Polymers for Near-field Electrospinning with Spatial Control

Antonio Osamu Katagiri Tanaka, Héctor Alán Aguirre Soto

Abstract

Near-field electrospinning (NFES) is identified to be a technique able to fabricate polymer nano and micro fibers with accurate placement. In the past years (2006-2019), several polymer solutions have been successfully electrospun into fibers through several variants of the conventional NFES process. Each NFES variant intents to tailor the process parameters in order to improve the fibers' properties. This paper presents a review on the research and related development of electrospun fibers, emphasizing the used polymers, solvents, and fiber characteristics. Relevant summary of polymer solutions and near-field electrospinning processing conditions is provided in this paper.

Keywords: polymer, solvent, near-field electrospinning, NFES, fibers, spatial control

Contents		5	NF	ES Variants	16
1 Introduction 1.1 Types of electrospinning 1.1.1 High voltage power supply . 1.1.2 Polymer reservoir 1.1.3 Stretching forces 1.1.4 Polymer yielding 1.1.5 Dispensing nozzle 1.1.6 Nozzle-to-substrate distance	2 2 2 2 2 2 2 2		5.1 5.2 5.3 5.4 5.5	Low-Voltage NFES (LV NFES) [8] Scanning Tip Electrospinning [9] 3D Electrospinning [26] Electrohydro-dynamic 3D Print- patterning or Electrohydro- dynamic Jetting [17] Multinozzle NFES [33–35] Electrohydro-dynamic Writ-	
2 "Electro - DC - monoaxial - solution - NF - spinning"	2			ing or Mechanoelectrospinning (MES) [24] Electrohydro-dynamic Direct-Write (EDW) [37]	
3 Polymer Solution	2		- 0	Mechano-Electrospinning [38]	
3.1 Polymers	3		5.6 5.7	Suspension NFES [39] Helix Electrohydro-dynamic Printing (HE-printing) [31]	16
4 NFES Parameters	4			Electrohydro-dynamic (EHD) jet	
4.1 Nozzle spinneret	4			printing [23]	16
4.2 Applied Voltage	5		5.8	Airflow-assisted Electrohydro-	
4.3 Nozzle-to-substrate distance	5			dynamic Direct-writing (EDW) [32]	16
4.4 Electric field	6 6		5.9	Tethered Pyro-Electrohydro- dynamic Spinning (TPES) [19]	16
Email addresses: oskatagiri@gmail.com (Antonio		6	Con	aclusion	16
Osamu Katagiri Tanaka), alan.aguirre@tec.mx (Héct Alán Aguirre Soto)	tor	7	NF	ES Achievements & Challenges	17

References

1.1.3. Stretching forces

17

Centrifugal force
Blowing forces
Electric force
Microfluidic forces
Mechanical force

1. Introduction

Even though electrospinning is an old invention [1], it is currently a trending topic among researchers [2–4]. One of the reasons electrospinning is to be studied is its potential to fabricate polymer nano-fibers from a variety of polymers. The technique allows the production of thin continuous fibers with ease, with diameters down to 3 nm in some cases, which is something difficult to achieve by other techniques. Furthermore, the basic setup can be modified with ease to fabricate different fibers with diversified functionalities with different materials. The produced fibers can be aligned or unaligned. Besides, the electrospinning equipment is inexpensive and of small size, compared to the equipment of standard spinning techniques. On the other hand, the understanding of the electrospinning process has improved in the last years [5].

1.1. Types of electrospinning

The main components of the electrospinning technique are the fluid control unit (e.g. syringe pump) and a voltage power supply. The process also requires a target electrode or combination of electrodes on which the fibers can be collected. Figure 1 describes a typical near-field electrospinning set-up [5]. Two sub-techniques can be derived from electrospinning depending on the distance between the dispensing electrode and the collector. The process in which the electrospun jet can be controlled near the tip is called NFES or near-field electrospinning [6]. Moreover, if the distance between the collector and the dispensing needle is greater, the configuration is known as FFES or far-field electrospinning [7].

1.1.1. High voltage power supply
Direct Current
Alternating Current

1.1.2. Polymer reservoirPolymer meltPolymer solution

1.1.4. Polymer yielding
Atomization (particles)
Spraying (droplets)
Spinning (fibers)

1.1.5. Dispensing nozzle

Coaxial / Monoaxial

Multinozzle

1.1.6. Nozzle-to-substrate distanceFar Field SpinningNear Field Spinning

2. "Electro - DC - monoaxial - solution - NF - spinning"

Near-field electrospinning is considered to be an outstanding technique to fabricate polymer fibers with spatial control and it has suffered several modifications to improve the precision and accuracy of the fiber deposition. This paper intents to collect the NFES variants of electrospunable polymer solutions with spatial control in recent research.

3. Polymer Solution

In electrospinning, it is typically agreed that the diameter of the fibers increased with higher concentration due to greater viscosity which withstands stretching. In near field electrospinning, similar observations have been reported where concentration increases, fiber diameter increased [9, 10]. However, in separate studies by Pan et al. [11, 12] using poly(γ -benzyl α , l-glutamate) and polyvinylidene fluoride (PVDF) reported reduction in fiber diameter with increasing concentration.

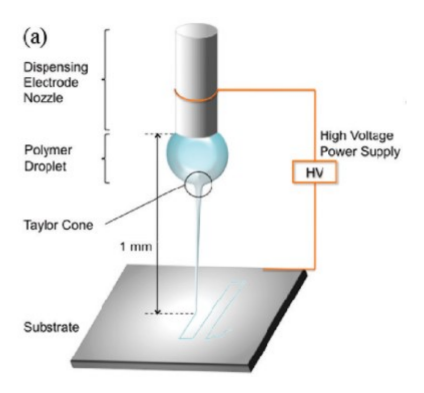


Figure 1: Typical near-field electrospinning set-up [8].

Table 1: Approximation process to estimate the critical polymer concentration. Several polymer concentrations are tried and the resulting jets are observed until a continuous stream is achieved.

Observation	Concentration Adjustement
Dripping, no stream	Increase
Splitting small droplets	Increase slightly
Steady stream	No concentration adjustment
Splitting large globs	Decrease slightly
Nozzle clogging	Decrease

3.1. Polymers

The polymer selection is in function on the intended application. For example, a fast dissolving hydrophilic polymer such as poly(ethylene oxide) (PEO) is used for fast drug delivery systems. Otherwise, slow dissolving polymers such as $poly(\varepsilon$ -caprolactone) (PCL) or poly(lactic-coglycolic acid) (PLGA) are implemented. [13]

The polymer molecular weight along with the polymer concentration and solvent selection have a direct effect on the solution viscosity, conductivity and surface tension, hence the solution behavior in the electrospinning process. The spunable viscosity range varies with the polymer and sol-

vent.

