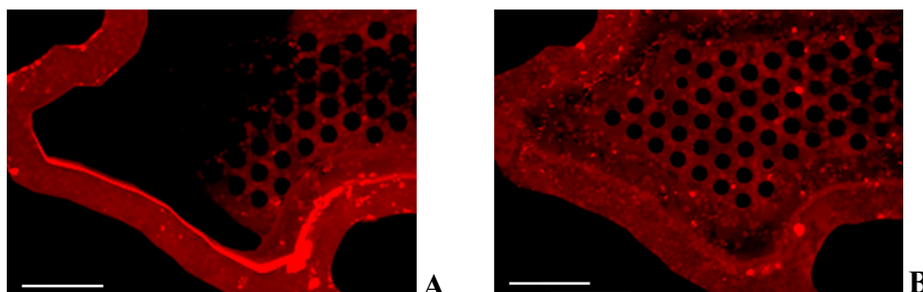
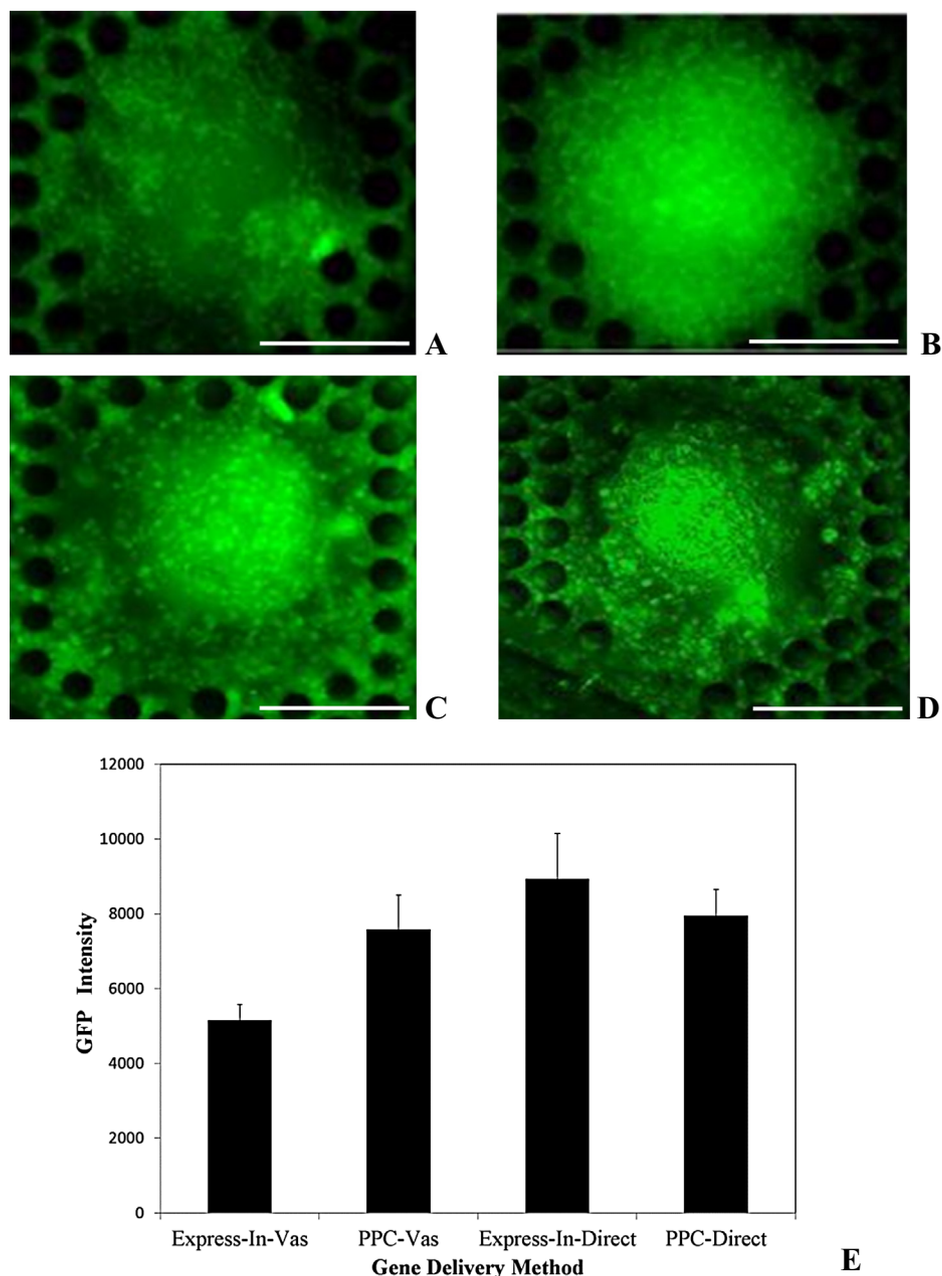


