Gelatin-Based Laser Direct-Write Technique for the Precise Spatial Patterning of Cells

Nathan R. Schiele, M.S.,¹ Douglas B. Chrisey, Ph.D.,¹,² and David T. Corr, Ph.D.¹

Laser direct-writing provides a method to pattern living cells in vitro, to study various cell–cell interactions, and to build cellular constructs. However, the materials typically used may limit its long-term application. By utilizing gelatin coatings on the print ribbon and growth surface, we developed a new approach for laser cell printing that overcomes the limitations of Matrigel®. Gelatin is free of growth factors and extraneous matrix components that may interfere with cellular processes under investigation. Gelatin-based laser direct-write was able to successfully pattern human dermal fibroblasts with high post-transfer viability (91% ± 3%) and no observed double-strand DNA damage. As seen with atomic force microscopy, gelatin offers a unique benefit in that it is present temporarily to allow cell transfer, but melts and is removed with incubation to reveal the desired application-specific growth surface. This provides unobstructed cellular growth after printing. Monitoring cell location after transfer, we show that melting and removal of gelatin does not affect cellular placement; cells maintained registry within 5.6 ± 2.5 μm to the initial pattern. This study demonstrates the effectiveness of gelatin in laser direct-writing to create spatially precise cell patterns with the potential for applications in tissue engineering, stem cell, and cancer research.

Introduction

A current limitation of many in vitro culture studies is the inability to precisely control the spatial relationship of cells to growth surface features, such as substrate modifications, or to other cells in culture. Precise control of cell location, for example, to within 10 μm, can be used to replicate the in vivo cellular microenvironment and spatial distribution, allowing investigation of cellular interactions in vitro. By controlling the cell’s location with respect to neighboring cells in culture, one can affect the mode of cellular signaling (direct cell contact, paracrine signaling, or endocrine signaling), as well as the types of cells in communication. Thus, custom in vitro co- or multicultures could be precisely designed to better understand cell–cell interactions as well as proximity-dependent cell fate decisions.

To achieve this level of precision in cellular assembly, a laser-based direct-writing technique, originally developed to print inks for passive microelectronic applications, was adapted to enable the patterning of biomaterials and living cells.¹ Previously, pulsed laser printing techniques have shown great promise for printing numerous types of mammalian cells.²⁻¹⁹ Pulsed laser cell depositions have shown high cell viability,⁴,⁵,⁸,¹¹⁻¹⁶ little to no DNA damage,⁶,¹⁶ unaltered apoptosis rates,⁵,¹⁶ little to no increase in heat shock protein expression,¹¹,¹⁵ and normal cell proliferation.⁸,¹¹,¹⁶

Current laser-based direct-write techniques have proven their efficacy and potential; however, many rely on commercially available basement membrane matrix, Matrigel® (BD Biosciences, Bedford, MA). Specifically, Matrigel has been used in the direct-write process either as a coating for the print ribbon, as a long-term growth surface on the receiving substrate, or both.²⁻⁴,⁶⁻⁸,¹⁰⁻¹³,¹⁵,¹⁶,¹⁸ Matrigel is useful for cell transfer as it cushions the impact at the receiving substrate, provides a scaffold for patterning three-dimensional cell constructs through layering, maintains a moist microenvironment, and possesses a wide array of extracellular matrix (ECM) proteins for cellular adhesion. However, despite these attributes, current laser direct-write techniques are limited in their scope and future application due to their reliance on Matrigel. The multiple intrinsic growth factor constituents of Matrigel—basic fibroblast growth factor, transforming growth factor-β, epidermal growth factor, insulin-like growth factor-1, and platelet-derived growth factor—can potentially act as extrinsic cues confounding the cellular processes under investigation, and thus may preclude or greatly limit the utility of laser direct-writing for precise cell cultures. Moreover, Matrigel is derived from murine tumors, and significant lot-to-lot variations exist in the constituents. Even small fluctuations in growth factor constituents can have a profound influence on cellular response. Further, for some applications the presence of
Gelatin is composed of acid-denatured collagen and has been used extensively for drug release and tissue engineering due to the biocompatibility, rapid biodegradability, known constituent purity, and the absence of growth factors. 23–26 In the present study, gelatin was uniformly spin coated onto the print ribbon and used to partially encapsulate trypsinized cells on the ribbon, providing a laser interaction and buffer zone to protect the cells. Moreover, on the receiving substrate and long-term growth surface, the gelatin coating cushions the impact of transfer while maintaining a moist microenvironment during the printing process. Further, gelatin melts at 37°C, 27 which allows it to be removed from the growth surface when placed in a standard cell culture incubator, thereby providing an unobstructed cellular growth surface. The ability to remove the gelatin layer could provide potential new applications; however, it could introduce an inherent obstacle for maintaining cell viability and pattern registry, and thus can be confounding extraneous growth factors, and thus can be utilized in studies involving cell types highly sensitive to external signals from ECM components and growth factors, such as cancer cells and stem cells. As such, gelatin-based laser direct-writing provides a solution for a variety of biomedical applications requiring precise cell patterning, particularly in the area of tissue engineering and regenerative medicine.

