

Video Article

# Electrospinning Fibrous Polymer Scaffolds for Tissue Engineering and Cell Culture

Jamie L. Ifkovits<sup>1</sup>, Harini G. Sundararaghavan<sup>1</sup>, Jason A. Burdick<sup>1</sup>

<sup>1</sup>Department of Bioengineering, University of Pennsylvania

Correspondence to: Jason A. Burdick at [burdick@seas.upenn.edu](mailto:burdick@seas.upenn.edu)

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## Abstract

As the field of tissue engineering evolves, there is a tremendous demand to produce more suitable materials and processing techniques in order to address the requirements (e.g., mechanics and vascularity) of more intricate organs and tissues. Electrospinning is a popular technique to create fibrous scaffolds that mimic the architecture and size scale of the native extracellular matrix. These fibrous scaffolds are also useful as cell culture substrates since the fibers can be used to direct cellular behavior, including stem cell differentiation (see extensive reviews by Mauck *et al.* and Sill *et al.* for more information). In this article, we describe the general process of electrospinning polymers and as an example, electrospin a reactive hyaluronic acid capable of crosslinking with light exposure (see Ifkovits *et al.* for a review on photocrosslinkable materials). We also introduce further processing capabilities such as photopatterning and multi-polymer scaffold formation. Photopatterning can be used to create scaffolds with channels and multi-scale porosity to increase cellular infiltration and tissue distribution. Multi-polymer scaffolds are useful to better tune the properties (mechanics and degradation) of a scaffold, including tailored porosity for cellular infiltration. Furthermore, these techniques can be extended to include a wide array of polymers and reactive macromers to create complex scaffolds that provide the cues necessary for the development of successful tissue engineered constructs.

## Video Link

The video component of this article can be found at <https://www.jove.com/video/1589/>

## Protocol

### A. Single Polymer Electrospinning

1. Prior to preparing the electrospinning solution, make a 0.5 wt% solution of the photoinitiator, Irgacure 2959 (I2959), in deionized water by dissolving at 37°C for several days. This step is not necessary if a photoreactive polymer is not being used.
2. Combine methacrylated hyaluronic acid (MeHA, see Burdick *et al.* for synthesis), poly(ethylene oxide) (PEO, 900 kDa), and I2959 in deionized water to prepare a solution with a final concentration of 2 wt% MeHA, 3 wt% PEO, and 0.05 wt% I2959. Use a vortex to mix the solution until it is clear. The polymer type and concentration, as well as the solvent used can be modified at this step depending on the desired scaffold properties.
3. Transfer the solution into a syringe and attach an 18 gauge, 6 inch long, blunt end needle to the end.
4. Program a syringe pump to eject at a rate of 1.2 mL/hr. Insert the syringe and needle into the device.
5. Attach the grounded lead of a high voltage power source to the collection apparatus. To collect randomly distributed fibers use a flat metal plate, or use a mandrel rotating at ~10 m/s to collect aligned fibers. Attach the positively charged lead to the needle. See **Figure 1** for a schematic of the electrospinning apparatus.
6. Adjust the needle or collecting device, such that there is a 15 cm distance between the two.
7. Start the flow on the syringe pump. When fluid is visualized on the tip of the needle, turn on the power source and set the voltage to 22 kV.
8. After the collection is complete (from several minutes for thin films, to up to 24 hours for a thicker mat), remove the scaffold from the collection apparatus and store it under vacuum overnight to ensure complete removal of the solvent.
9. Visualize the fiber morphology using scanning electron microscopy (SEM), a representative scaffold is shown in **Figure 1** for both aligned and non-aligned structures.

Note: The sample flow rate, distance to collection device, and voltage are dependent on the polymer and solvent combination and must be optimized for each system, usually by observing the scaffold morphology with SEM.

### B. Photocrosslinking and Photopatterning

1. Cut out 5mm X 5mm samples from the scaffold mat.

2. Prepare for crosslinking by placing each scaffold on a foil covered glass slide, placing the photomask directly on the scaffold, covering with a clean glass slide and clipping both ends with binder clips. Note: four samples can be done at once, and a transparency without a pattern can be used for crosslinking if a pattern is not desired.
3. Purge scaffold set up in a nitrogen chamber. It is important to keep the construct free from oxygen, which can inhibit crosslinking.
4. Place scaffold setup in nitrogen chamber under  $\sim 10 \text{ mW/cm}^2$  365 nm light with a collimating adapter for 5 minutes. See schematic in **Figure 2**.
5. Remove each scaffold and place in a 12 well plate.
6. Add 2 mL of deionized water into each well, parafilm the plate to prevent evaporation of the water and place at 37°C for 24 hours. Change the water three times.
7. Visualize pore formation under light microscope (see example in **Figure 2**). The scaffolds are now ready for use.

Note: Photocrosslinking is not necessary for many types of polymers, however photopatterning can only be used with photoreactive polymers.

