Patterned Hydrogel Substrates for Cell Culture with Electrohydrodynamic Jet Printing

Michael J. Poellmann, Kira L. Barton, Sandipan Mishra, Amy J. Wagoner Johnson*

Cells respond to and are directed by physiochemical cues in their microenvironment, including geometry and substrate stiffness. The development of substrates for cell culture with precisely controlled physiochemical characteristics has the potential to advance the understanding of cell biology considerably. In this communication, E-jet printing is introduced as a method for creating high-resolution protein patterns on substrates with controlled elasticity. It is the first application of E-jet printing on a soft surface. Protein spots as small as 5 μm in diameter on polyacrylamide are demonstrated. The patterned hydrogels are shown to support cell attachment and spreading. Polyacrylamide substrates patterned by E-jet printing may be applied to further the study of cellular mechanobiology.
In this paper, we introduce electrohydrodynamic jet (E-jet) printing\textsuperscript{[9,10]} as a new method for printing ECM proteins with micro-scale pattern features on polyacrylamide substrates. In E-jet printing, an ink solution is placed in a sealed reservoir with a conductive nozzle. With the right balance between capillary force and the applied back-pressure on the fluid, the ink forms a spherical meniscus at the tip of the nozzle. Applying a potential difference between the nozzle and the substrate creates an electric field, pulling the fluid meniscus into a cone. It is through jetting from the tip of this cone that E-jet printing achieves high resolution.\textsuperscript{[5]} Droplet size and jetting frequency depend on the back-pressure, the separation distance between nozzle and substrate, and the applied voltage. The system can be operated in either direct current (DC) mode with droplets at a constant frequency or jetting in a continuous stream, or in pulsed-voltage mode with droplets controlled by a user-defined square wave voltage signal.\textsuperscript{[11]}

Most notably, this paper represents the first application of E-jet on a soft substrate. Protein printing is demonstrated on polyacrylamide substrates with different elasticities. We also demonstrate cell attachment and spreading on printed fibronectin. The feature sizes obtained are significantly smaller than single cells, and are comparable to or smaller than other methods for patterning hydrogel cell substrates, including inkjet printing\textsuperscript{[12,13]} and microcontact printing\textsuperscript{[6]}.

E-jet printing is a promising technique for creating substrates with subcellular-scale protein patterns on surfaces with well-defined mechanical properties for use in mechanobiology studies.

**Experimental Section**

**Substrate Preparation**

Glass cover slips were cleaned in a dilute soap solution then rinsed several times with de-ionized (DI) water and once with ethanol. The glass was prepared for gel attachment by treating for 10 min under 2 vol% 3-(trimethoxysilyl)propyl methacrylate (Sigma) in 95 vol% ethanol, pH 5, and placed on a 60 °C hotplate for 10 min. A prepolymer solution consisting of 8 vol% acrylamide (Bio-Rad), 0.2 vol% N,N,N′-methylenebis(acrylamide) (bis) (Bio-Rad), and 0.2 vol% acrylic acid (Sigma) was prepared in 0.137 M NaCl and 0.050 M 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) (Sigma), pH 8. Polymerization was initiated by adding 1/80 vol% ammonium persulfate (Bio-Rad) and 1/1000 vol% N,N,N′,N′-tetramethylethenediamine (Bio-Rad). For softer (ca. 9 kPa) and stiffer (ca. 35 kPa) gels, 6.0%/0.15% and 10.0%/0.25% acrylamide/bis (Bio-Rad), 2 vol% 3-(trimethoxysilyl)propyl methacrylate (Sigma) in 95 vol% 0.05 M 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (Sigma), rinsed with DI water, placed on a 60 °C hotplate for 5 min. Activated gels were stored in a dry environment prior to printing.