Materials and Methods

Laser system

The matrix-assisted pulsed laser evaporation direct-write (MAPLE DW) system used in these experiments incorporates a pulsed excimer laser (TeoSys, Crofton, MD) operating at a wavelength of 193 nm argon-fluorine (ArF), coupled with computer-aided design (CAD)/computer-aided manufacturing (CAM) control (Fig. 1). The laser beam has a near-Gaussian distribution, a pulse width of 8 ns, and a repetition rate that can be varied from 1 up to 300 Hz. The beam is transmitted to the ribbon through an intracavity variable aperture, a series of mirrors, two irises to set the spot size, and lastly though a 15x objective to focus the beam. An x-y motorized ribbon stage, mounted above the x-y motorized receiving stage, is controlled via a software interface that allows for user-specified motion of the ribbon, independent of the receiving substrate. Patterns used for cell deposition can be drawn in a commercial two-dimensional CAD package and converted into motion and laser-firing code. For this study, user-specified pattern arrays were written in a g-code format, which controlled the motion of the x-y motorized receiving stage. CAD/CAM control allows the user to specify the distance between transferred spots of cells within the machine code and thus prescribe the location on the receiving substrate and proximity to other cells within the user-defined pattern. Independent ribbon control allows the user to target specific cells, or groups of cells, on the print ribbon, thereby offering even greater control of cell number within the pattern. Additionally, the number of cells in each transferred spot can be modulated through adjustments in spot size and cell density. Cell density in the transferred spot can be adjusted by changing the density of trypsinized cells on the print ribbon, and by adjusting the laser beam diameter using a series of irises, the

![FIG. 1. Schematic of the matrix-assisted pulsed laser evaporation direct-write system used to print trypsinized cells. (Inset) An image of the cells on the print ribbon highlights the optical clarity of the gelatin, which is necessary to observe cells, thereby allowing for the targeting of specific cells for transfer, and verification of proper placement on the growth surface.](image-url)
size of the transferred spot of cells (diameter ~20–500 μm) can be prescribed. Further, the distance between the print ribbon and receiving substrate can be ranged from ~1 cm down to ~100 μm using a z-stage translator. An in situ energy meter is used to record the energy of every laser pulse to track shot-to-shot repeatability and ensure that the appropriate energy is delivered to the ribbon.

The MAPLE DW system also contains an in situ charged-couple device camera that shares the optical path with the laser as it travels through the final objective. This allows the user to observe the trypsinized cells on the ribbon before transfer (Fig. 1, inset), target specific cells in real time, and verify the transfer of cells from the ribbon to the receiving dish.

Print ribbon and receiving dish preparation

Gelatin preparation. A 10 wt% concentration of gelatin was prepared for the receiving substrate by mixing 1.0 g of porcine skin-derived type A gelatin (Sigma-Aldrich, St. Louis, MO) with 10 mL of Dulbecco’s modified Eagle’s medium (DMEM) and heated to obtain a homogenous mixture. A 20 wt% concentration of gelatin was created for the ribbon by mixing 4 g of gelatin with 20 mL of sterile cell culture-grade water.

