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Jingwei Xie
*Marshall University, xiej@marshall.edu*

Matthew R. MacEwan

Xiaoran Li

Shelly E. Sakiyama-Elbert

Younan Xia

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Neurite outgrowth on nanofiber scaffolds

Chemical methods for nanofabrication

From golden cages to golden pyramids
Neurite Outgrowth on Nanofiber Scaffolds with Different Orders, Structures, and Surface Properties

Jingwei Xie, Matthew R. MacEwan, Xiaoran Li, Shelly E. Sakiyama-Elbert, and Younan Xia*

Department of Biomedical Engineering, Washington University, St. Louis, Missouri 63130

Structural and biomolecular patterns play an important role in the embryonic development of the nervous system. In building the intricate neural networks, axons must be precisely guided to the synaptic targets and various cell populations have to be spatially distributed into a specific pattern. The effectiveness of these two processes critically depends on the presence of patterned cues to guide neurite outgrowth. The cues can typically be divided into two main categories: chemical cues based on neurite attractive/repulsive molecules (e.g., netrins, slits, semaphorins, and ephrins),1 and physical cues that may include applied tension/stress, electrical polarization, magnetic field, and topography.2–5

The effects of topographic cues on neurite extension have been extensively investigated through the use of substrates containing microgrooves or microchannels of different depths and widths, typically generated using microlithography. Neurites have been demonstrated to grow parallel to a channel when the channel was 20–40 μm wide but perpendicular to a channel when the width increased to the range of 40–60 μm.6 Also, neurites have been aligned perpendicular to shallow grooves of 1 μm in width and hundreds of nanometers in depth.7 In addition, neonatal rat DRG neurons have been cultured on poly(dimethyl siloxane) (PDMS) substrates patterned with grooves and coated with poly-L-lysine (PLL) and laminin, and it was shown that the neurites could extend across several adjacent grooves.8 Another recent study suggested that topography might play a critical role in axon formation and that trophic support, such as immobilized nerve growth factor (NGF), could enhance axon growth only after initiation.9 Although soft lithographic techniques allow for the fabrication of topographical features with precise dimensions for investigating the role of topography in axonal growth at the micro- and nanoscale, PDMS has little value for translational research as it is not biodegradable.9

Electrospinning is an effective and widely utilized method for producing continuous fine fibers from polymer solutions or melts for a variety of applications.10–12 So far this technique has been successfully applied to more than 100 types of natural and synthetic polymers.12 Due to their small diameters and large surface areas, electrospun nanofibers can be employed to mimic the extracellular matrix (ECM) for cell attachment and nutrient transportation and have been intensively investigated as scaffolds for various tissue engineering applications.10–12

ABSTRACT  Electrospun nanofibers can be readily assembled into various types of scaffolds for applications in neural tissue engineering. The objective of this study is to examine and understand the unique patterns of neurite outgrowth from primary dorsal root ganglia (DRG) cultured on scaffolds of electrospun nanofibers having different orders, structures, and surface properties. We found that the neurites extended radially outward from the DRG main body without specific directionality when cultured on a nonwoven mat of randomly oriented nanofibers. In contrast, the neurites preferentially extended along the long axis of fiber when cultured on a parallel array of aligned nanofibers. When seeded at the border between regions of aligned and random nanofibers, the same DRG simultaneously expressed aligned and random neurite fields in response to the underlying nanofibers. When cultured on a double-layered scaffold where the nanofibers in each layer were aligned along a different direction, the neurites were found to be dependent on the fiber density in both layers. This biaxial pattern clearly demonstrates that neurite outgrowth can be influenced by nanofibers in different layers of a scaffold, rather than the topmost layer only. Taken together, these results will provide valuable information pertaining to the design of nanofiber scaffolds for neuroregenerative applications, as well as the effects of topology on neurite outgrowth, growth cone guidance, and axonal regeneration.

KEYWORDS: electrospun nanofibers · patterning · coating · neurite outgrowth · guidance

*Address correspondence to xia@biomed.wustl.edu.

