



Controlled Deposition and Collection of Electro-spun Poly(ethylene oxide) Fibers

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Abstract

Electro-spinning is a process by which sub-micron polymer fibers can be produced with an electrostatically driven jet of polymer solution (or polymer melt). Electro-spun fibers are typically collected in the form of non-woven mats, which are of interest for a variety of applications, including semi-permeable membranes, filters, composite reinforcement, and scaffolding used in tissue engineering. A characteristic feature of the electro-spinning process is the onset of a chaotic oscillation of the electro-spinning jet. The current work demonstrates the feasibility of dampening this instability and controlling the deposition of sub-micron polymer fibers (<300 nm in diameter) on a substrate through use of an electrostatic lens element and collection target of opposite polarity. Real-time observations of the electro-spinning process have been made with high speed, high magnification imaging techniques. Fiber mats and yarns electro-spun from polyethylene oxide have been analyzed by wide angle electron diffraction optical microscopy and environmental electron microscopy.

ACKNOWLEDGMENTS

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Contents

1.	Introduction	1
2.	Experimental	4
3.	Theory	4
4.	Results and Discussion	6
5.	Conclusion	12
	References	15
	Distribution List	17
	Report Documentation Page	19

Figures

1.	Standard Electro-spinning Apparatus	2
2.	Field Lines Calculated for a Syringe and Grounded Target Geometry .	3
3.	High Speed Image of Electro-spinning Process	3
4.	Forces Exerted on a Charge Element by Neighboring Charge Elements	6
5.	Multiple Field Electro-spinning Apparatus	7
6.	Field Lines Calculated for Multiple Field Electro-spinning Geometry .	8
7.	Image of Electro-spinning Jet 4 inches Below Capillary Orifice	9
8.	Fibers Electro-spun From 10% Solution Concentration of PEO in Water	10
9.	Electron Micrograph of PEO Fibers Electro-spun With Multiple Field Method	10
10.	Electro-spun Fibers Collected on a Rotating Drum With the Multiple Field Electro-spinning Method	11
11.	2D WAXD Pattern Obtained From PEO Electro-spun Yarns	12
12.	Plot of the Integrated Scattering Intensity as a Function of $2q$ From Figure 11b	13
13.	Powder WAXD Pattern for PEO Powder	13

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CONTROLLED DEPOSITION AND COLLECTION OF ELECTRO-SPUN POLY(ETHYLENE OXIDE) FIBERS

1. Introduction

For almost 100 [1] years, it has been known that polymer fibers can be generated from an electrostatically driven jet of polymer solution (or polymer melt). This process, known as electro-spinning, has received much attention in the last decade because of its ability to consistently generate polymer fibers that range from 5 to 500 nanometers in diameter. Because of the small pore size and high surface area inherent in electro-spun textiles [2,3,4], these fabrics show promise for exploitation in soldier protective clothing and filtration applications [5]. Other exciting applications that are being explored include scaffolding for tissue growth [6], optical and electronic applications [7,8].

The schematic for a typical electro-spinning apparatus is depicted in Figure 1 and its associated field lines in Figure 2. Polymer solution is forced through a syringe at a rate of about 0.5 ml/hr, resulting in the formation of a drop of polymer solution at the tip of the needle. A high voltage (5 to 15 kV) is applied to the syringe, causing the surface of the drop to distort into the shape of a cone, as depicted in Figure 1. When a critical voltage is exceeded (typically 5 kV for the solutions discussed in this report), a jet of solution erupts from the apex of the cone and is accelerated toward the electrically grounded collection target by the macroscopic electric field (see Figure 2). As this jet travels through the air, the solvent evaporates, leaving behind a polymer fiber to be collected on an electrically grounded target. Figure 3 is an image of the electro-spinning process obtained through high-speed photography. The image clearly depicts the random motion of the electro-spinning jet as it travels toward the target. Baumgarten [9] first depicted the chaotic nature of the electro-spinning jet motion by using high speed photography. Recent work by Reneker et al. [10] suggests that this chaotic motion, or “bending instability,” results (at least in part) from repulsive forces originating from the charged elements within the electro-spinning jet. Because of the chaotic motion of the electro-spinning jet as it travels to its target, deposition of electro-spun fibers on a stationary target is essentially random. This is perfectly acceptable for membrane and filter applications, which take advantage of the small pore size obtained by the random morphology of the non-woven electro-spun mat.

Collecting electro-spun fibers in the form of a yarn or “tow” (i.e., bundle of fibers) for post-processing to improve mechanical performance or depositing of electro-spun fibers on a substrate in specific places or patterns is problematic because of the random nature of the fiber deposition. Some efforts to improve control of the electro-spinning jet and deposition process include the use of both

mechanical and electrostatic means. It has been shown that electro-spun fibers can be collected in a textile where the fibers will be more or less oriented parallel to the direction of rotation [11] if the target is a drum rotating at high revolutions per minute. However, the area coated with electro-spun fibers is still quite large. It has also been shown that electro-spun fibers retain a significant portion of their charge upon deposition [2,12], and during the appropriate conditions [12], it is possible to have electro-spun fibers deposit preferentially on an aluminum screen forming a three-dimensional grid structure.