Receiving dish preparation. A 100-mm-diameter Petri dish was mounted on the x–y computer-controlled motorized stage, and a fiduciary mark was laser micromachined into the bottom of the dish for use in tracking cellular pattern registry.

The marked Petri dishes were plasma cleaned for 1 min, and coated with 1.5 mL of poly-L-lysine (PLL) hydrobromide (Sigma-Aldrich). After 5 min, the excess PLL was aspirated from the dish and the dish was left to dry in a laminar flow hood for 1 h. The PLL-coated receiving dish was then mounted on a bench-top spin coater, and 1.0 mL of 10% gelatin, warmed to 60 °C, was pipetted onto the receiving dish while spinning at 4000 rpm, for 25 s. The dish was placed into a refrigerator (4 °C) for 5 min at which time, 10 mL of DMEM, also at 4 °C, was pipetted over the dish and excess DMEM was aspirated. The receiving dish was placed into a standard cell culture incubator (37 °C, 5% CO₂, and 95% RH) for ~20 min.

Ribbon preparation. A 50-mm-diameter UV transparent quartz flat disk (“ribbon”) (Edmund Optics, Barrington, NJ) was cleaned with 70% ethanol, dried, and mounted on a bench-top spin coater. Gelatin (20%) was warmed to 60 °C and pipetted (1.5 mL) onto the ribbon while spinning at 2000 rpm, for 20 s. The ribbon was then placed directly into a standard cell culture incubator for 3 min.

Cell culture and direct-write

Human dermal fibroblast cells (ATCC, Manassas, VA) were grown in a standard cell culture incubator in 75-cm² T-flasks (BD Biosciences) using culture medium (89.5% DMEM, supplemented with 10% fetal bovine serum and 0.5% penicillin-streptomycin). One milliliter of cells from a cell suspension (density ~5.9×10⁵ cells/mL) was pipetted into the 20% gelatin-coated ribbon, incubated for 7 min, and placed in a laminar flow hood for 4 min. The ribbon was then tilted to remove excess culture medium, blotted dry, inverted, and mounted onto the ribbon stage. The laser beam diameter was set to 50 μm to achieve a fluence of ~1.0 J/cm². The 10% gelatin-coated receiving dish was mounted onto the receiving stage and moved to within ~500 μm of the ribbon using the z-stage translator. Independent ribbon control allowed the user to target specific spots on the ribbon for transfer to ensure a fully populated cellular array, and the user-generated g-code controlled the receiving dish, to pattern customized arrays of cells. A 2×2 array program with 500-μm spacing between spots was executed, pulsing the laser once for each corresponding targeted transfer spot, depositing the cells in proximity to the fiduciary mark on the receiving dish.

After cells were transferred, from the ribbon to the receiving dish, they were imaged on an inverted optical microscope (Carl Zeiss, Thornwood, NY) with phase contrast (10×). Once imaged, the receiving dish was placed in an incubator for 15 min to allow for initial cell attachment to the growth surface, at which time, 10 mL of fresh cell culture medium, warmed to 37 °C, was added and the dish was returned to the incubator.