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applications. In particular, aligned nanofibers are well-suited for neural tissue engineering as the anisotropic properties of aligned nanofibers may provide spatial guidance for neurite outgrowth and axonal elongation in vitro. For example, previous studies have shown that aligned nanofibers were better suited for culturing neural stem cells in vitro than scaffolds consisting of randomly oriented nanofibers. Other studies have also demonstrated that aligned nanofibers were able to direct neurite extension from cultured DRG and guide axonal growth or glia migration. In addition, one prior study showed that neurite outgrowth was significantly increased on aligned nanofibers immobilized with laminin relative to untreated samples. In all these studies, however, neurite extension was only examined for single-layered scaffolds that only contained either aligned or randomly oriented nanofibers. Given the intricate structure of a neural network, it will be interesting and significant to investigate neurite extension and axonal regeneration on more complex scaffolds, such as single-layered scaffolds containing both aligned and disordered fibers with an interface between them and bilayered scaffolds composed of nanofibers with different orders or orientations.

In the past, we and other groups have modified electrospinning to provide a simple and versatile method for generating two- and three-dimensional assemblies of nanofibers with well-defined and controllable orders, structures, and surface properties. In the present study, we cultured embryonic chick DRG on some of these assemblies consisting of nanofibers electrospun from poly(e-caprolactone) (PCL), a biocompatible and biodegradable polymer, and then examined the outgrowth of neurites in vitro. These studies not only provide a better understanding of neurite outgrowth, growth cone guidance, and axonal regeneration on nanofiber scaffolds but also offer valuable information with regard to the design of new scaffolds for neuroregenerative applications.

RESULTS AND DISCUSSION

In our previous studies, we were able to generate uniaxially aligned fibers across the void gap of a metal frame and multilayered fiber mats with controllable hierarchical structures by layer-by-layer stacking. In the present work, scaffolds made of randomly oriented nanofibers were fabricated by directly depositing the electrospun fibers onto glass coverslips. Figure 1A shows a scanning electron microscopy (SEM) image of a typical sample of randomly oriented PCL nanofibers. Figure 1B shows the Fourier fast transfer (FFT) analysis, indicating that the nanofibers were randomly oriented because the pixel intensities (labeled with light blue color) were independent of direction. We then cultured DRG on the random PCL nanofibers and found that DRG adhered poorly to them and could easily fall off the scaffold due to washing during the process of immunostaining. Only a few of the DRG could adhere to random PCL nanofibers and grow.
contrast, all of the DRG adhered well to the random PCL nanofibers after coating with laminin. Hence, laminin coating could significantly promote adhesion of DRG to random PCL nanofiber scaffolds. Panels C and D of Figure 1 show fluorescence micrographs of the typical morphology of DRG after they had been seeded on bare and laminin-coated, random PCL nanofibers and then cultured for 6 days. The neurites grew radially outward from the main body without preference to any specific direction, exhibiting a circular appearance. This observation is consistent with the results of previous studies. In addition, it seems that DRG on bare and laminin-coated random PCL nanofibers exhibited a similar neurite field in terms of profile and neurite length.

Using an approach similar to what we used in our previous studies, aligned PCL nanofibers were collected on a metal frame and then transferred onto a glass coverslip by lifting them up from the underneath. Figure 2A shows SEM image of a typical sample of aligned PCL nanofibers, and the inset shows an FFT pattern, suggesting that the fibers were uniaxially aligned. When DRG were cultured on the aligned PCL nanofibers, we found that the DRG adhered well to the scaffold even without laminin coating and neurites preferred to grow along the long axis of the fiber (Figure 2B). The fluorescence micrograph in Figure 2B also indicates that some of the neurites initially did not grow along the direction of fiber alignment but could turn their growth directions and eventually grew parallel to the fiber alignment. Similarly, DRG adhered well to aligned PCL nanofibers whose surface had been coated with laminin, and the neurite outgrowth followed the fiber alignment direction (Figure 2C). We also found that some of the neurites could dramatically change their growth directions (Figure 2D), showing even sharper turns than on the aligned fibers without a laminin coating. This observation indicates that the laminin coating could greatly enhance the guidance of neurite outgrowth by the underlying nanofibers.