Solutions with low viscosity are prone to insufficient polymer chain entanglements to produce fibers. [13] On the other hand, if the solution is too viscous, then the surface tension cannot easily be overcome by the electric field. In both cases, the result can be droplets or particles forming rather than fibers; see Table 1.

3.2. Solvents

The solvent used must be capable of dissolving the polymer of interest at an appropriate concentration to form fibers, and must posses a suitable volatility. A low-volatility solvent like water may

fail to evaporate completely over the distance between the spinneret and the collector. When the fibers form, they will hence contain residual water owing to this incomplete evaporation. The residue solvent will subsequently evaporate from the fibers upon storage, resulting in ribbon-like (flattened) fibers, wrinkles on the fiber surface or fused fibers. On the other hand, a high-volatility solvent may evaporate very quickly, leading to larger fiber diameters (less time for elongation before solidification) and clogging of the spinneret (due to drying of the liquid at the spinneret before jetting, or drying of the Taylor cone during jetting). Solvents commonly used for electrospinning include ethanol, chloroform, dichloromethane and hexafluoroisopropanol.

Mixtures of miscible solvents can be used to ensure that sufficient polymer can be dissolved to give a solution of appropriate viscosity and volatility with suitable dielectric constant range to allow fiber formation. However, care must be taken because using a mixture of solvents with very different volatilities can result in porous fiber structures, as reported by Katsogiannis et al. for organic solvent mixtures with dimethyl sulfoxide (DMSO). [14] DMSO evaporates much more slowly than the organic solvents used, which results in its incorporation into the fibers. The DMSO will eventually evaporate, yielding porous fibers.

It is also important to take into account the surface tension of the solution. Solvents with very high surface tensions (e.g. water) can result in instability arising during the spinning process, and a broad range of fiber diameters in the products. If necessary, a surfactant can be added to reduce the surface tension, but this will be incorporated into the fibers produced.

4. NFES Parameters

To spin nano fibers at close distances, the initial diameter of the jet is required to be as small as possible since stretching of the thread is limited. Kameoka et al. [15] demonstrated that a small initial spinning radius can be achieved using an atomic force microscope tip with a small polymer

solution drop at the tip.

Near-field electrospinning, has exhibited to be capable fabricate nano fibers over and nano fiber patterns [16]. Nevertheless, having a small polymer solution drop at the nozzle tip limits the length of the fibers that can be fabricated in a continuous manner. Using a spinneret with a reservoir (e.g. syringe) of solution generally produces fibers with diameter of a few micrometers [17, 18], since it creates a limit to which the nozzle inner diameter can be reduced to allow the solution to flow through.

Coppola et al. [19] have showed a NFES variant that allows polymer nano fibers to be deposited directly from a polymer drop, averting the issue of nozzle clogging. The fibers are also prone soaking after deposition thus giving the fibers a semi-circular cross-section as depicted in Xue et al.'s [18] work.

4.1. Nozzle spinneret

The thinnest nozzles in literature so far are about 100 μm in diameter, for instance Chang et al. [9] used a 100 μm inner diameter needle tip to electrospin poly(ethylene oxide) (PEO) and Camillo et al. [20] used a micro-diameter tip Tungsten spinneret in a 26G needle to electrospin co-polymer, poly[2-methoxy-5-(2-methoxy-5)]ethylhexyloxy)-1,4-phenylenevinylene PPV) with poly(ethylene oxide) (PEO). The nozzle most commonly comprises a simple narrowbore, blunt-end metal needle. The diameter of the needle can vary, but most commonly researches work with internal diameters below 1 mm. This translates to needles of gauge 18–22. In general, this simple spinneret design can be used to achieve successful spinning. A bluntend rather than a tapered-end for the needle exit is important as the size distribution of the products increase with an increase in needle tip angle. However, it should be noted that there will be some interactions between the solvent and polymer molecules in the solution and the metal surface of the spinneret. There will exist some attractive forces between the polar groups in the polymer and the electropositive metal surface, which can act counter to the drawing force of the electric field and can pull the polymer solution back into the spinneret. It has been found that coating the spinneret exterior in a non-conductive and non-stick polymer such as Teflon can reduce these interactions. [21] As a result, the electrical energy can be more efficiently used to elongate and narrow the polymer jet, and narrower fibers can be produced. In addition, strong attractive forces between the polymer jet and the metal spinneret can result in fibers becoming attracted to the needle, leading to lower yields and potentially to blocking of the exit orifice. This effect too can be ameliorated using an epoxy coating. [22]

4.2. Applied Voltage

In recent literature, near field electrospinning has been studied to reduce the fiber diameter and to improve the fiber deposition accuracy. Camillo et al. [20] demonstrated that the application of a modified fine tip nozzle enables the fabrication of 100 nm diameter fiber at a nozzle-to-substrate distance of 500 μm and an applied voltage of 1.5 kV. On the other hand, Bisht et al. [8] and Chang et al. [9] came to the conclusion higher voltages yield thicker micro-fibers with a loss in jet stability.

This discrepancy in literature between the applied voltage and resulting fiber diameter is due to the relationship with other variables such as nozzle-to-substrate distance and solution deposition rate. For instance, if a high voltage is applied at a low deposition rate then electrospraying is achieved, meaning the formation of several noncontinuous fibers. The applied voltage shall be sufficient to break the surface tension and initiate the jet, but low enough to avoid multiple jets at the nozzle tip.

Bisht et al. [8] achieved the fabrication of thinner fibers with spatial control by reducing the applied voltage to 200-600~V at a nozzle-to-substrate distance of 0.5-1~mm. The low voltage setting does not create enough charge to break the polymer solution surface tension to initiate the electrospinning process.

Bisht et al. [8] and Chang et al. [9] initiated the electrospun fibers by mechanically pull the poly-

mer solution at the nozzle tip using a micro-probe tip. Chang and coworkers reduced the applied voltage from 1.5 kV to 600 V with a nozzle-to-substrate distance of 500 μm to yield a fiber diameter between 3 μm and 50 nm. With an applied voltage of 200 V and a nozzle-to-substrate distance of 1 mm, PEO nano fibers were deposited with a diameter about 20 nm.

In near-field electrospinning, the applied voltage has an impact on the produced fiber morphology. For instance, a voltage higher or lower to the optimum voltage will translate into an increase in fiber diameter. Song et al. [23] demonstrated that a decrease in voltage from 400 to 500 V can reduce the fiber diameter from 160 to about 60 nmwith a nozzle-to-substrate distance of 20 μm . The optimum voltage is achieved when a balance is attained between the stretching of the jet and the speed at which it hits the substrate. The increase of voltage yields thinner fibers as it causes greater stretching, and a greater jet acceleration.