DNA damage analysis and cell viability

To determine whether gelatin-based laser direct-write induces double-strand DNA breaks, the presence of phosphorylated-H2AX (phospho-H2AX) was quantified immunocytochemically on cell transfer arrays (n = 4). H2AX is a histone that is phosphorylated when a double-strand DNA break occurs and aids in recruiting proteins responsible for double-strand break repair. The colocalization of nuclear stain and immunostaining for phospho-H2AX has been previously used to verify DNA damage of human fibroblasts when exposed to UV radiation. Phospho-H2AX was seen as early as 15 min after DNA damage and through 12 h depending on radiation dosage. Analysis of phospho-H2AX for this study was completed using a Celomics® Phospho-H2AX Activation Kit (Thermo Scientific, Waltham, MA) and phospho-H2AX primary antibody and Goat anti-mouse fluorescently conjugated secondary antibody. Immunostaining was completed on a negative control of nontransferred fibroblasts, a positive control of non-laser transferred fibroblasts that had been exposed for 1 h to a 25 mg/mL concentration of etoposide (Sigma Aldrich) to induce single and double-strand DNA breaks, and four arrays of laser printed cells. Cells were fixed 3 h after laser transfer or seeding (non-laser transferred controls) to allow for cellular adherence to the growth surfaces. Hoechst dye was used to stain for cellular DNA, to identify the cell nucleus, and to verify colocalization of DNA and phospho-H2AX. Immunofluorescent images were taken on an inverted fluorescent microscope (Carl Zeiss).

Cell viability was tracked up to 24 h post-transfer and was determined by the ability of the cells to return to their normal adherent cell morphology from their trypsinized state. A total of 369 cells from 15 separate laser transfers were analyzed for viability.

Gelatin surface analysis

To characterize the topography of the receiving dishes, they were imaged using an MFP-3D atomic force microscope (AFM; Asylum Research, Santa Barbara, CA). A receiving dish was coated with PLL and filled with the cell culture
medium as a negative control. A second dish was coated in PLL, then spin coated with gelatin, filled with 4°C cell culture medium, and stored at 4°C as a positive control, and the experimental dishes were prepared as described in the receiving dish preparation, but no cells were transferred. The dishes were incubated for 15 min post-transfer, and filled with 10 mL of the warmed cell culture medium as previously prescribed, and returned to the incubator for 15 min before AFM imaging. Additional time points of 1 and 12 h of incubation were also imaged. All dishes were rinsed with ultra-pure cell culture water at 4°C before AFM imaging to remove potential artifacts generated by the proteins in the cell culture medium, then dried in a laminar flow hood, cut into sections, and mounted on glass slides. Imaging was completed in AC mode with a 50-μm scan size.

**Cell pattern registry following laser direct-write**

Images of the patterned cells and fiduciary mark in the receiving dishes were taken immediately after transfer (0 min), at 15 min, 30 min, and 1 h after laser direct-write. A semiautomated MATLAB® code was developed to track the trypsinized cell centroids in relation to the stationary micromachined fiduciary mark. The distance from cell centroid to fiduciary mark was calculated for each time point, then compared to the initial (0 min) time point. The absolute value of the change in distance between 0–15 min, 0–30 min, and 0–60 min indicates the distance a cell moved from its original patterned location. A total of 437 cells, from 15 laser transfers into seven separate receiving dishes, were analyzed for registry to the initial pattern. All data were expressed as the mean ± standard deviation of the mean. Because no outliers were present, comparisons were made between time points using two-tailed, heteroscedastic unpaired Student's t-test parametric analyses, with a level of \( p < 0.05 \) to establish statistical significance.

**Cell morphology and ECM production analysis**

Immunocytochemistry was performed to investigate if either the gelatin surface treatment or the laser direct-write process affects the cells' longer-term functionality (e.g., morphology and ECM production) in culture. Cells were laser direct-written into two transfer arrays (\( n = 2 \) separate receiving dishes) with 600-μm spacing between transfer spots, and cultured for 8 days. Additional cultures were carried out on fibroblasts pipetted into a PLL-coated dish and a PLL- and gelatin-coated dish (\( n = 1 \) for each) prepared as prescribed for a receiving dish to serve as nonlaser transferred controls. The cell culture medium was exchanged every 3 days. After 8 days in culture, cells were fixed in 4% paraformaldehyde and treated using standard immunocytochemistry protocols and anti-human antibody: anti-40k to observe fibronectin. Cell nuclei were observed with Hoechst dye and F-actin stress fibers with phalloidin (Invitrogen, Carlsbad, CA). Images were taken on an inverted fluorescent microscope (Carl Zeiss).

**Results**

Human dermal fibroblast cells were successfully laser patterned using gelatin. Independent ribbon control allowed the user to target specific spots on the ribbon for transfer and the user-generated g-code enabled customized arrays of cells to be patterned (Fig. 2).