## C. Dual Polymer Electrospinning with Fluorescent Fiber Visualization

1. Prepare a 5 wt% solution of PEO (200 kDa) in 90% ethanol. Stir at 700 rpm at 50°C for at least 2 hours prior to electrospinning.
2. Transfer the PEO solution to a syringe and add DAPI for a final concentration of 10 mg/mL concentration in the electrospinning solution. Wrap the syringe in aluminum foil to protect it from light.
3. Prepare the same solution described in step A2, except also add methacryloxyethyl thiocarbamoyl rhodamine B (MeRho, 683.24 g/mol) for a final concentration of 25  $\mu\text{M}$  in the electrospinning solution. Wrap the syringe in aluminum foil to protect it from light.
4. Carefully use scotch tape to secure methacrylated glass coverslips (see Khademhosseini *et al.*) to the surface of the mandrel. Note: fibers will become tethered to methacrylated glass coverslips.
5. Attach one end of a 5 mm long piece of silicone tubing with luer lock attachments to one of the syringes and attach the other end to a needle. Insert the syringe into the syringe pump and put the needle through the hole in the fanner. Repeat with the other syringe and fanner on the opposite site of the mandrel. The fanners translate the length of the mandrel and are used to ensure an equal distribution of both polymers in the resulting scaffold. See schematic in **Figure 3**.
6. Adjust the parameters described in Section A appropriately according to **Table 1**. Ensure that the needle tips are centered on the mandrel.

**Table 1. Dual Polymer Electrospinning Parameters**

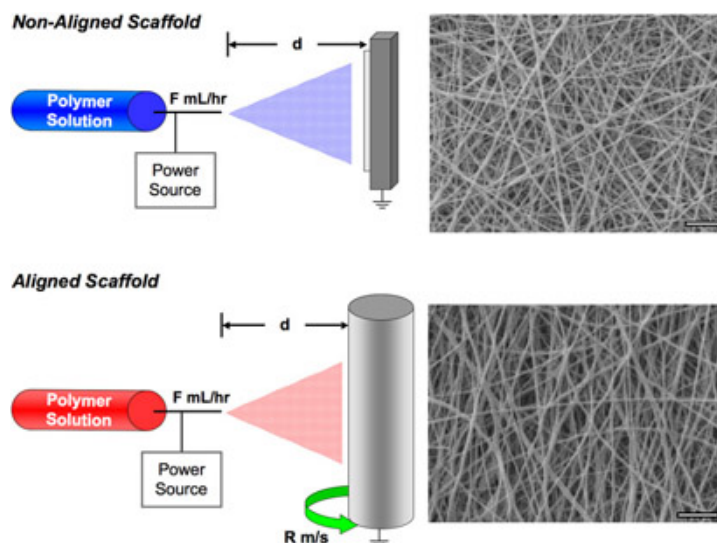
Syringe	Fanner to Needle Tip (cm)	Needle Tip to Mandrel (cm)	Flow Rate (mL/hr)	Applied Voltage (kV)
MeHA	6	15	1.2	+22
PEO	6	10	1.2	+15

7. Once everything is aligned properly, turn off the lights and turn on the mandrel and the syringe pumps. Remove the aluminum foil from the syringes. When fluid is visible on the tips of both needles, simultaneously turn on the power supplies and plug in the fanners.
8. When collection is complete, turn off the power supplies and mandrel and unplug the fanners. Carefully remove the coverslips and tape using a razor blade.
9. Visualize the fibers using a fluorescent microscope equipped with filters for rhodamine and DAPI. See example in **Figure 3**. Note: Fluorescent fibers are most clear to view when electrospun for short periods of time (<3 minutes), however, this process can be extended for an unlimited amount of time in order to create thicker constructs, especially without the use of the glass coverslips.