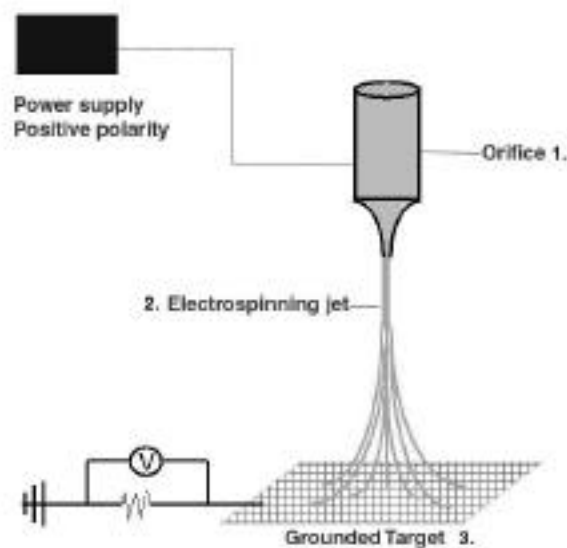


Figure 1. Standard Electro-spinning Apparatus.

Finally, Jaeger et al. [13] have demonstrated that it is possible to stop the precession of the electro-spinning jet about the tip of the syringe, which contributes to the chaotic motion of the electro-spinning jet. They accomplished this by using a single charged ring of like voltage and polarity placed concentrically about the syringe needle. Although this method of electro-spinning has the effect of stabilizing the jet at the point of initiation, the jet still undergoes bending instability as it proceeds toward the ground plane after passing through this single electrode region.

The objective of the current research has been to construct a novel electro-spinning apparatus that uses electrostatic fields (other than the one responsible for jet initiation) to dampen the bending instability inherent in the electro-spinning process. This apparatus has allowed for much greater control of the deposition and collection of electro-spun fibers. Wide angle X-ray diffraction (WAXD) analysis has been performed on electro-spun poly(ethylene oxide) (PEO) yarns collected by this apparatus, which indicated the presence of some

molecular orientation and a poorly developed crystalline microstructure within the electro-spun fibers.

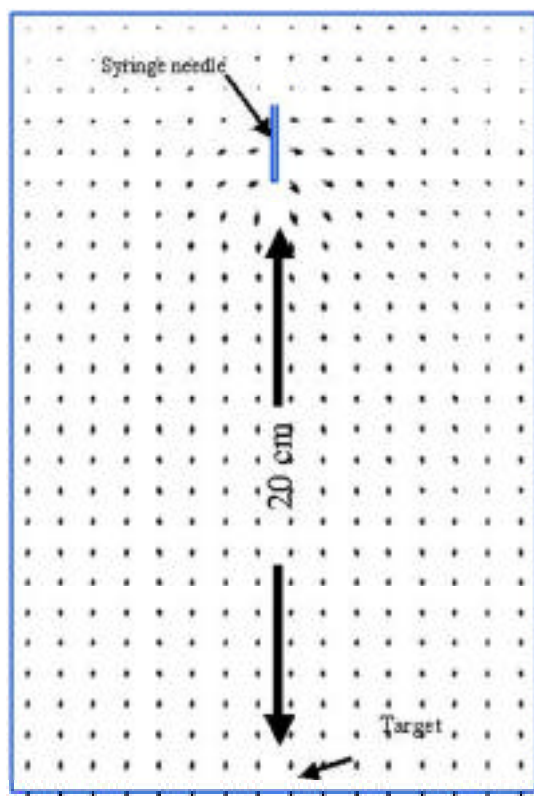


Figure 2. Field Lines Calculated for a Syringe and Grounded Target Geometry.

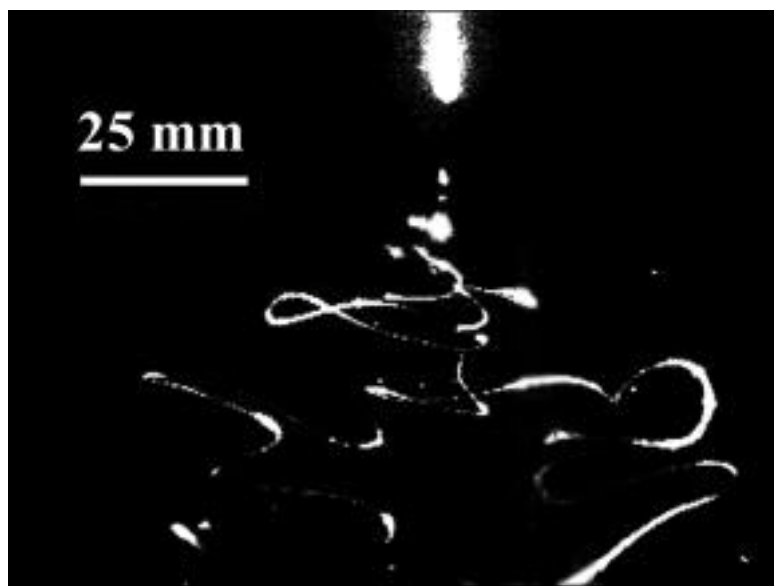


Figure 3. High Speed Image of Electro-spinning Process.