Fig. 4. Polymers in synthetic tumor network. A. Express-In shows non-uniform distribution and significant aggregation. B. PPC shows uniform distribution and minimal aggregation. Scale bars: 250  $\mu\text{m}$ .



**Fig. 5.** Delivery system screening. A. Express-In based GFP transfection following vascular injection. Non-uniform and minimal GFP expression is observed on the periphery of the tumor while the core remains untransfected. B. PPC based GFP transfection following vascular injection. Uniform and intense GFP expression and transfected core are observed. C. Express-In based GFP transfection following direct injection. D. PPC based GFP transfection following direct injection. Both polymers demonstrate uniform expression following direct injection. E. GFP intensity comparison for PPC and Express-In following vascular and direct injection. PPC and Express-In perform similarly following direct injection. PPC performs significantly better than Express-In following vascular injection. Data is shown as mean  $\pm$  S.D. with experiments performed in triplicates. Scale bars: 250  $\mu$ m.

bolus injection of drug vs. a constant infusion and the tradeoff between efficacy and toxicity. Conditions of gradients of nutrients and oxygen diffusion can also be tested in these systems which will again allow more realistic test conditions. A significant advantage of using microfluidic based systems is the savings in reagents and time compared to standard well plate assays.

The developed synthetic tumor network device and assay provides an ideal *in vitro* platform to test the efficiency of delivery systems under conditions mimicking physiological situations. Different from other microfluidic *in vitro* tumor models reported in the literature [25–35], the developed synthetic tumor network model replicates the morphology, fluidics and leaky vasculature observed *in vivo*, specifically (a) *in vivo* based vascular morphology, (b) engineered leaky gaps between the vessels and the tumor, and (c) 3D culture of tumor cells.

The leakiness of the vasculature used in this study was 2  $\mu$ m. However, this can be readily modified from a few nanometers to several tens of micrometers to account for heavily leaky vessels or non-leaky portions of the vasculature. The developed synthetic tumor network model can be used to study the mechanisms of drug delivery vehicle transport, drug–cell interactions, tumor transfection, and tumor–endothelium interactions.

## 5. Conclusion

Well plate assays routinely used to assess performance of drug delivery systems do not predict *in vivo* responses. In this study, synthetic tumor networks clearly demonstrated its utility in accurately predicting *in vivo* behavior. Both the GFP gene delivery nanoparticles

studied here – PPC and Express-In – showed similar high efficiency transfection results using intra-tumoral injection. In contrast, intra-peritoneal administration *in vivo* showed uniform transfection for PPC and poor transfection for Express-In similar to the results obtained from the synthetic tumor network assays.

Synthetic tumor network assay allows replication of *in vivo* conditions comprising of morphology from *in vivo* vascular networks, co-culture of endothelial cells under physiological fluid flow and 3D culture of tumor cells, as well as the leakiness of the tumor vasculature in an *in vitro* model. The developed system and assay can be used to study cell–cell and cell–particle interactions and will have significant applications in basic and applied research, where it can be used to characterize and develop next generation delivery vehicles, and in drug discovery where it can be used to study the efficacy of the drug in realistic tumor microvascular networks.

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