**E-Jet Printing**

The glass-bonded and activated hydrogels were placed on a 5-axis computer-controlled stage and connected to ground. IgG ink solutions consisted of 5 mg·mL\textsuperscript{-1} rabbit Immunoglobulin-G (Pierce), 20 vol% glycerin, and 0.1 vol% Tween-20 (Sigma) in phosphate buffered saline (PBS) (Lonza). Fibronectin inks contained 0.5 mg·mL\textsuperscript{-1} human fibronectin (BD Biosciences) in salt-free 0.050 M HEPES, pH 8 with 0.1% Tween-20 with 20 or 40 vol% glycerin for printing with 5 μm and 2 μm inner diameter nozzles, respectively. Ink was loaded in a pressure-sealed syringe clamped above the substrate with Luer-tipped micropipette nozzles (World Precision Instruments). Prior to printing, the nozzles were sputter-coated with a 5 nm layer of Au/Pd for conductivity and treated for 3 min in 1H,1H,2H,2H-perfluorodecane-1-thiol (Sigma). Voltage, substrate alignment, nozzle separation distance, and stage displacement were controlled through a custom-developed LabVIEW interface. Printing was monitored with an Infinity 3 camera (Luminera Scientific) with an Edmund Industrial Optics lens. Following printing, substrates were stored in a sealed container at room temperature for at least 4 h, then thoroughly rinsed in PBS to rehydrate and remove unreacted NHS groups. For cell culture, gels were sterilized by briefly soaking in 70 vol% ethanol then transferred to sterile 6 cm plates and rinsed several times with cell media.

**Pattern Characterization**

Hydrogels were rehydrated in phosphate buffered saline. IgG proteins were detected with Dylight 488-conjugated goat anti-rabbit secondary antibodies (Pierce) following treatment with a solution of 1 vol% goat serum (Pierce). Fibronectin was visualized after treatment with sheep anti-human fibronectin IgG (R&D Systems) and NI493-conjugated donkey anti-sheep IgG (R&D Systems). Images were obtained with a QImaging Retiga 2000R camera mounted on a Leica DMi 4000 B inverted fluorescent microscope. A Zeiss 710 confocal microscope was used for 3D imaging a 3 × 10 array of 10 μm diameter spots and a 2 × 6 array of 20 μm spots. Depth of penetration was determined using Zeiss ZEN software.

**Cell Culture**

Hydrogels patterned with fibronectin were sterilized in 70% ethanol and rinsed several times with PBS and media prior to cell seeding. D1 ORL UVA mouse mesenchymal stem cells (ATCC) and primary rat mesenchymal stem cells (Millipore) were grown at 37 °C in a humidified incubator under Dulbecco’s Modified Eagle Medium-low glucose, modified to contain 4 × 10\textsuperscript{-3} M l-glutamine, 1 × 10\textsuperscript{-3} M sodium pyruvate, and 3.7 g·L\textsuperscript{-1} sodium bicarbonate (UIUC Cell Media Facility) with 10% added fetal bovine serum (Gibco) and 100 U·mL\textsuperscript{-1} penicillin-streptomycin (Mediatech). The D1 cells were lifted at passage 10–12 with 0.05% trypsin-EDTA (Invitrogen), seeded on substrates at a density of 10\textsuperscript{4} cells per dish, and observed with phase contrast microscopy after 3 d. Primary cells were lifted at passage 3, seeded at a density of 5 × 10\textsuperscript{4} cells per dish.
After 24 h, the cells were fixed with neutral buffered formalin (Sigma), treated with 1 vol% Triton X-100 (Sigma), and stained with rhodamine phalloidin (Invitrogen) following the fibronectin antibody treatment. Image brightness, contrast, and coloring was adjusted using NIH ImageJ software was used to make composite fluorescent images.

Results and Discussion

The polyacrylamide substrates contain an added 0.2% acrylic acid into the polymer backbone. Those groups are subsequently modified with EDC/NHS to activate the gel for protein attachment. When proteins contact the activated hydrogel, amine-containing amino acids displace the NHS groups and form covalent bonds directly to the polymer backbone.[14] Fully-hydrated substrates shrink during printing, which is undesirable because it is critical to maintain a consistent distance between the printer nozzle and the substrate. To control for this, we baked the substrates on a hotplate to remove most of the water prior to printing. Aside from maintaining a constant stand-off height, a dry environment also better protects the activity of NHS groups.[14] Following printing, the remaining NHS groups are hydrolytically cleaved when the substrates are rehydrated with PBS, making unpatterned regions resistant to further protein adsorption. The gel activation and protein conjugation are summarized in Figure 1.