2. Experimental

PEO fibers were electro-spun from a 10% (wt) concentration of PEO in water. PEO of 400,000 molecular weight was used in making the solution. Two ES30P power supplies of positive polarity from Gamma High Voltage, Inc., were used to apply voltages of +5 to +15 kV to the vertically oriented syringe tip and rings. A Glassman Series EH high voltage power supply of negative polarity was used to apply a voltage of -9 to -12 kV to the collection plate situated at the bottom of the apparatus. Polymer solution was fed to the syringe needle tip through a Teflon[®] tube with a 0.125-inch inner diameter via a Harvard 2000 syringe pump. Representations of electric field lines associated with specific electro-spinning apparatus were calculated with the computer program, Student's Quick Field[™]. Electron micrographs of the electro-spun fibers were obtained with a Phillips Electroscan environmental scanning electron microscope.

High speed images were taken with a Photometrics cooled charge coupled device camera with a 3- by 2-k chip. The camera was attached to a Questar Schmidt-Cassegrain telescope in order to achieve high magnification from a distance of 40 cm. The electro-spinning jet was illuminated from behind with a Continuum Surelite II yttrium-aluminum-garnet laser. The laser emitted light at a wavelength of 532 nm, with a pulse duration of 10 ns. Optical micrographs of collected fibers were obtained with a Wild Makroscope M420 stereo-microscope.

Yarns of electro-spun fibers used in WAXD experiments were collected by "combing." This process involved passing two wooden splints spaced approximately 2.0 cm apart repeatedly through the electro-spinning jet about 2 cm above the collection target. WAXD analysis was performed with a Rigaku UltraX 18-kW rotating anode X-ray source and a Bruker Hi-Star 2D area detector.

3. Theory

When an external electric field is applied to a polymer solution, ions in the solution will aggregate around the electrode of opposite polarity. Positive ions travel to the negatively charged electrode and negative ions travel toward the positive electrode. This results in an excess of charge of opposite polarity in the volume of solution near an electrode. Consider a drop of polymer solution suspended at the tip of a metal syringe. When a voltage is applied to the metal syringe, the ions in the solution of like polarity will be forced to aggregate at the surface of the drop. The electric field generated by the surface charge will cause the drop to distort into the shape of a cone [14]. If the electric potential of the surface charge exceeds a critical value [14], the electrostatic forces will overcome

the solution surface tension. A thin jet of solution will erupt from the surface of the cone and travel toward the nearest electrode of opposite polarity or electrical ground. Although the details of charge motion in the electro-spinning jet are not well understood, it is believed that excess charge is essentially static with respect the moving coordinate system of the jet [10]. This means that the electro-spinning jet can be thought of as string of charge elements connected by a visco-elastic medium, with one end fixed at the point of origin and the other end free.

As discussed in the introduction, the free end of the electro-spinning jet follows a chaotic path as it travels toward the grounded collection plate, as seen in Figure 3. This chaotic motion, or instability, is the result of a complicated interaction of variables that include viscosity, surface tension, electrostatic forces, air friction, and gravity. Recent work by Reneker et al. [10] has attempted to create a model of jet motion that takes these variables into account. This effort has met with limited success, although detailed understanding of the jet motion remains elusive. For the present discussion, it is appropriate to focus on the electrostatic forces acting on the charged elements composing the jet and the visco-elastic response of the polymer solution to these electrostatic forces.

In their work, Reneker et al. [10] proposed the following mechanism for the onset of jet instability. Upon initiation, the jet of polymer solution is rapidly accelerated away from the syringe toward the grounded target by electrostatic forces. This has the effect of providing a longitudinal stress that stabilizes the jet, keeping it initially straight. At some distance from the point of initiation, the jet of polymer solution begins undergoing stress relaxation. The point along the jet where this occurs depends on the spinning voltage, which is proportional to the strength of the macroscopic electric field [10]. By increasing the voltage, the electric field strength increases the length of the stable jet [9]. Once stress relaxation occurs, Reneker et al. [10] proposed that electrostatic interaction between the charged elements of the jet begins to dominate the ensuing motion, initiating and perpetuating the chaotic motion of the jet. This initiation of the bending instability was described in the following manner.

If one considers three unconnected point charges of equal value in a line (see Figure 4), it can be seen that the center charge, B, is acted upon by two forces of equal magnitude and opposite direction given by the equation,

$$F = kq_a q_b / r^2 = kq_c q_b / r^2 \quad 1.$$

in which q_a , q_b , and q_c are charges of equal sign and magnitude, r is the separation between charges, and k is Coulomb's constant. Should a perturbation cause q_b to move out of line, there is a net lateral force on q_b , the magnitude of which is given by the equation

$$F_l = 2F \cos \theta \quad 2.$$

in which q is the angle that is formed by r and line perpendicular to jet axis.