Another workaround to break the polymer solution surface tension is to initialize the NFES process with a higher voltage and then lower the voltage once the jet is created. Huang et al. [24] implemented the previous and yield ordered fibers with a distance between adjacent fibers of 50 μm . In most cases, a positive voltage is applied to the spinneret.

4.3. Nozzle-to-substrate distance

In NFES, the fiber morphology can be altered by the control of the height between the nozzle and the substrate (collector). With the decrease of the nozzle-to-substrate distance, the electric field strength increases; however it can cause incomplete solvent volatilisation and possible short circuits between the collector and the nozzle tip.

An optimal nozzle-to-substrate distance shall be defined to ensure the fabrication of dry continuous fibers. If the solvent is not well evaporated, the produced fibers are prone to defects; on the other hand if solidification happens too fast, the solids can block the spinneret which can prevent a continuous fiber yield. Furthermore, the polymer jet will discharge itself as soon as possible, therefore long distances can result in low yields.

Typically, metal nozzle tips are used, with small inner diameters. From literature, needles with small diameters produce thinner fibers. A thin nozzle tip can help the reduction of the fiber diameter, but also it is more likely to become blocked.

4.4. Electric field

Recent literature suggests that the fiber morphology depends on the electric field profile created by the applied voltage during NFES. Since the electric field is an induced force that attracts the solution jet towards the desired location within the collector.

Bisht et al. [8] and Min et al. [25] have reported the ability to electrospin nano fibers with high accuracy. Min et al. [25] implemented a NFES setup with multiple "field-effect transistors" on a flexible polyacrylate collector with an x-y stage velocity of 13.3 cm/s to fabricate fibers with a diameter about 289 nm and a distance between adjacent fibers of 50 μm .

On the other hand, Bisht et al. [8] showed evidence of fabricated fibers with low-voltage NFES with high accuracy and precision. Bisht et al.'s suspended fibers were deposited over carbon posts with a distance between adjacent fibers of $100 \ \mu m$ with diameter of $30 \ \mu m$ [8].

The employment of guided electrodes in NFES, adapts the fabrication process to yield a more accurate fiber deposition. For instance, Kim et al. [26] manufactured ink patterns on a paper with silver nano particles. The printed patterns aid the fibers to land on the desired location. Kim et al. [26] electrospun the fibers with a distance between adjacent fibers of 150 μm .

Xu et al. [27] created a straight jet from the nozzle tip to the substrate using a guiding electrode underneath the collector. The purpose of the guiding electrode is to adjust the path of the NFES jet. With the guiding electrode implementation, the fiber's spread was reduced from 74 μm to 7 μm .

4.5. Substrate

Due to the close distance between the grounded substrate and the charged spinneret in NFES, the set up is prone to electrical shorts. In NFES, when a short circuit takes place, the electrospinning process is interrupted resulting in the fabrication of discontinuous fibers. Two workarounds to avoid electrical shorts is to lower the applied voltage and to install less conductive substrates [28, 29].

Liu et al. [28] discovered that the fiber alignment is improved by using a glass-cooper foil substrate, however the well aligned fibers are spoiled after prolonged depositions due to residual charges. Additionally, the effect of residual charges is amplified with the used collector substrate contains a conductive layer and a non-conductive layer [28].

On the other hand, Choi et al. [29] implemented a hydrophobic substrate to deposit the fibers with plasma treatment to increase the conductivity of selected areas. NFES was carried put with precise deposition as the fibers were placed as per the desired design within the hydrophilic substrate.

Polymer(s)	Solvent(s)	NFES Variant	Process Parameters and Fiber Characterization	Ref.
Poly(ethylene ox-	Deionized wa-	Low-Voltage NFES	Solution Concentration: 1, 2, and 3 $wt\%$ PEO	[8]
ide) (PEO; $MW =$	ter	(LV NFES)	Nozzle: 27 gauge type 304; stainless steel needle	
$4,000,000 \ g/mol)$			Solution deposition rate: lower than $1\mu L/h$	
			Nozzle-to-substrate distance: 1mm	
			Substrate composition: Pyrolyzed SU-8 carbon and Si	
			Applied voltage: polymer jet initiated at 400-600 V and	
			dispensed at 200-400 V	
			x-y stage velocity: $10-40mm/s$	
			Fiber Diameter: 50-425nm	
			Distance between adjacent fibers: Not determined	
Poly[2-	acetonitrile	Typical NFES pro-	Solution Concentration:	[30]
methoxy-5-(2-	toluene mixture	cess	$10mg$ of MEH-PPV in $2mL$ of toluene; $500\mu L$ of MEH-	
ethylhexyloxy)-1,4-	(65/35); acetic		PPV solution with $250mg$ of PEO in $3.5mL$ of acetonitrile /	
phenylenevinylene	acid toluene		toluene (65 / 35); $500\mu L$ of MEH-PPV solution with $250mg$	
(MEH-PPV; MW	(17/83); pure		of PEO in $3mL$ of acetic acid / toluene (17 / 83). The re-	
$= 380,000 \ g/mol)$	toluene		sulting MEH-PPV/PEO concentration is 0.08 $wt\%$	
with Poly(ethylene			Nozzle: mm-diameter tip Tungsten spinneret in a 26 gauge	
oxide) (PEO; MW			needle	
$= 300,000 \ g/mol)$			Solution deposition rate: $50\mu L/h$	
			Nozzle-to-substrate distance: $500\mu m$	
			Substrate composition: SiO2/Si (oxide thickness = 800	
			nm)	
			Applied voltage: around $1.3kV$	
			x-y stage velocity: $50cm/s$	
			Fiber Diameter: 100nm	
			Distance between adjacent fibers: around $100\mu m$	

∞	

Table 2 continued Poly(ethylene ox-	Water	Scanning Tip Elec-	Solution Concentration: 7wt% PEO	[9]
ide) (PEO; MV =		trospinning and	Nozzle: Needle outer diameter of $200\mu m$ and inner diame-	LJ
$300,000 \ g/mol)$		NFES	ter of $100\mu m$	
, 3,			Solution deposition rate: $0.1\mu L/h$	
			Nozzle-to-substrate distance: $500 \mu m$	
			Substrate composition: Not determined	
			Applied voltage: polymer jet initiated at $1.5 \ kV$ and dis-	
			pensed at $600V$	
			x-y stage velocity: $120mm/s$	
			Fiber Diameter: $709\pm131nm$; $49-74nm$ when applied volt-	
			age is $800V$	
			Distance between adjacent fibers: Not determined	
			Notes: $108m$ yield in $15min$ with a fiber diameter of	
			$709 \pm 131 nm$	
Poly(vinylidine flu-	N,N Dimethyl-	Helix Electrohydro-	Solution Concentration: $1.8g$ PVDF in $4.1g$ of DMF and	[31]
orid) (PVDF; MW	formamide	dynamic Printing	4.1g of acetone. The resulting concentration is 18% PVDF.	
$= 440,000 \ g/mol)$	(DMF)	(HE-printing)	Nozzle: Needle outer diameter of $510\mu m$ and inner diameter	
			of $260\mu m$	
			Solution deposition rate: $400nL/min$	
			Nozazle-to-substrate distance: 10-50mm	
			Substrate composition: Poly(dimethylsiloxane) (PDMS)	
			on Ecoflex	
			Applied voltage: $1.5-3kV$	
			x-y stage velocity: 0-400mm/min	
			Fiber Diameter: about $1.5-3\mu m$	
			Distance between adjacent fibers: Not determined	