Similar to our previous study, we quantified the average neurite length, the maximum neurite length, and eccentricity of the neurite field using MATLAB (Figure 3). There was no significant difference for the neurite length and eccentricity between bare and laminin-coated samples for random fibers. However, the average length of the neurites increased from 857 μm for laminin-coated, random fibers to 1085 μm for aligned, bare fibers, and further to 1542 μm for aligned, laminin-coated fibers (Figure 3A). The maximum length of neurites projecting from DRG cultured on aligned nanofibers was longer, approximate 2-fold greater, than that of neurites projecting from DRG cultured on random nanofibers (Figure 3B). Laminin-coated, aligned fibers further enhanced the maximum neurite length: approximately 500 μm longer than the bare, aligned nanofibers. Accordingly, the value of eccentricity increased from 0.54 for laminin-coated, random fibers to 0.82 for aligned, bare fibers, while it was 0.89 for laminin-coated aligned fibers (Figure 3C). These results clearly demonstrate that aligned nanofibers could promote DRG adhesion and enhance the neurite guidance and extension as compared to random fibers. Laminin coating could further enhance the neurite extension for aligned fibers.

By making use of the collector composed of two metal strips separated by an air gap, we were able to fabricate scaffolds containing both aligned and random nanofibers. Figure 4A shows SEM image of a typical sample where the nanofibers had a clear transition from aligned to random orientation. In this case, the nanofibers were deposited on the metal strips as randomly oriented mats while they were uniaxially aligned across the gap. The insets show FFT patterns taken from these two regions, confirming the alignment and randomness of their fibers. Figure 4B shows a fluorescence micrograph of the typical morphology of DRG after they had been seeded on random PCL nanofibers and then cultured for 6 days. The neurites grew radially outward from the main body without preference to any specific direction, exhibiting a circular appearance. This observation is consistent with the results of previous studies. In addition, it seems that DRG on bare and laminin-coated random PCL nanofibers exhibited a similar neurite field in terms of profile and neurite length.
cence micrograph of the typical morphology of DRG cultured at the border between the random and aligned fibers. Interestingly, the neurites grew without any preference in orientation on the side with random fibers and grew along the fiber alignment on the side with aligned fibers. Overall, the DRG shows a “Janus feature”, with one side of the neurites uniaxially aligned and the other side radially pointing in different directions. Similar to cultures on uniformly aligned fibers, we observed enhancement of neurite extension and guidance of neurite outgrowth when the nanofibers were precoated with laminin before DRG culture (Figure 4C,D). Figure 5A shows SEM image of DRG cultured at the border between aligned and random fibers of a laminin-coated scaffold. It can be clearly seen that the neurites on one side of the DRG were randomly distributed on the disordered fibers (Figure 5B), while those on the other side grew preferentially along the fiber alignment direction (Figure 5C,D). Figure 5D also shows cell migration along the direction of fiber alignment.

Through layer-by-layer stacking, we were able to fabricate multilayered scaffolds of aligned fibers with a specific orientation in each layer. Figure 6 shows SEM images of typical double-layered meshes with low (distance between fibers \( \approx 5 \, \mu m \)), mediate (distance between fibers \( \approx 1 \, \mu m \)), and high (distance between fibers \( \approx 0 \)) densities of fibers where the nanofibers in different layers were rotated by approximately 90° from each other.

The FFT patterns shown in Figure 6A,B suggest that the mesh consisted of two layers of uniaxially aligned nanofibers with their orientation rotated by an angle of 80 and 90°, respectively. However, the FFT pattern in the inset of Figure 6C shows a pattern similar to that of uniaxially aligned nanofibers due to a large thickness for the top layer.