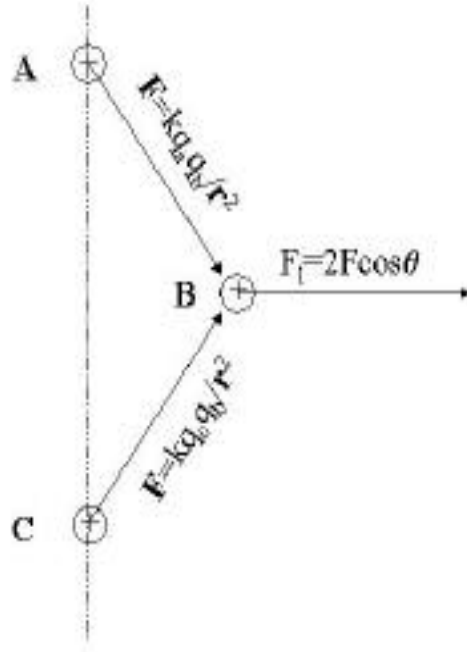


Figure 4. Forces Exerted on a Charge Element by Neighboring Charge Elements.

This lateral force, F_l , leads to an inherent instability predicted by Earnshaw's theorem [15]. Since an electrostatically driven jet can be thought of as a line of charged elements, it is thought that the instability predicted by Earnshaw is responsible for the onset of the chaotic motion of the free end of the electro-spinning jet [10]. Because any force, either electrostatic or mechanical in nature, to counter F_l is absent, the initially small perturbations grow unfettered, leading to the large looping motions that are commonly observed [2,9,10].

4. Results and Discussion

It is clear from the discussion in the previous section that electrostatic interactions between individual charge elements in the jet and between charge elements and the macroscopic electric field are primarily responsible for initiation and perpetuation of the bending instability. With this knowledge, it should be possible to design an electro-spinning apparatus that can dampen or eliminate the bending instability through control of the shape and strength of the macroscopic electric field that exists as a result of the potential difference

between the point of jet initiation and the collection target (see Figures 5 and 6). The electro-spinning apparatus discussed here is similar to one first described by Melcher and Warren [16], who were studying capillary instability and jet disintegration in electrostatically driven jets of low molecular weight fluids.

The apparatus depicted in Figure 5 is different from the setup illustrated in Figures 1 and 2 in two ways. The first innovation is a series of charged rings used as an electrostatic “lens” element that changes the shape of the macroscopic electric field from the point of jet initiation to the collection target. The field lines converge to a center line above the collection target. When the charged jet passes through this field, it is forced to the center in a manner that is analogous to a stream of water that is poured into a funnel. The second difference is that the collection target has a potential bias whose polarity is opposite that of the lens element and syringe. This allows for a continuous increase in the electric field strength and a corresponding increase in downward force on the jet as it approaches the collection target (i.e., $\text{Force} = q \cdot E$ in which E is the field strength and q is the magnitude of the charge element in Coulombs).

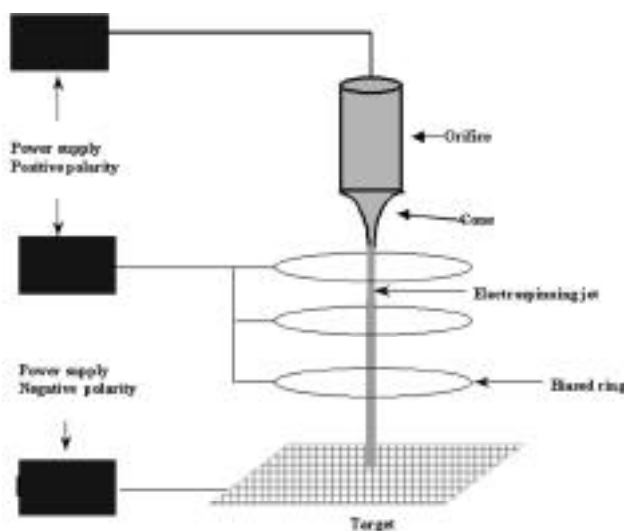


Figure 5. Multiple Field Electro-spinning Apparatus.

Figure 5 is a schematic of the multiple field electro-spinning apparatus. The apparatus consists of three high voltage power supplies. The first power supply is positive in polarity and is connected to the syringe. This is the source that supplies the critical voltage needed for jet initiation, which will be designated as the spinning voltage for the rest of this discussion. The second power supply provides the voltage for eight copper rings that are connected in series. It also provides positive polarity which will be referred to as the ring voltage. The third power supply is connected to the collection target and it provides negative polarity that will be referred to as the target voltage. The rings are 10 cm in diameter and are spaced at 1.9-cm intervals, approximately. The total distance

from the tip of the syringe to the collection target is about 20 cm. Values of +9 kV for the spinning voltage, +4 to +5 kV for the ring voltage, and 11 kV for the target voltage were usual for a typical experiment.