**DNA damage analysis and cell viability**

Arrays of laser-transferred cells demonstrated no colocalization of Hoechst dye for cellular DNA and the immunostain for phospho-H2AX, indicating the absence of double-strand DNA breaks. A negative control showed only Hoechst dye for DNA and a positive control showed the colocalization of Hoechst dye with phospho-H2AX (Fig. 3).

Viability was determined by the ability of a cell to exhibit its normal adherent morphology from its trypsinized state after laser transfer. At 24 h after laser direct-write, a total of 13 cells (\( n = 369 \) cells from 15 separate laser transfers) did not exhibit normal adherent morphology and were deemed nonviable. Overall, fibroblasts maintained an average viability of 91% (±3% standard error of the mean) post-transfer at 24 h after laser direct-write.

---

**FIG. 2.** Representative time course of an array of human dermal fibroblasts after gelatin-based laser direct-writing, illustrating that **(a)** immediately after patterning, fibroblasts remain in their trypsinized morphology, **(b)** at 30 min after transfer some cells begin to spread and attach, **(c)** at 45 min after transfer, the fibroblasts display normal adherent morphology, and **(d)** at 24 h post-transfer, the cells display their normal spindle-like adherent morphology (2×3 array, 500-μm spacing, transfer spot size ~300 μm).
**Time course of gelatin removal**

AFM analysis of surface topography indicated that the 10% gelatin coating on the receiving dish changed with respect to incubation time. Once incubated, the gelatin-layer was disrupted and surface droplet formation was observed at 15 min, an indication of gelatin melting. After 1 h of incubation, surface topography of the receiving dish was similar to the negative control (gelatin-free, PLL-only). After 12 h of incubation, there were no further changes in surface topography compared to either the PLL-control or 1-h incubated dish, suggesting that gelatin melting and removal occurred within 1 h of incubation (Fig. 4).

**Cell pattern registry following laser direct-write**

Cell tracking analysis after transfer demonstrated a high degree of pattern registry (Fig. 5). Cells were tracked immediately after transfer (0 min), and up to 60 min after transfer, which corresponds to the time course of gelatin melting and cellular attachment. After 15 min of incubation after laser transfer, cell centroid distances from the fiduciary mark differed from the initial time point by $4.3 \pm 3.0 \mu m$ (mean ± standard deviation of the mean). After 30 min of incubation, cell centroid distances differed by $5.0 \pm 3.4 \mu m$ from the initial time point; at 60 min the difference was $5.6 \pm 2.5 \mu m$. No significant differences were observed between any of the time points (Fig. 6).

A small number of cells (15.6% of the 437 cells transferred) rinsed away from the transfer spots in the initial cell culture medium application or during multiple movements from the incubator to the microscope for imaging. A total of 39 cells could not be tracked from the initial time point after 15 min of incubation; 18 additional cells could not be tracked at 30 min, and 11 cells at 60 min. A transfer efficiency of 84.4% was maintained for the 15 transfer arrays at 60 min.

**Cell morphology and fibronectin synthesis**

After laser direct-write and subsequent culture for 8 days, cell morphology appeared no different from the nonlaser direct-written controls. All conditions (laser direct-written cells, and cells pipetted on controls consisting of both a PLL-only dish and a PLL-gelatin receiving dish) presented elongated and spread cell morphology with pronounced F-actin stress fibers (Fig. 7). Long-term cell functionality was demonstrated with antibody staining for fibronectin, which showed similar structure and amounts of ECM protein synthesis across all conditions (Fig. 7).