When DRG were cultured on the double-layered meshes, the DRG could adhere to all the scaffolds without the application of any laminin. Figure 7A shows a fluorescence micrograph illustrating the typical morphology of DRG cultured on the mesh consisting of bare PCL fibers with a low fiber density. Very few neurites were observed, and they tended to grow along the directions of the orthogonal fibers. Figure 7B shows a fluorescence micrograph of DRG cultured on the same kind of scaffold, but with laminin coating. The substrate was precoated with poly(ethylene glycol) (PEG) by physical adsorption before the nanofiber mesh was deposited. Compared to the sample without laminin coating, more and longer neurites were observed. More interestingly, we demonstrated for the first time that some of the neurites grew along the long axis of the fibers in one layer and suddenly make a sharp turn to follow the long axis of fibers in the other layer. This can be clearly seen in the inset of Figure 7B. Figure 7C shows the typical morphology of DRG cultured on PCL fiber mesh with a mediate fiber density. Many short neurites were observed, and some of the neurites grew along the long axis of underlying fibers and turned to

Figure 4. (A) SEM image of disorder-to-aligned fiber mat. The FFT patterns in the inset indicate that the fibers were aligned on one side and randomly oriented on the other side. (B) Typical morphology of DRG cultured at the border between random and aligned PCL nanofibers (bare). (C) Typical morphology of DRG cultured at the border between random and aligned PCL fibers coated with laminin. The dashed line indicates the borderline between aligned (right side) and randomly oriented (left side) fibers. (D) An enlarged view of (C).

Figure 5. (A) SEM images of DRG cultured at the interface between aligned and random PCL fibers with laminin coating. (B–D) Enlarged views of regions B–D indicated in image (A).
the direction of fiber alignment in the top layer. In contrast, more and longer neurites appeared after coating with laminin, and the neurites seemed to form an orthogonal pattern resembling the underlying nanofiber mesh (Figure 7D). For the samples with a high fiber density, fewer and shorter neurites were noticed on bare scaffolds as compared to the laminin-coated samples (Figure 7E,F). However, the neurites were still able to form an orthogonal pattern similar to the underlying nanofiber mesh. We found that there was no significant difference in average neurite length on laminin-coated (516 μm) and bare scaffolds (470 μm) (Figure 10A). In contrast, the maximum neurite length for A/R with laminin coating was significantly longer than the sample without coating (Figure 10B). Accordingly, the eccentricity for the laminin-coated sample (0.60) was greater than the bare sample (0.33) (Figure 10C). A similar observation was made when using aligned fibers with a high fiber density on the top layer and random fibers on the bottom layer (Figure 9A). The average neurite length was similar for samples with and without laminin coating (Figure 10A). The maximum neurite length was slightly longer for laminin-coated R/A samples as compared to bare R/A samples (Figure 10B). Accordingly, their eccentricities were 0.70 and 0.59, respectively (Figure 10C). A similar observation was made when using aligned fibers with a high fiber density on the top layer and random fibers in the bottom layer (Figure 9A). The role of the fibers in the bottom layer under certain circumstances.

Understanding neurite contact guidance is of critical importance for the design of synthetic nerve grafts. It is well-known that contact guidance of neurites can be exerted by topographic features. Also, contact guidance of neurite outgrowth has been extensively investigated using surface features patterned by techniques such as photolithography and microcontact printing. Although electrospun nanofibers have been examined for the guidance of neurite outgrowth, most of these studies were limited to the use of either random or aligned fibers. In the present work, we have moved one step forward by examining the effects of various nanofiber assemblies with complex structures on the neurite extension and outgrowth guidance.
Previous studies showed that neurite outgrowth could be guided along the direction of aligned electrospun nanofibers.\textsuperscript{15–17} Our results are in line with the observations of these reports. We also demonstrated that aligned nanofibers could enhance the neurite extension based on the analysis of maximum neurite length. We further established that neurites exhibited two different patterns on two opposite sides of DRG when cultured at the borderline between random and aligned nanofibers. Our study of neurite outgrowth on double-layered fiber meshes may provide valuable information to help us better understand the contact guidance of neurite outgrowth by topographic cues. For example, why do the neurites grow along the fiber alignment direction? During neurite outgrowth, neurite filopodia and lamellipodia which emerge from the growth cone probe the surrounding extracellular microenvironment for neural growth cone pathfinding, and these cytoskeleton structures rich in microtubules and actin filaments yield traction forces that push and pull the neurite forward.\textsuperscript{27,28} The direction of neurite extension is caused by the strength of the traction force exerted by the filopodia, and this force is determined by the extent to which these protein filaments can accumulate, assemble, and orient in the direction of a cell protrusion.\textsuperscript{6,29} In view of our results, we believe the mechanisms used by the aligned nanofibers to control the neurite orientation can be summarized as follows. When DRG were cultured on the surface of a scaffold of random nanofibers, the strength of the traction force exerted by the filopodia was uniform in all directions. As a result, neurites emerged from the DRG main body in all directions and exhibited a radial distribution surrounding DRG main body (Figure 1). When cultured on the surface of aligned nanofibers, the strength of the traction force exerted by the filopodia was not uniform in all directions any more due to the anisotropic properties of the aligned fibers. In this case, neurites may prefer to extend along the fibers instead of stepping across them (Figure 2B–D). One previous study, however, reported that microtubules and actin filaments within the cytoplasm are too rigid to allow considerable deformation of filopodia to accommodate alterations of topography.\textsuperscript{30} This conclusion is in agreement with our findings.
contradiction to what we have observed in the present work (Figure 6E).