Figure 6 is a diagram depicting the macroscopic electric field generated by the apparatus described previously. The arrows indicate the direction of the electric field lines. The electric field is nearly uniform in direction down the center of the apparatus. At the top of the apparatus, there is a tendency for the field lines near the edge of the initiating electrode to point to the rings, which are at a lower potential. About 8 cm from the initiating electrode, the field lines start to converge at the center of the apparatus, indicating a net restoring force that acts on the charge jet. The field strength and corresponding restoring force increase as the jet approaches the target. Upon starting, there is some jet instability near the initiating electrode because of the edge effects. As the jet proceeds toward the target, the instability is dampened under the influence the converging electric field lines. Since the jet is continuous, the dampening of the instability and acceleration of the jet down stream act to stabilize the part of the jet near the initiating electrode.

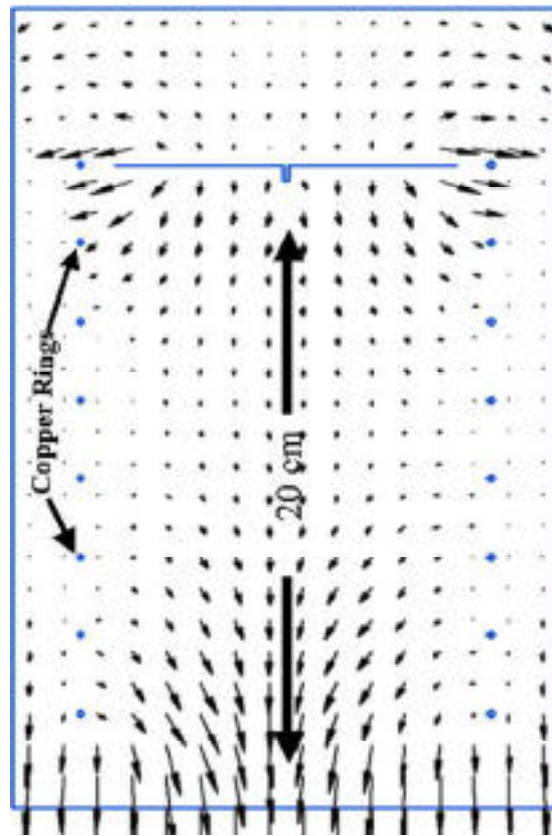


Figure 6. Field Lines Calculated for Multiple Field Electro-spinning Geometry.

Figures 7a and 7b are high-speed images of the electro-spinning jet generated with a 7% concentration solution of PEO in water with the apparatus described previously. These images were obtained about 10 cm below the syringe tip (half way to the target), which is well into the region where the bending instability normally occurs (see Figure 3). Figure 7a is an image of a straight jet with no evidence of any bending instability; Figure 7b illustrates the effect of decreasing the ring voltage from +5 kV to about +2.5 kV. In this case, the jet assumes the shape of a tight corkscrew as a result of relaxing the constraining electric field provided by the rings. When the ring voltage is increased again to 5 kV, the jet straightens.

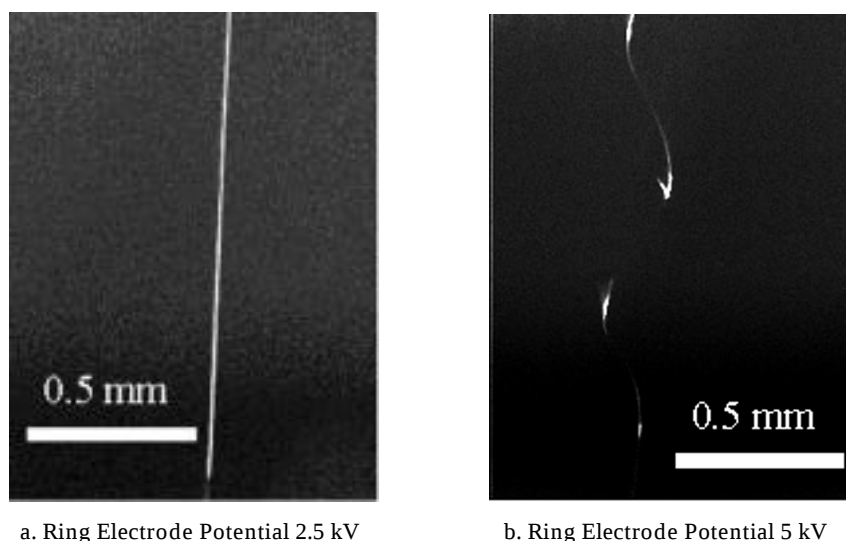
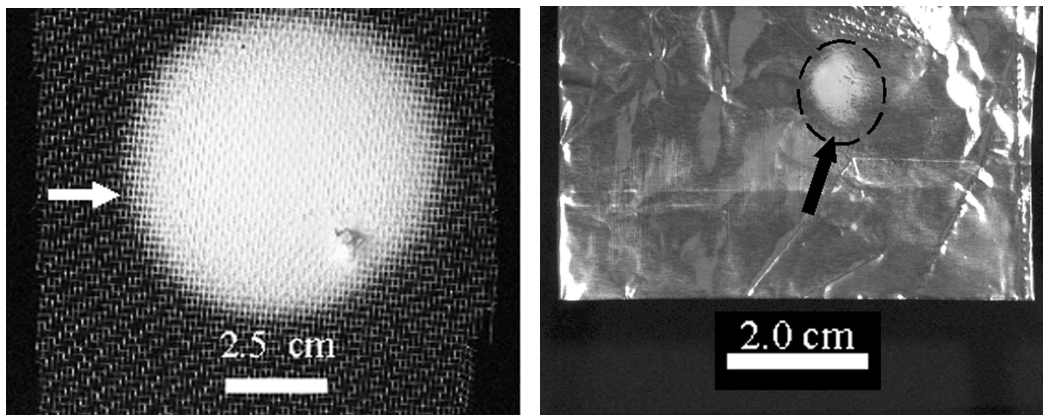