**Discussion**

Laser direct-writing is a powerful tool for the precise patterning of cells to investigate cell-to-cell interactions and responses with respect to spatial cues, and for rapidly building complex cellular constructs. However, the reliance
FIG. 4. Atomic force microscopic images illustrating the transient response of gelatin-coated growth surfaces with incubation duration. (a) PLL-only dish, prepared without gelatin to serve as a (-) control; (b–e) PLL-coated receiving dishes spin coated with 10% gelatin, and incubated for (b) 0 min (+) control, (c) 15 min, (d) 1 h, and (e) 12 h. Within 15 min of incubation, gelatin melting was observed, and at 1 h of incubation a similar surface topography to the negative control was seen, with no further changes observed at 12 h, suggesting an unobstructed growth surface with gelatin melting and removal within 1 h. PLL, poly-L-lysine.

FIG. 5. Representative image of indexed centroids of laser patterned trypsinized cells and the fiduciary marker (a) immediately after laser direct-write (0 min), and (b) after 30 min of incubation after laser direct-write. At 30 min the trypsinized cells begin to spread out and display their normal adherent morphology.
of this technique on Matrigel, used on the print ribbon and/or receiving substrate, introduces growth factors and ECM constituents that can influence cellular growth and differentiation post-transfer, which precludes or confounds its application, particularly with sensitive cell types. Further, Matrigel limits the potential of laser direct-writing for in vivo human tissue engineering applications in that Matrigel will not receive FDA approval in the foreseeable future, due to its human tissue engineering applications in that Matrigel will not receive FDA approval in the foreseeable future, due to its mouse Engelbreth-Holm-Swarm tumor origin and immunogenicity.35,36

To overcome these limitations, this study detailed a new gelatin-based laser direct-write method, developed to precisely pattern viable mammalian cells, in a completely Matrigel-free manner, using human dermal fibroblasts. This technique maintained high cell viability after transfer, and exhibited no evidence of double-strand DNA breaks, thereby demonstrating the nondetrimental nature of the laser-transfer process to the cells. Further, this method showed a high degree of cell registry on the receiving substrate. Registry to the initial pattern after 1 h of incubation was maintained to within the diameter of a cell (5.6 ± 2.5 µm), despite the melting and removal of the temporary gelatin layer from the growth surface.

This melting and subsequent removal of the gelatin from the receiving substrate can be utilized to provide a homogeneous growth surface that allows for normal cellular interactions and growth, uninhibited by surface treatments. This ability to pattern cells on a uniform surface is unique to laser printing as compared to other 2-D patterning techniques, such as photolithography37 or micropatterning with stencils38 that rely on specific cellular adhesion sites that may restrict normal cellular growth and interactions. On the receiving dish, the gelatin coating melted at 37°C and was removed in the course of 1 h of incubation, leaving the desired growth surface (e.g., tissue culture plastic, PLL-coated culture plastic, user- and application-specific culture surfaces, or other ECM protein coatings), further ensuring that the function of gelatin is only transient, and thus greatly reducing any unintended effect it may have on normal cellular proliferation.

In the original Matrigel-based MAPLE DW technique, to load the print ribbon with cells, the trypsinized cells must begin to form initial attachments through cellular adhesion to the Matrigel coating on the ribbon. Thus, the cells are no longer fully trypsinized, and the focal adhesions that adhere the cells to the print ribbon may be disrupted or damaged during the transfer process. Rather than allowing the cells to spread and loosely attach to the ribbon, we exploit the slow melting of gelatin to partially encapsulate the trypsinized cells in the print ribbon’s gelatin coating. This partial encapsulation of cells on the ribbon represents a fundamental change from the original MAPLE DW technique. Since the trypsinized state of the cells is maintained during printing, the cells are balled-up rather than spread out and attached, and no focal adhesions will be disrupted or damaged during laser transfer. Further, this technique can be applied to virtually any type of cell. Thus, single heterogeneous ribbons of multiple cell types can be prepared, thereby making possible the rapid printing of multiple cell types.

An unexpected advantage of the gelatin for the print ribbon was its optical clarity, which allowed for observation of both the target location on the receiving dish as well as the cells on the print ribbon. Additionally, the firm cell–ribbon attachment enables the targeting of specific cells for transfer, which cannot be achieved with other biological laser printing techniques in which cells are suspended and free floating on the print ribbon in a combination of cell culture medium and glycerol.10,11,14,15 The coupling of optical clarity and firm cell attachment on the ribbon provides unprecedented user control and selective cell targeting.