During development, the ECM directs the axons of maturing neurons to their innervating targets through a combination of contact-mediated (e.g., laminin) and soluble factors such as neurotrophic factor NT-3. Lamnin, an ECM glycoprotein, promotes neurite outgrowth via multiple cell-adhesion sites. Binding of laminin to neuronal surface integrins is required for proper neural crest migration. Laminin also supports neurite outgrowth from cultured explants and modulates the guidance of growth cones in response to extracellular cues. The electrospun nanofiber itself can be functionalized by encapsulation or attachment of biochemical cues to improve its biomimetic capability. One recent study showed coupling laminin with electrospun poly(L-lactic acid) (PLLA) nanofibers could enhance the neurite extension of PC12 cells. In this study, we demonstrated that immobilization of laminin on the surface of PCL nanofibers through electrostatic interaction between poly-L-lysine and laminin could enhance neurite guidance and extension. PEG coating on the substrate might be beneficial to the neurite guidance by nanofibers in that PEG is one of the best-known nonfouling biomaterials with extremely low energy and non-adhesive surfaces. We also found that neurites could make a sharp turn upon encountering the fibers in another layer with a different orientation when DRG were cultured on a double-layered nanofiber scaffold (with laminin coating and low fiber density) supported on a PEG-coated polystyrene substrate. Combined together, these results clearly suggest that electrospun nanofibers can serve as a class of versatile scaffolds for controlling the fabrication of neural networks.

CONCLUSION

We have demonstrated that electrospinning can generate a variety of nanofiber assemblies, which can serve as a new platform for investigating the outgrowth of neurites. When DRG were cultured on nanofiber scaffolds, the neurites were evenly distributed on scaffolds consisting of random fibers and grew preferentially along the fibers on uniaxially aligned samples. When seeded at the border between aligned and random fibers, the neurites originating from the same DRG could simultaneously grow into both aligned and random structures. Moreover, by stacking electrospun nanofibers into double-layered meshes, the neurites could be guided to grow into complex patterns, and the nanofibers in both layers could provide the guidance. For all scaffolds consisting of PCL nanofibers, the effectiveness of guidance could be further improved by coating the surface with laminin. Taken together, these results could contribute to a better design of new scaffolds for nerve repair and lead to a more thorough understanding of neurite outgrowth behavior on electrospun nanofibers.

EXPERIMENTAL METHODS

Fabrication and Characterization of Nanofiber Assemblies. The nanofibers were produced by electrospinning, and the setup was similar to what we used in previous studies. Poly(e-caprolactone) (PCL) (Mn = 65,000 g/mol; Sigma-Aldrich, St. Louis, MO) was dissolved in a solvent mixture consisting of dichloromethane (DCM) and N,N-dimethylformamide (DMF) (Fisher Chemical, Waltham, MA) with a ratio of 8.2 (v/v) at a concentration of 20% (w/v). Polymer solution was pumped at a flow rate of 0.5 mL/h using a syringe pump. A dc high voltage of 12 kV was applied between the nozzle (a 22-gauge needle) and a grounded collector. Different collectors were used to generate different types of nanofiber assemblies. Random nanofibers were directly collected using cover glass slips. A stainless steel frame (with an open void of 2 cm × 5 cm) was used as the collector. Subsequently, the aligned nanofibers were easily transferred to the cover glass slips by lifting off the fibers. Samples containing both random and aligned fibers next to each other were obtained by using two metal frames separated by an air gap. Fibers were deposited in the random and aligned form on the metal part and across the air gap, respectively.