Figure 7. Image of Electro-spinning Jet 4 inches Below Capillary Orifice.

Figures 8a and 8b are images of fiber mats electro-spun from a 10% concentration of PEO in water. Figure 8a was produced with the standard electro-spinning method, without biased lens elements. The distance from tip to target was 17 cm and the spinning voltage was about +7 kV at the syringe tip, and the fibers were collected on an aluminum screen that was electrically grounded. The fiber mat in Figure 8b was obtained with the multiple field electro-spinning apparatus via positively biased rings orifice and a negatively biased foil collection target. Comparison of these two shows the significant decrease in the diameter (from ~7 cm in 8a to ~1 cm in 8b) of the area of coverage achieved with the multiple field electro-spinning apparatus.

This reduction in the area of coverage is the result of dampening the bending instability. The multiple field technique can also affect the size of the individual fibers. Figure 9 is an electron micrograph of fibers electro-spun from 10% PEO in water with the multiple field technique. The average diameter of the fibers is 270 nm, which compares to an average diameter of 400 nm reported for PEO

electro-spun from a 10% solution with the standard method of electro-spinning, which consisted of a positively biased syringe tip (7 kV), a grounded collection target, and a tip to target distance of about 17 cm [2]. It is thought that the reduction in the fiber diameter is attributable to the overall change in potential from syringe tip to target being much greater for the multiple field apparatus (21 kV) used to collect the sample in the figure [8,9]. The fibers in Figure 9 are uniform in their overall morphology and lack beads or junctions, which indicates that most of the solvent has evaporated by the time the fibers are collected on the target.



a. Standard Electro-spinning Technique Collected on Wire Mesh (diameter ~7 cm).

b. Multiple Field Technique Collected on Copper Foil (diameter ~1 cm).

Figure 8. Fibers Electro-spun From 10% Solution Concentration of PEO in Water (arrows indicate area of target covered by electro-spun fibers).

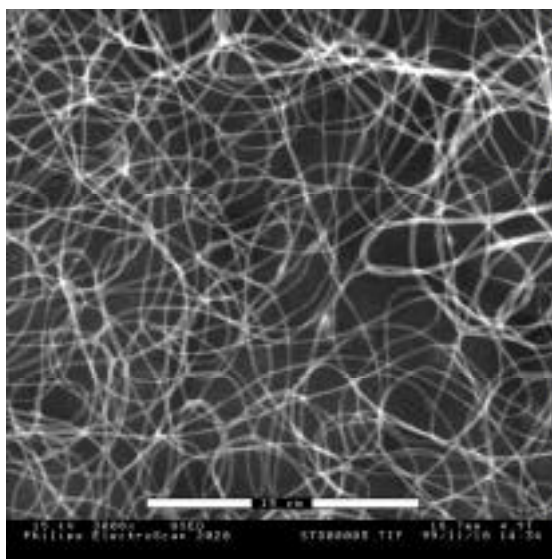


Figure 9. Electron Micrograph of PEO Fibers Electro-spun With Multiple Field Method. (Fibers were electro-spun from 10% solution concentration of PEO in water.)

When the chaotic motion of the jet is dampened, it becomes possible to deposit electro-spun fibers on a substrate in a more targeted fashion. When the target is a rotating drum covered with copper foil and charged to a potential of -11 kV, the electro-spun fibers are collected in a strip that is approximately 0.6 cm wide. Figure 10 shows two such strips collected within 0.6 cm of each other. Upon starting, some slight adjustment of the spinning voltage, ring voltage, and the solution feed rate are required in order to optimize the control of the deposition process. The strip on the right side of the image was collected during this optimization process and is slightly less well defined. After 1.5 hours, the drum was shifted slightly to the right and the second well-defined strip was collected during optimized conditions. The collection time for each strip was 1.5 hours.