Cell centroid location tracking showed that trypsinized fibroblast cells remained in their specified locations even as the fresh growth medium was added following laser direct-write and the gelatin melted. The high degree of registry (5.6 ± 2.5 µm at 60 min post-transfer) to the initial pattern demonstrates the precision necessary to fabricate cellular constructs for in-depth investigations of cellular behavior and interactions. Although labor intensive, in situ cell tracking through 24 h post-transfer avoided having to fix and stain cells, which would have offered only representative temporal snapshots. Similarly, the in situ viability analysis, used to verify the passivity of the gelatin-based MAPLE DW transfer process, granted continuous observation and analysis over 24 h. This provides a conservative measure of cell viability, and avoids the use of live–dead staining, which can overestimate the viability, as dead cells may inadvertently be rinsed away while live cells remain adherent.

Phospho-H2AX allowed DNA damage analysis to be conducted on transfer conditions that would exactly match transfer conditions used for virtually any other laser-based cell patterning application. Although comet assays can quantify the amount of DNA damage, they require gel electrophoresis, which greatly limits, if not precludes, the investigation of DNA damage while maintaining the intended cellular pattern and transfer conditions. The ability of phospho-H2AX immunochemistry to assay cells while in
their pattern arrays enables the analysis of DNA damage without compromising or sacrificing the structural information of the culture. Future studies can therefore investigate the spatial distribution of DNA damage.

Long-term cell functionality was verified with antibody staining for the synthesis of the ECM protein, fibronectin, in which laser-transferred cells showed no difference from pipetted (nonlaser transfer) controls. Within these controls, cells grown on receiving dishes with 10\% gelatin coatings exhibited similar cell morphology and long-term cellular protein synthesis as those without gelatin. These findings suggest that neither the laser transfer process nor the 10\% gelatin coating on the receiving dish had a negative influence on the cells' behavior in culture, thereby indicating that the long-term functionality of the cells is maintained in gelatin-based laser direct-writing.

The results presented herein demonstrate that gelatin-based MAPLE DW is an improvement over previous materials used for conducting in-depth investigations of cellular interactions based on spatial proximity, composition, and geometric location that are devoid of the influence of extraneous ECM proteins or growth factors. Our future applications of gelatin-based laser direct-writing will investigate the spatial dependence of cells in culture on ECM protein production, as well as the microenvironmental factors that affect stem cell differentiation. The ability to pattern cells and have control over growth substrate constituents without introducing extraneous growth factors enables future studies involving sensitive cell types, and for investigating the effect of spatial patterning on cellular growth factor secretion and cellular protein production. Gelatin-based laser direct-writing is a powerful approach in terms of precision, customization, and reproducibility. Through this relatively simple change to the MAPLE DW process, the utility of laser cell patterning has expanded, and laser direct-writing can play a more definitive role in future investigations in basic cell biology, as well as applications in tissue engineering, cancer, and regenerative medicine research.

Acknowledgments
The authors would like to thank Chris Bjornsson, Ph.D., and Gaurav Jain, M.S., of Rensselaer Polytechnic Institute for their assistance with AFM imaging, as well as Livingston Van de Water, Ph.D., Courtney Betts, M.S., and Paula McKeown-Longo, Ph.D., of Albany Medical College for their...
assistance and generous gifts of fibronectin antibody (anti-40k), phalloidin, and Hoechst dye. This study has been partially supported by the National Institutes of Health (1R56DK088217-01) and internal start-up funds at Rensselaer Polytechnic Institute.

Disclosure Statement

No competing financial interests exist.

References


Address correspondence to:

David T. Corr, Ph.D.
Department of Biomedical Engineering
Rensselaer Polytechnic Institute
Jonsson Engineering Center, Rm. 7042
110 8th Street
Troy, NY 12180-3590

E-mail: corrd@rpi.edu

Received: July 28, 2010
Accepted: September 17, 2010
Online Publication Date: October 22, 2010