The activated substrates are electrically grounded and placed on a vacuum chuck attached to a computer-controlled translating stage for printing. The protein ink (described below) is loaded into a reservoir, below which there is a glass capillary nozzle that is sputter-coated with gold-palladium for enhancing conductivity. Ink droplet size and jetting frequency are controlled by computer-defined voltage and back-pressure. The E-jet system is illustrated in Figure 2.

We first printed rabbit immunoglobulin G (IgG) as a model protein in DC mode with the ink solutions containing 5 mg·mL⁻¹ IgG, 20% glycerin, and 0.05% Tween-20. Glycerin increases the viscosity of the solution and mitigates evaporation from the meniscus, which can cause the nozzle to clog. Tween-20 helps prevent protein aggregation. Printing typically occurs with a back-pressure of 1–3 psi and an applied voltage of 250–350 V. Nozzles have 5 μm inner diameter and are kept at a consistent stand-off height of 30 μm above the substrate. Droplets of various sizes are demonstrated on hydrogels with elastic moduli.

![Figure 1](image1.png)  
**Figure 1. Summary of bioconjugation reactions.** Polyacrylamide substrates contain 0.2% acrylic acid. The substrates are activated with NHS using EDC (top). Immediately following activation the substrates are partially dehydrated. Proteins printed on the dried substrates form covalent bonds to the gel (bottom). Exposed amine groups on the proteins displace the immobilized NHS groups.

![Figure 2](image2.png)  
**Figure 2. Diagram of E-jet setup.** The balance between the applied voltage, back pressure applied to the ink reservoir, and capillary pressure in the nozzle determines droplet size and frequency. The activated polyacrylamide hydrogel is grounded and attached to a translating and tilting computer-controlled stage.
ranging from 9.6 to 34.4 kPa, a range that is known to influence MSC differentiation. Following printing, IgG-patterned gels are washed and treated with fluorescent antibodies for visualization (Figure 3a-c). Confocal microscopy shows that proteins are located at the surface of gels where they would be accessible to cells (Figure 3c), but also penetrate approximately 30 μm into the gel.

ECM protein inks consisted of 0.5 mg·mL⁻¹ fibronectin and 0.05% Tween-20. To obtain features with higher resolution, we use 2 μm nozzles. We increase the glycerin content to 40% to reduce the potential for clogging the smaller nozzle. Substrates were treated with a primary antibody that binds fibronectin and a secondary antibody conjugated to a fluorophore (green) (Figure 3d-f). Spots and lines smaller than 5 μm across were obtained, with the smallest spots obtained using pulse control (Figure 3e). Substrates were seeded with mesenchymal stem cells following printing. The cells were allowed to attach and spread overnight before fixing and imaging. Primary mesenchymal stem cells were stained with rhodamine phalloidin, which binds to filamentous actin and appears red (Figure 4a-c). Additionally, D1 cells were patterned and imaged using phase contrast microscopy after 3 days of culture (Figure 4d,e). Cells were observed to adhere only to regions where fibronectin had been printed.

E-jet printing possesses superior qualities when compared with conventional inkjet printing, namely, a higher feature resolution. E-jet printed features are an order of magnitude smaller than inkjet printing, enabling patterning of single cells. Only microcontact printing offers comparable resolution, achieving line widths of less than 10 μm. However, the main disadvantage of microcontact printing is its inflexibility. Once fabricated, stamps cannot be altered, and fabricating a new stamp requires a clean room facility and a relatively high level of expertise. E-jet printing, in comparison, has high process flexibility and the pattern can be controlled at the point of printing. Additionally, E-jet systems can be assembled at relatively low cost.
Conclusion

We report a technique for patterning cell adhesion proteins with high resolution on polyacrylamide hydrogels using E-jet printing. The use of a hydrogel substrate is a novel application for E-jet, which is capable of creating patterns with dimensions significantly smaller than a single cell. This method can be employed to create patterned ECM proteins on surfaces with precisely controlled physical properties for cell culture.

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