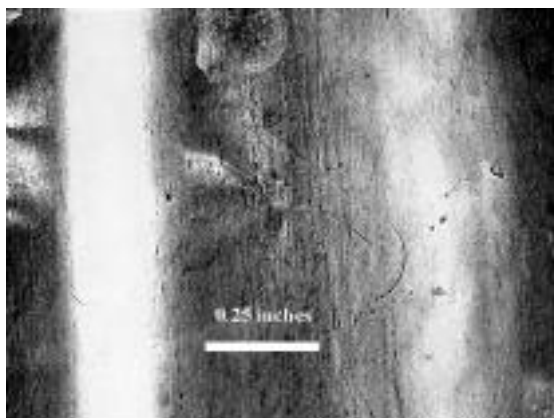


Figure 10. Electro-spun Fibers Collected on a Rotating Drum With the Multiple Field Electro-spinning Method.

It is also possible to collect electro-spun fibers in the form of a yarn with the multiple field apparatus. This was accomplished by a combing technique, in which two wooden splints spaced about 1 inch apart were repeatedly passed between the last ring and the collection plate, through the electro-spinning stream. This process allows for the collection of fibers that are more or less macroscopically oriented parallel to the direction of the combing. After 1 hour, enough electro-spun material was collected for the purpose of WAXD analysis. The fibers were removed from the splints and gently twisted into the yarns seen in Figure 11a. Figure 11b is the WAXD pattern obtained for the PEO yarns depicted in Figure 11a. The pattern shows six diffraction arcs that are characteristic of the monoclinic crystal structure of PEO. The two equatorial reflections correspond to the 120 crystallographic planes, and the four arcs in the quadrants correspond to the 112 planes. The particularly intense ring that occurs at $2\theta = 28.44^\circ$ corresponds to the 111 reflections of the silicon powder standard.

Figure 12 is a plot of the integrated intensity of the 2D WAXD pattern in Figure 11b as a function of 2θ . When this figure is compared to a powder WAXD diffraction pattern of PEO (see Figure 13), it can be seen that the 120 and 112

reflections of the fiber pattern are relatively broad and weak with respect to the background. In addition, the powder WAXD pattern of the PEO also contains numerous higher order reflections that are not present in the fiber pattern. The broad, weak nature of the fiber pattern reflections, together with the lack of higher order reflections, indicates that the crystalline microstructure of the electro-spun fibers is not well developed. The fact that the fiber pattern reflections (see Figure 11b) show distinct arcs rather than isotropic circles like those reported for electro-spun polyethylene by Larrondo et al. [17] suggests that some degree of molecular orientation results from this multiple field electro-spinning process. Note that no attempts were made to improve the overall crystallinity or the crystal orientation through post-process, although it may be possible to improve the degree of order in the multiple field-spun fibers with standard techniques, such as annealing under tension.

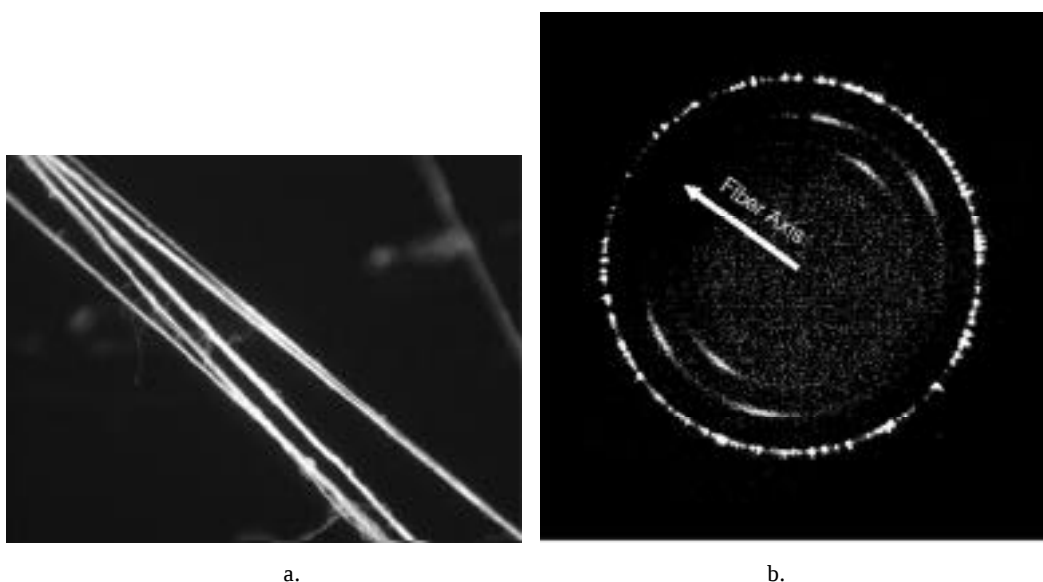


Figure 11. 2D WAXD Pattern Obtained From PEO Electro-spun Yarns.

5. Conclusion

It has been demonstrated that it is possible to control the deposition of electro-spun fibers through the use of an electrostatic lens element and biased collection target. By the application of a secondary external field of the same polarity as the surface charge on the jet, it is possible to control or eliminate the bending instability inherent in conventional electro-spinning experiments. This mechanism allows for greater control of the deposition of electro-spun fiber on a surface and for collection of electro-spun fibers in other forms besides non-woven mats. Analyses of yarns of electro-spun polyethylene oxide via WAXD

techniques indicate the presence of some molecular orientation and a poorly developed crystalline microstructure in the fibers.