0		

Table 2 continued	Dimethyl	Flootnobyzdno	Solution Concentration: POSS-PCU and POSS-PCL-	[17]
Polyhedral	Dimethyl	Electrohydro-		[17]
Oligomeric	acetamide	dynamic 3D Print-	PCU used in $20\%w/w$ concentration in DMAC	
Silsesquioxane-	(DMAC) and		Nozzle: needle of 750 μm in diameter	
Poly(Carbonate-	1-Butanol	Electrohydro-	Solution deposition rate: less than $1\mu L/min$	
Urea)Urethane		dynamic Jetting	Nozzle-to-substrate distance: about between $500\mu m$ to	
(POSS-PCU)			2mm	
and Polyhe-			Substrate composition: Not determined	
dral Oligomeric			Applied voltage: $8.0-10.0kV$	
Silsesquioxane			x-y stage velocity: $10mm/s$	
Poly(Caprolactone-			Fiber Diameter: $5-50\mu m$	
Poly(Carbonate-			Distance between adjacent fibers: $250\mu m$	
Urea)Urethane)			·	
(POSS-PCL-PCU)				
(Dry Polycarbon-				
(DI) I DIVCAIDOII-				
`				
ate MW = 2000				
`	Distilled water	Electrohydro-	Solution Concentration: $6wt\%$ PEO	[24]
ate $MW = 2000$ g/mol) Poly(ethylene ox-	Distilled water	Electrohydro- dynamic Writing	Solution Concentration: $6wt\%$ PEO Nozzle: Not determined	[24]
ate MW = 2000 g/mol) Poly(ethylene oxide) (PEO; MW =	Distilled water	•	Nozzle: Not determined	[24]
ate MW = 2000 g/mol)	Distilled water	dynamic Writing or Mechanoelectro-		[24]
ate $MW = 2000$ g/mol) Poly(ethylene oxide) (PEO; $MW =$	Distilled water	dynamic Writing	Nozzle: Not determined Solution deposition rate: 1200nL/min Nozzle-to-substrate distance: 7.5mm	[24]
ate $MW = 2000$ g/mol) Poly(ethylene oxide) (PEO; $MW =$	Distilled water	dynamic Writing or Mechanoelectro-	Nozzle: Not determined Solution deposition rate: 1200nL/min Nozzle-to-substrate distance: 7.5mm Substrate composition: Not determined	[24]
ate $MW = 2000$ g/mol) Poly(ethylene oxide) (PEO; $MW =$	Distilled water	dynamic Writing or Mechanoelectro-	Nozzle: Not determined Solution deposition rate: $1200nL/min$ Nozzle-to-substrate distance: $7.5mm$ Substrate composition: Not determined Applied voltage: polymer jet initiated at 2 kV and dis-	[24]
ate $MW = 2000$ g/mol) Poly(ethylene oxide) (PEO; $MW =$	Distilled water	dynamic Writing or Mechanoelectro-	Nozzle: Not determined Solution deposition rate: $1200nL/min$ Nozzle-to-substrate distance: $7.5mm$ Substrate composition: Not determined Applied voltage: polymer jet initiated at 2 kV and dispensed at $0.8\text{-}1kV$	[24]
ate $MW = 2000$ g/mol) Poly(ethylene oxide) (PEO; $MW =$	Distilled water	dynamic Writing or Mechanoelectro-	Nozzle: Not determined Solution deposition rate: $1200nL/min$ Nozzle-to-substrate distance: $7.5mm$ Substrate composition: Not determined Applied voltage: polymer jet initiated at 2 kV and dis-	[24]

		14010 01 0.1	winding (LDW)	0.21111111	
				Solution deposition rate: $30\mu L/h$	
				Nozzle-to-substrate distance: 2mm	
				Substrate composition: Silicon	
				Applied voltage: about $2kV$	
				x-y stage velocity: $1-20mm/s$	
				Fiber Diameter: $3.73 \pm 1.37 \mu m$	
				Distance between adjacent fibers: $5.13 \pm 6.67 \mu m$	
	Poly(Vinylidene	Acetone and	3D Electrospinning	Solution Concentration: $17wt\%$ PVDF; $1.7g$ of PVDF,	[26]
	Fluoride) (PVDF;	Dimethyl Sul-		5g of acetone, $0.5g$ of Capstone FS-66, $5g$ of DMSO	
	MW = 534,000	foxide (DMSO)		Nozzle: Needle inner diameter of $100\mu m$	
	g/mol)			Solution deposition rate: $14 nL/min$	
				Nozzle-to-substrate distance: $750\mu m$	
				Substrate composition: A4 size commercial printing pa-	
10				per (Double A)	
				Applied voltage: $1.9kV$	
				x-y stage velocity: $10mm/s$	
				Fiber Diameter: Not determined	
				Distance between adjacent fibers: Not determined	
	Poly(9-Vinyl Car-	Styrene	Typical NFES pro-	Solution Concentration: $3.96wt\%$ PVK in styrene	[25]
	bazole) (PVK; MW		cess	Nozzle: Needle inner diameter of $100\mu m$	
	$= 1,100,000 \ g/mol)$			Solution deposition rate: $500nL/min$	
				Nozzle-to-substrate distance: around 2.5mm	
				Substrate composition: Si/SiO2	
				Applied voltage: $3-4kV$	
				x-y stage velocity: $13.3cm/s$	
				Fiber Diameter: $289.26 \pm 35.37nm$	
$Au_{!}$				Distance between adjacent fibers: $50\mu m$	

Notes: 15m yield in 2min

0.21mm

Solution Concentration: 8wt% PEO

Nozzle: Outer airflow passage diameter: 1mm Airflow

gas pump pressure: 25kPa Inner liquid passage diameter:

Table 2 continued
Poly(ethylene ox-

ide) (PEO; MW =

 $300,000 \ g/mol)$

Deionized wa-

ter and ethanol

with a volume

ratio of 3:1

Airflow-assisted

writing (EDW)

Direct-

Electrohydro-

dynamic

[32]

Table 2 continued				
Polystyrene (PS;	1,2,4-Trichloro	Electrohydro-	Solution Concentration: 1 to $5wt\%$ PS	[23]
MW Not deter-	benzene	dynamic (EHD) jet	Nozzle: Glass nozzle inner diameter of $2\mu m$ and outer di-	
mined)		printing	ameter of $2.66\mu m$	
			Solution deposition rate: Not determined	
			Nozzle-to-substrate distance: $20, 30, 40\mu m$	
			Substrate composition: Si	
			Applied voltage: $500 \text{ to } 400V \text{ in } 25V \text{ increments}$	
			x-y stage velocity: $0.01-10mm/s$	
			Fiber Diameter: about $60-170\mu m$	
			Distance between adjacent fibers: Not determined	
Poly(ethylene ox-	Not determined	Typical NFES pro-	Solution Concentration: $3wt\%$ PEO	[16]
ide) (PEO; $MW =$		cess	Nozzle: Not determined	
$300,000 \ g/mol)$			Solution deposition rate: Not determined	
			Nozzle-to-substrate distance: $500\mu m$	
			Substrate composition: Si	
			Applied voltage: $1000V$	
			x-y stage velocity: $20cm/s$	
			Fiber Diameter: $300nm$	
			Distance between adjacent fibers: $25\mu m$	
Poly(ethylene ox-	Distilled water	Multinozzle NFES	Solution Concentration: $5wt\%$	[33]
ide) (PEO; $MW =$			Nozzle: four-nozzle and six-nozzle array with needle spacing	
$2,000,000 \ g/mol)$			changes from $1.5mm$ to $3.5mm$	
			Solution deposition rate: $1-3\mu L/min$	
			Nozzle-to-substrate distance: 2mm	
			Substrate composition: Not determined	
			Applied voltage: 1.7-2.7kV	
			x-y stage velocity: Not determined	
			Fiber Diameter: 5.47µm	
			Distance between adjacent fibers: 3-5 mm	

-	_	
Ν	೨	
	_	

August 22, 2019

Table 2 continued				
Poly(ethylene ox-	Distilled water	Multinozzle NFES	Solution Concentration: $5wt\%$	[34]
ide) (PEO; $MW =$			Nozzle: Dual-28G-needle array with needle inner diameter	
$2,000,000 \ g/mol)$			of $0.18mm$ and outer diameter of $0.36mm$; with needle spac-	
			ing changes from $2.0mm$ to $3.0mm$	
			Solution deposition rate: $0.2\mu L/min$	
			Nozzle-to-substrate distance: 3.0-4.0mm	
			Substrate composition: Not determined	
			Applied voltage: $2.0-3.0kV$	
			x-y stage velocity: $20mm/s$	
			Fiber Diameter: Not determined	
			Distance between adjacent fibers: $218-326\mu m$	
Poly(ethylene ox-	Distilled water	Multinozzle NFES	Solution Concentration: $5 wt\%$	[35]
ide) (PEO; $MW =$			Nozzle: Dual-28G-needle array with needle inner diameter	
$2,000,000 \ g/mol)$			of $180\mu m$ and outer diameter of $360\mu m$; with needle spacing	
			changes of $2.0mm$	
			Solution deposition rate: $0.2\mu L/min$	
			Nozzle-to-substrate distance: 4.0mm	
			Substrate composition: chromium-plated glass	
			Applied voltage: $2.5kV$	
			x-y stage velocity: $20mm/s$	
			Fiber Diameter: Not determined	
			Distance between adjacent fibers: 2.3002-2.7224mm	
Poly(ethylene ox-	Not determined	Typical NFES pro-	Solution Concentration: $2wt\%$	[27]
ide) (PEO; $MW =$		cess	Nozzle: G30 needle with inner diameter of 0.15mm	
$4,000,000 \ g/mol)$			Solution deposition rate: Not determined	
			Nozzle-to-substrate distance: 1-3mm	
			Substrate composition: Silicon	
			Applied voltage: $1250V$	
			x-y stage velocity: Not determined	
			Fiber Diameter: Not determined	
			Distance between adjacent fibers: $20\mu m$	

\vdash	
ಲು	

August 22, 2019

Table 2 continued				
Gelatin	Acetic Acid and	Typical NFES pro-	Solution Concentration: $11wt\%$ gelatin, $30wt\%$ water,	[18]
(porcine skin; MW	Ethyl Acetate	cess	35.4wt% acetic acid, $23.6wt%$ ethyl acetate	
$Not \ determined)$			Nozzle: 19G needle tip with outer diameter of 1.08mm	
			Solution deposition rate: Not determined	
			Nozzle-to-substrate distance: 1.25mm	
			Substrate composition: Poly(Dimethylsiloxane) (PDMS)	
			films	
			Applied voltage: $1000V$	
			x-y stage velocity: Not determined	
			Fiber Diameter: around $2-3\mu m$	
	,		Distance between adjacent fibers: $40\mu m$	
Poly(ethylene ox-	Water/Ethanol	Typical NFES pro-	Solution Concentration: PEO concentrations of 16% and	[36]
ide) (PEO; MW = 1)	$(\mathrm{v}/\mathrm{v}=60/40)$	cess	18%	
$300,000 \ g/mol)$			Nozzle: $40\mu m$	
			Solution deposition rate: Not determined	
			Nozzle-to-substrate distance: 1mm	
			Substrate composition: Planar silicon	
			Applied voltage: $1.7kV$	
			x-y stage velocity: $0.36m/s$	
			Fiber Diameter: $5.15\mu m$	
D 1 / +1 1	TTT / TT / 1	T21 . 1 1	Distance between adjacent fibers: Not determined	[0=1
Poly(ethylene ox-	Water/Ethanol	Electrohydro-	Solution Concentration: 14wt% PEO	[37]
ide) (PEO; MW = 1)	$(\mathrm{v}/\mathrm{v}=3/1)$	dynamic Direct-	Nozzle: Stainless needle with inner diameter of $210\mu m$ and	
$300,000 \ g/mol)$		Write (EDW)	outer diameter of $400\mu m$	
			Solution deposition rate: $50\mu L/h$	
			Nozzle-to-substrate distance: 2mm	
			Substrate composition: Poly(ethylene terephthalate) (PET)	
			Applied voltage: $3kV$	
			x-y stage velocity: $700mm/s$	
			Fiber Diameter: $15-35\mu m$	
			Distance between adjacent fibers: $70\mu m$	