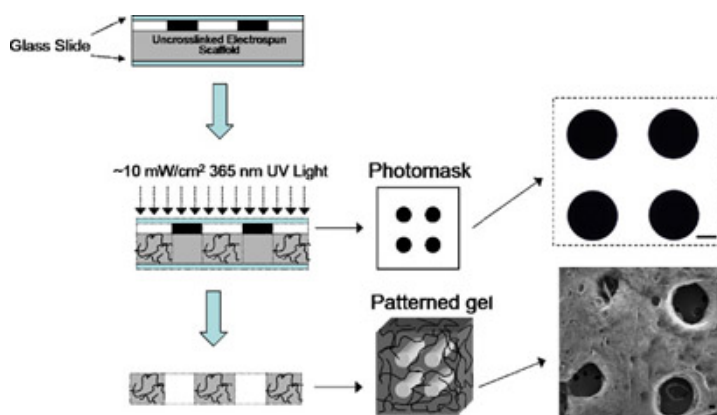
## D. Seeding Cells on Scaffolds

1. Place each coverslip with electrospun polymer attached into an individual well of an appropriately sized well plate. Note: cellular interactions with the fibers are easily visible by electrospinning onto methacrylated glass coverslips. Also, MeRho can again be used for fiber visualization.
2. Incubate the scaffolds in PBS overnight to ensure complete removal of solvent and any possible leachable by-products.
3. Remove the PBS from the wells.
4. To sterilize the scaffolds, place them under a germicidal lamp in a laminar flow hood for 30 minutes. If a full scaffold is being seeded, flip the scaffold over and place under a germicidal lamp for another 30 minutes.
5. Perform standard cell culture and prepare a concentrated cell suspension at the desired cell density (e.g., 6,000 cells  $\text{cm}^{-2}$ ). For example, use a 100 mL of cell suspension for a 22 mm X 22 mm coverslip. Place in the incubator for 1 hour.
6. Add the appropriate amount of cell culture media to each well. Place the scaffolds back into the incubator.
7. Stain the cells using a commercially available Live/Dead kit. Note: other types of cell staining, such as DAPI (cell nuclei) and fluorescently labeled phalloidin (actin stress fibers) can be applied to visualize the cells.
8. Visualize the cells and fibers using a fluorescent microscope equipped with filters for TRITC and FITC. See example in **Figure 4**.

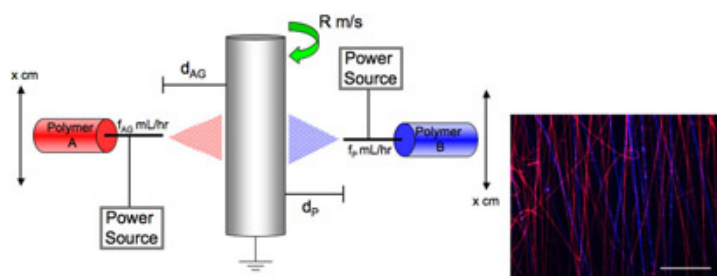
## Representative Results:



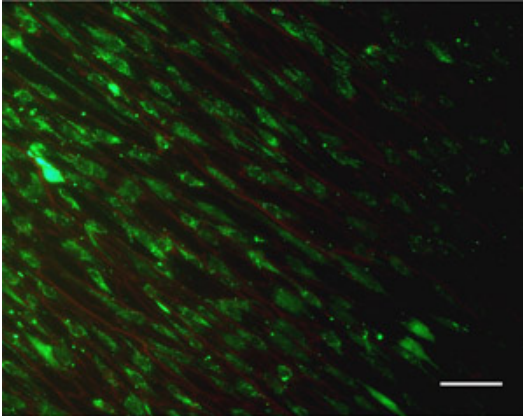
**Figure 1.** Schematic illustrating the device setup for non-aligned scaffold formation (top) and aligned scaffold formation (bottom). Example scanning electron microscopy images of each type of scaffold are shown. Scale bar =  $5\mu\text{m}$ . Please [click here](#) to see a larger version of figure 1.



**Figure 2.** Schematic of patterning method. Patterns are formed by placing a photomask between the light source and scaffold during photocrosslinking and then washing away unreacted polymer. Photomask and SEM image of scaffolds after pore formation and lyophilization. Scale bar =  $100\mu\text{m}$ . Please [click here](#) to see a larger version of figure 2.



**Figure 3.** Schematic of the multi-polymer electrospinning setup and necessary processing parameters, as well as a representative fluorescent image of a mixture of two fiber populations (example: red MeHA and blue PEO) which were simultaneously electrospun to form a multi-polymer scaffold. Scale bar =  $100\mu\text{m}$ . Please [click here](#) to see a larger version of figure 3.



**Figure 4.** An example Live/Dead staining image of human mesenchymal stem cells and their interactions with the electrospun fibers. Scale bar = 100  $\mu$ m. Please [click here](#) to see a larger version of figure 4.

## Discussion

Electrospinning was used to prepare fibrous scaffolds from polymers. Photocrosslinkable scaffolds based on hyaluronic acid were used as an illustrative example, where light exposure is needed for crosslinking. With the use of reactive macromers, such as MeHA, channels that have previously demonstrated enhanced cellular distribution were incorporated into the scaffolds with the use of a mask during photocrosslinking to form macro and micro-porous scaffolds. Moreover, two distinct polymers were simultaneously electrospun using our custom apparatus. The presence of the two distinct fiber populations was verified with the addition of a fluorescent dye to each electrospinning solution, which was later viewed using a fluorescent microscope. Multi-polymer scaffolds can be used to enhance cellular distribution, as well as better tune the mechanics and degradation of a scaffold for a particular application relative to single polymer electrospun scaffolds. Furthermore, sterilization and cell seeding of the scaffolds has been demonstrated. Many of these techniques are versatile and can be used for a range of different polymers to fabricate diverse scaffolds for applications in tissue engineering and for the culture of cells.

## Acknowledgements

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