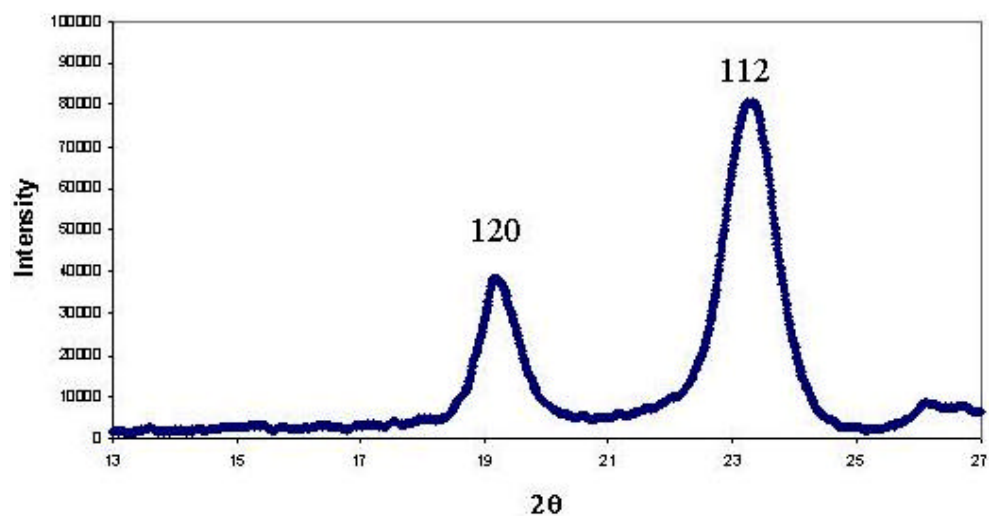


Figure 12. Plot of the Integrated Scattering Intensity as a Function of 2θ From Figure 11b.

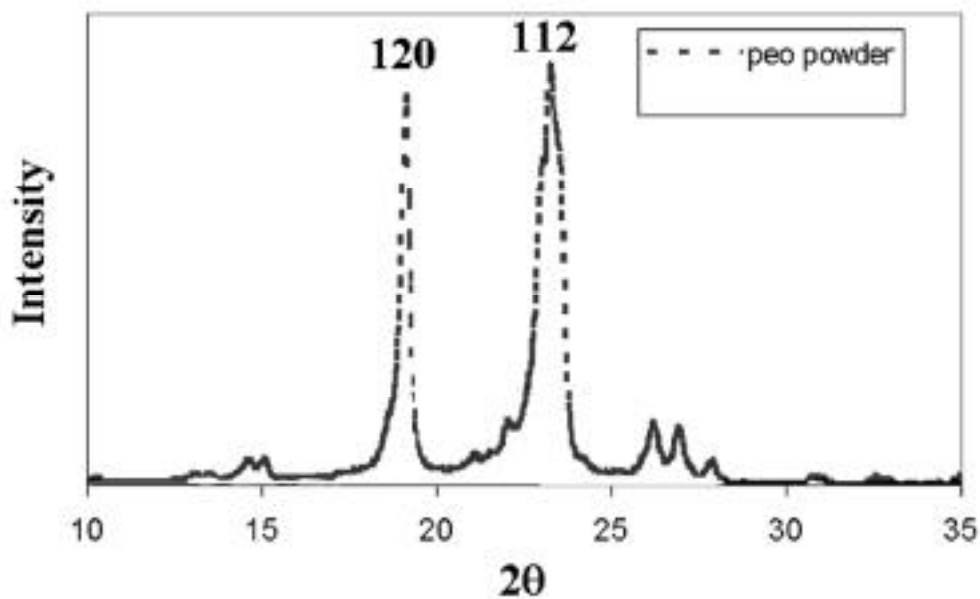


Figure 13. Powder WAXD Pattern for PEO Powder.

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13. ABSTRACT (Maximum 200 words) Electro-spinning is a process by which sub-micron polymer fibers can be produced with an electrostatically driven jet of polymer solution (or polymer melt). Electro-spun fibers are typically collected in the form of non-woven mats, which are of interest for a variety of applications, including semi-permeable membranes, filters, composite reinforcement, and scaffolding used in tissue engineering. A characteristic feature of the electro-spinning process is the onset of a chaotic oscillation of the electro-spinning jet. The current work demonstrates the feasibility of dampening this instability and controlling the deposition of sub-micron polymer fibers (<300 nm in diameter) on a substrate through use of an electrostatic lens element and collection target of opposite polarity. Real-time observations of the electro-spinning process have been made with high speed, high magnification imaging techniques. Fiber mats and yarns electro-spun from polyethylene oxide have been analyzed by wide angle electron diffraction optical microscopy and environmental electron microscopy.					
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