۰		4	
H	\wedge		

August 22, 2019

Poly(ethylene ox-	Deionized wa-	Mechano-	Solution Concentration: $3wt\%$ PEO	[38]
ide) (PEO; $MW =$	ter	Electrospinning	Nozzle: Stainless steel nozzle with inner diameter of $160\mu m$	
$300,000 \ g/mol)$			and outer diameter of $310\mu m$	
, 3,			Solution deposition rate: $50nL/min$	
			Nozzle-to-substrate distance: 2-5mm	
			Substrate composition: Silicone	
			Applied voltage: polymer jet initiated at $2kV$ and dispensed at $1kV$	
			x-y stage velocity: $200-400mm/s$	
			Fiber Diameter: from 344 ± 32 to $214\pm27nm$	
			Distance between adjacent fibers: Not determined	
Poly(co-Glycolic)	Dimethyl Car-	Tethered Pyro-	Solution Concentration: Not determined	[19]
acid (PLGA; MW	bonate (DMC)	Electrohydro-	Nozzle: nozzle-free	
$Not \ determined)$		dynamic Spinning		
		(TPES)	directly on a flat substrate	
			Nozzle-to-substrate distance: Taylor's cone is focused	
			and put in direct contact with the collector	
			Substrate composition: Poly(tetrafluoroethylene)	
			(PTFE) coated glass slide	
			Applied voltage: pyro-electric field of between 2.7	
			$x10^7 \ V/m \text{ and } 5.5x10^7 \ V/m$	
			x-y stage velocity: Not determined	
			Fiber Diameter: 304.7nm	
			Distance between adjacent fibers: Not determined	

Table 2 continued				
Poly(ethylene oxide) (PEO; MW = $4,000,000$ g/mol) with Tetrabutylammonium tetrafluoroborate (TBF; MW Not determined) and SU-8 2002	N,N Dimethyl- formamide (DMF)	Typical NFES process	Solution Concentration: SU-8/PEO/TBF blend with 0.75wt% PEO, 1wt% TBF; the blend is diluted with 30vol% DMF µmµm Solution deposition rate: Not determined Nozzle-to-substrate distance: Not determined Substrate composition: Brass disk with a diameter of 38mm Applied voltage: 980V x-y stage velocity: Not determined Fiber Diameter: Not determined Distance between adjacent fibers: Not determined	[6]
Poly(ethylene oxide) (PEO; 200,000 g/mol)	Water:Ethanol (3:2)	Suspension NFES	Solution Concentration: $14wt\%$ PEO Nozzle: stainless steel needle (25 G) with inner diameter of $0.25mm$ Solution deposition rate: $3nL/s$ Nozzle-to-substrate distance: between 0.5 and $10mm$ with $0.5mm$ increments Substrate composition: Planar silicon electrodes Applied voltage: $1.6kV$ x-y stage velocity: 50 , 150 , and $250mm/s$ Fiber Diameter: $300nm$ Distance between adjacent fibers: 0.1 and $0.5mm$	[39]
Poly(ethylene oxide) (PEO; MW = $400,000 \ g/mol$)	Deionized water	Typical NFES process	Solution Concentration: $10wt\%$ PEO Nozzle: $32G$ metal needle Solution deposition rate: (Jet impact speed of $5mm/s$) Nozzle-to-substrate distance: $0.5mm$ Substrate composition: p-type silicon wafer Applied voltage: $400V$ x-y stage velocity: $5mm/s$ Fiber Diameter: $Not\ determined$ Distance between adjacent fibers: $50\mu m$	[40]

5. NFES Variants

[SECTION UNDERWORK]

Nanofibers are fibers with diameters in the nanometer range. The development of nanofibers has greatly enhanced the scope for meeting up the modern world challenges.

Currently there are two types of electrospinning systems available for producing nanofiber: needle based electrospinning and needleless electrospinning. This paper summarizes the basic mechanism of various types of needle based and needleless spinning systems described in various literatures by many researchers.

5.1. Low-Voltage NFES (LV NFES) [8]

Some differences have been discovered between LV-NFES and conventional NFES. Low voltage near field electrospinning produces thinner fibers with lower voltages. Moreover, when implementing a moving stage, the fibers are affected by the mechanical stretching. Bisht et al. [8] reported that thinner diameters are yield with the increase of the x-y stage velocity, and larger diameters by decreasing the stage velocity.

5.2. Scanning Tip Electrospinning [9]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.3. 3D Electrospinning [26] Electrohydro-dynamic 3D Print-patterning or Electrohydro-dynamic Jetting [17]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.4. Multinozzle NFES [33–35]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.5. Electrohydro-dynamic Writing or Mechanoelectrospinning (MES) [24] Electrohydro-dynamic Direct-Write (EDW) [37] Mechano-Electrospinning [38]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.6. Suspension NFES [39]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.7. Helix Electrohydro-dynamic Printing (HE-printing) [31]
Electrohydro-dynamic (EHD) jet printing [23]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.8. Airflow-assisted Electrohydro-dynamic Direct-writing (EDW) [32]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

5.9. Tethered Pyro-Electrohydro-dynamic Spinning (TPES) [19]

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua.

6. Conclusion

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua. Ut enim ad minim veniam, quis nostrud exercitation ullamco laboris nisi ut aliquip ex ea commodo consequat. Duis aute irure dolor in reprehenderit in voluptate velit esse cillum dolore eu fugiat nulla pariatur. Excepteur sint occaecat cupidatat non proident, sunt in culpa qui officia deserunt mollit anim id est laborum.

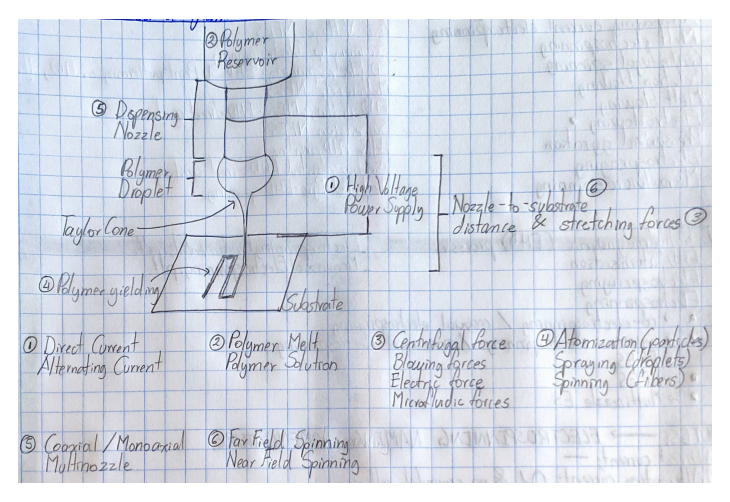


Figure 2:

7. NFES Achievements & Challenges

Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut labore et dolore magna aliqua. Ut enim ad minim veniam, quis nostrud exercitation ullamco laboris nisi ut aliquip ex ea commodo consequat. Duis aute irure dolor in reprehenderit in voluptate velit esse cillum dolore eu fugiat nulla pariatur. Excepteur sint occaecat cupidatat non proident, sunt in culpa qui officia deserunt mollit anim id est laborum.