The electrospun PCL nanofibers were coated with laminin (Millipore, Temecula, CA) as the following. The electrospun fibers were immersed in a 0.1% poly-L-lysine (PLL) (Sigma-Aldrich) solution for 1 h at room temperature, followed by washing with phosphate buffered saline (PBS) buffer (Invitrogen) three times. Subsequently, the nanofiber sample was immersed in a laminin solution (26 µL, 50 µg/mL laminin solution diluted with 5 mL of...
DRG Culture. Embryonic day 8 (E8, stage HH35-36) chicks were removed from the eggs and decapitated. DRG were dissected from the thoracic region and collected in Hank’s buffered salt solution (HBSS) prior to plating. DRG were then placed onto the nanofiber scaffolds (1 DRG per sample) and incubated for 6 days in modified neurobasal (NB) media containing NB media, 1% ABAM, 1% N-2 supplement (InVitrogen, Carlsbad, CA), and 30 ng/mL rhβ-nerve growth factors (NGF) (R&D Systems, Minneapolis, MN).

SEM Characterization of DRG. The DRG were fixed in 3.7% formaldehyde for 30 min. Subsequently, it was dehydrated in ethanol with a series of concentrations (30, 50, 70, 90, 95, and 100%) and dried in vacuum. Finally, the sample was coated with platinum using a sputter prior to imaging by SEM. The accelerating voltage was 15 kV for imaging.

DRG Immunostaining. After 6 days incubation, the DRG were immunostained with the marker antineurofilament 200 (Sigma-Aldrich). Briefly, the DRG were fixed in 3.7% formaldehyde for 45 min and permeabilized by 0.1% Triton X-100 for 30 min. Then it was blocked in PBS containing 5% normal goat serum (NGS) (InVitrogen) for 1 h. Primary antibody diluted with PBS that contained 2% NGS was applied to the cells overnight at 4 °C. The antineurofilament 200 marker was detected using Alexa Fluor 488 goat antimouse IgG (1:200, InVitrogen) secondary antibody. After staining, fluorescent images were taken using a QICAM Fast Cooled Mono 12-bit camera (Q Imaging) attached to an Olympus microscope with Capture 2.90.1 (Olympus). Eccentricity of the neurite field, maximum length of extending neurites, and average length of extending neurites were simultaneously calculated from fluorescent images using custom-designed image processing software constructed in MATLAB (MathWorks Inc.). Neurite field eccentricity and length of neurite extension were specifically calculated for each sample as established measures of directed neurite growth and rate of neurite growth, respectively. Quantitative analysis was accomplished by separately fitting both the leading edge of the neurite field and the perimeter of the DRG cell mass, identified using 10 user-selected points, to a standard elliptical eq 1:41

\[
\frac{(x - h)^2}{a^2} + \frac{(y - k)^2}{b^2} = 1
\]

where point \((h, k)\) is the center of the ellipse and \(a\) and \(b\) are the ellipse’s semimajor and semiminor axes. Eccentricity of the neurite field was then calculated using eq 2:

\[
\text{Ecc.} = \sqrt{a^2 - b^2} / a
\]

Values of \(a\) and \(b\) were obtained from the specific elliptical equation fit to the leading edge of the neurite field. Average and maximum length of neurite extension was then calculated as the average distance and the greatest distance between the elliptical curve identifying the border of the DRG cell mass and the elliptical curve identifying the leading edge of the neurite field along a line oriented radially from the center of the DRG cell mass. Mean values and standard deviation were reported (\(n = 4 - 8\)). Statistical analysis of average neurite length, maximum neurite length, and eccentricity of neurite field was performed using the Scheffe’s post hoc test by analysis of variance at a 95% confidence level.

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Supporting Information Available: Figures S1–S3. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES AND NOTES