References

- [1] F. Anton, Process and apparatus for preparing artificial threads (1930). doi:https://patents.google.com/?q=D01D5%2f0076.
- [2] Z.-M. Huang, Y. Z. Zhang, M. Kotaki, S. Ramakrishna, A review on polymer nanofibers by electrospinning and their applications in nanocomposites,

- Composites Science and Technology 63 (15) (2003) 2223-2253. doi:10.1016/S0266-3538(03)00178-7.
- [3] D. H. Reneker, A. L. Yarin, Electrospinning jets and polymer nanofibers, Polymer 49 (10) (2008) 2387–2425. doi:10.1016/J.POLYMER.2008.02.002.
- [4] J. D. Schiffman, C. L. Schauer, A Review: Electrospinning of Biopolymer Nanofibers and their Applications, Polymer Reviews 48 (2) (2008) 317–352. doi:10.1080/15583720802022182.
- [5] Q. Li, Chapter 7: Liquid Crystal-Functionalized Nano- and Microfibers Produced by Electrospinning - Liquid Crystals Beyond Displays: Chemistry, Physics, and Applications, John Wiley & Sons, 2012. doi:9781118078617.
- [6] A. Cisquella-Serra, M. Magnani, Álvaro Gual-Mosegui, S. Holmberg, M. Madou, M. Gamero-Castaño, Study of the electrostatic jet initiation in near-field electrospinning, Journal of Colloid and Interface Science 543 (2019) 106–113. doi:10.1016/J.JCIS.2019.02.041.
- [7] S. K. Nataraj, K. S. Yang, T. M. Aminab-Polyacrylonitrile-based nanofibers—A havi, state-of-the-art review. Progress in Polymer Science 37 (3)(2012)487 - 513.

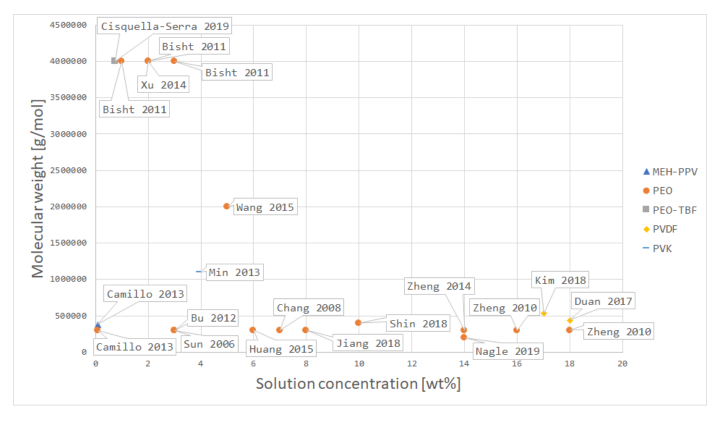


Figure 3:

- doi:10.1016/J.PROGPOLYMSCI.2011.07.001.
- [8] G. S. Bisht, G. Canton, A. Mirsepassi, L. Kulinsky, S. Oh, D. Dunn-Rankin, M. J. Madou, Controlled Continuous Patterning of Polymeric Nanofibers on Three-Dimensional Substrates Using Low-Voltage Near-Field Electrospinning, Nano Letters 11 (4) (2011) 1831–1837. doi:10.1021/nl2006164.
- [9] C. Chang, K. Limkrailassiri, L. Lin, Continuous near-field electrospinning for large area deposition of orderly nanofiber patterns, Appl Phys Lett (2008) 3doi:10.1063/1.2975834.
- [10] J. Zheng, Y. Z. Long, B. Sun, Z. H. Zhang, F. Shao, H. D. Zhang, Z. M. Zhang, J. Y. Huang, Polymer nanofibers prepared by low-voltage near-field electrospinning, Chinese Physics B 21 (4) (2012) 1–6. doi:10.1088/1674-1056/21/4/048102.
- [11] C.-T. Pan, C.-K. Yen, Z.-H. Liu, H.-W. Li, S.-W. Kuo, Y.-S. Lu, Y.-C. Lai, Poly(γ -benzyl α , l-glutamate) in Cylindrical Near-Field Electrospinning Fabrication and Analysis of Piezoelectric Fibers (2014).
- [12] C.-T. Pan, C.-K. Yen, S.-Y. Wang, Y.-C. Lai, L. Lin, J. C. Huang, S.-W. Kuo, Near-field electrospinning enhances the energy harvesting of hollow PVDF piezoelectric fibers, RSC Advances 5 (103) (2015) 85073–85081. doi:10.1039/C5RA16604G.
- [13] S. Chakraborty, I.-C. Liao, A. Adler, K. W. Leong, Electrohydrodynamics: A facile technique

- to fabricate drug delivery systems, Advanced Drug Delivery Reviews 61 (12) (2009) 1043–1054. doi:10.1016/j.addr.2009.07.013.
- [14] K. A. G. Katsogiannis, G. T. Vladisavljević, S. Georgiadou, Porous electrospun polycaprolactone (PCL) fibres by phase separation, European Polymer Journal 69 (2015) 284–295. doi:10.1016/j.eurpolymj.2015.01.028.
- [15] J. Kameoka, H. G. Craighead, Fabrication of oriented polymeric nanofibers on planar surfaces by electrospinning, Applied Physics Letters 83 (2) (2003) 371– 373. doi:10.1063/1.1592638.
- [16] D. Sun, C. Chang, S. Li, L. Lin, Near-Field Electrospinning (2006). doi:10.1021/nl0602701.
- [17] A. Gupta, A. M. Seifalian, Z. Ahmad, M. J. Edirisinghe, M. C. Winslet, Novel Electrohydrodynamic Printing of Nanocomposite Biopolymer Scaffolds, Journal of BIOACTIVE AND COMPATIBLE POLYMERS 22 (2007). doi:10.1177/0883911507078268.
- [18] N. Xue, X. Li, C. Bertulli, Z. Li, A. Patharagulpong, A. Sadok, Y. Y. S. Huang, Rapid Patterning of 1-D Collagenous Topography as an ECM Protein Fibril Platform for Image Cytometry, PLoS ONE 9 (4) (2014) e93590. doi:10.1371/journal.pone.0093590.
- [19] S. Coppola, V. Vespini, G. Nasti, O. Gennari, S. Grilli, M. Ventre, M. Iannone, P. A. Netti, P. Ferraro, Tethered Pyro-Electrohydrodynamic Spinning

- for Patterning Well-Ordered Structures at Microand Nanoscale, Chem. Mater 26 (2014) 3360. doi:10.1021/cm501265j.
- [20] D. D. Camillo, V. Fasano, F. Ruggieri, S. Santucci, L. Lozzi, A. Camposeo, D. Pisignano, Near-field electrospinning of conjugated polymer light-emitting nanofibers, Nanoscale 5 (2013) 11637–11642. doi:10.1039/C3NR03094F.
- [21] Q. Xiang, Y.-M. Ma, D.-G. Yu, M. Jin, G. R. Williams, Electrospinning using a Teflon-coated spinneret, Applied Surface Science 284 (2013) 889–893. doi:10.1016/j.apsusc.2013.08.030.
- [22] Q. Wang, D. G. Yu, S. Y. Zhou, C. Li, M. Zhao, Electrospun amorphous medicated nanocomposites fabricated using a Teflon-based concentric spinneret, E-Polymers 18 (1) (2018) 3–11. doi:10.1515/epoly-2017-0110.
- [23] C. Song, J. A. Rogers, J.-M. Kim, H. Ahn, Patterned polydiacetylene-embedded polystyrene nanofibers based on electrohydrodynamic jet printing, Macromolecular Research 23 (1) (2015) 118–123. doi:10.1007/s13233-015-3024-2.
- [24] Y. Huang, Y. Duan, Y. Ding, N. Bu, Y. Pan, N. Lu, Z. Yin, Versatile, kinetically controlled, high precision electrohydrodynamic writing of micro/nanofibers, Scientific Reports 4 (1) (2015) 5949. doi:10.1038/srep05949.
- [25] S.-Y. Min, T.-S. Kim, B. J. Kim, H. Cho, Y.-Y. Noh, H. Yang, J. H. Cho, T.-W. Lee, Large-scale organic nanowire lithography and electronics, Nature Communications 4 (1) (2013) 1773. doi:10.1038/ncomms2785.
- [26] J. Kim, B. Maeng, J. Park, Characterization of 3D electrospinning on inkjet printed conductive pattern on paper, Micro and Nano Systems Letters 6 (1) (2018) 12. doi:10.1186/s40486-018-0074-1.
- [27] J. Xu, M. Abecassis, Z. Zhang, P. Guo, J. Huang, K. Ehmann, J. Cao, Accuracy Improvement of Nano-fiber Deposition by Near-Field Electrospinning, International Workshop on Microfactories IWMF2014 (9th) (2014).
- [28] Z. H. Liu, C. T. Pan, L. W. Lin, J. C. Huang, Z. Y. Ou, Direct-write PVDF nonwoven fiber fabric energy harvesters via the hollow cylindrical near-field electrospinning process (2014) 25003— 25014doi:10.1088/0964-1726/23/2/025003.
- [29] W. S. Choi, G. H. Kim, J. H. Shin, G. Lim, T. An, Electrospinning onto Insulating Substrates by Controlling Surface Wettability and Humidity, Nanoscale Research Letters 12 (2017). doi:10.1186/s11671-017-2380-6
- [30] D. D. Camillo, V. Fasano, F. Ruggieri, S. Santucci, L. Lozzi, A. Camposeo, D. Pisignano, Near-field electrospinning of conjugated polymer light-emitting nanofibers, Nanoscale 5 (2013) 11637–11642. doi:10.1039/C3NR03094F.

- [31] Y. Duan, Y. Ding, Z. Xu, Y. Huang, Z. Yin, Helix Electrohydrodynamic Printing of Highly Aligned Serpentine Micro/Nanofibers., Polymers 9 (9) (sep 2017). doi:10.3390/polym9090434.
- [32] J. Jiang, X. Wang, W. Li, J. Liu, Y. Liu, G. Zheng, J. Jiang, X. Wang, W. Li, J. Liu, Y. Liu, G. Zheng, Electrohydrodynamic Direct-Writing Micropatterns with Assisted Airflow, Micromachines 9 (9) (2018) 456. doi:10.3390/mi9090456.
- [33] H. Wang, S. Huang, F. Liang, P. Wu, M. Li, S. Lin, X. Chen, Research on Multinozzle Near-Field Electrospinning Patterned Deposition, Journal of Nanomaterials 2015 (2015) 1–8. doi:10.1155/2015/529138.
- [34] Z. Wang, X. Chen, J. Zeng, F. Liang, P. Wu, H. Wang, Controllable deposition distance of aligned pattern via dual-nozzle near-field electrospinning, AIP Advances 7 (3) (2017) 035310. doi:10.1063/1.4974936.
- [35] Z. Wang, X. Chen, J. Zhang, Y.-J. Lin, K. Li, J. Zeng, P. Wu, Y. He, Y. Li, H. Wang, Fabrication and evaluation of controllable deposition distance for aligned pattern by multi-nozzle near-field electrospinning, AIP Advances 8 (7) (2018) 075111. doi:10.1063/1.5032082.
- [36] G. Zheng, W. Li, X. Wang, D. Wu, D. Sun, L. Lin, Precision deposition of a nanofibre by nearfield electrospinning, Journal of Physics D: Applied Physics 43 (41) (2010) 415501. doi:10.1088/0022-3727/43/41/415501.
- [37] J.-Y. Zheng, H.-Y. Liu, X. Wang, Y. Zhao, W.-W. Huang, G.-F. Zheng, D.-H. Sun, Electrohydrodynamic Direct-Write Orderly Micro/Nanofibrous Structure on Flexible Insulating Substrate, Journal of Nanomaterials 2014 (2014) 1–7. doi:10.1155/2014/708186.
- [38] N. Bu, Y. Huang, X. Wang, Z. Yin, Materials and Manufacturing Processes Continuously Tunable and Oriented Nanofiber Direct-Mechano-Electrospinning Written by ously Tunable and Oriented Nanofiber Direct-Written by Mechano-Electrospinning (2012).doi:10.1080/10426914.2012.700145.
- [39] A. R. Nagle, C. D. Fay, Z. Xie, G. G. Wallace, X. Wang, M. J. Higgins, A direct 3D suspension nearfield electrospinning technique for the fabrication of polymer nanoarrays, Nanotechnology 30 (19) (2019) 195301. doi:10.1088/1361-6528/ab011b.
- [40] D. Shin, J. Kim, J. Chang, Experimental study on jet impact speed in near-field electrospinning for precise patterning of nanofiber, Journal of Manufacturing Processes 36 (2018) 231–237. doi:10.1016/J.JMAPRO.2018.10.011.