1 Contents

2	1	1 Introduction								
3	2	2 Experiment								
4		2.1	Sample Preparation	3						
5		2.2	Morphology and microstructure characterizations	4						
6			2.2.1 Polarized Optical Microscope (POM)	4						
7			2.2.2 Small Angle X-ray Scattering(SAXS)	4						
8			2.2.3 Wide Angle X-ray Scattering(WAXS)	4						
9		2.3	Dynamic Property Characterization	4						
10		2.4	Thermal Property Characterization	5						
11			2.4.1 Thermogravimetric Analysis (TGA)	5						
12			2.4.2 Differential Scanning Calorimetry (DSC)	5						
13	3	Resi	t and Analysis 6							
14		3.1	Crystallization behavior	6						
15	4 Conclusion									
16	5 Acknowlagement									
17		Abstract								
18		Lore	m ipsum dolor sit amet, consectetuer adipiscing elit. Ut purus elit, vestibulum ut, placerat ac, adipiscing vit	ae,						
19	fel	is. Cu	rabitur dictum gravida mauris. Nam arcu libero, nonummy eget, consectetuer id, vulputate a, magna. Don	vida mauris. Nam arcu libero, nonummy eget, consectetuer id, vulputate a, magna. Donec						
20	vehicula augue eu neque. Pellentesque habitant morbi tristique senectus et netus et malesuada fames ac turpis egestas.									
21	Mauris ut leo. Cras viverra metus rhoncus sem. Nulla et lectus vestibulum urna fringilla ultrices. Phasellus eu tellus									
22	sit	sit amet tortor gravida placerat. Integer sapien est, iaculis in, pretium quis, viverra ac, nunc. Praesent eget sem vel leo								
	ultrices bibandum. Aaneen faucibus. Marbi dolor pulla, malasuada au pulvinar at mollis as, pulla Curabitur austar									

semper nulla. Donec varius orci eget risus. Duis nibh mi, congue eu, accumsan eleifend, sagittis quis, diam. Duis eget
orci sit amet orci dignissim rutrum.

6 1 Introduction

There is a great deal of interests in the development of viable green technologies aimed at the enhancement of biodegradable and biocompatible polymer materials, such as the blend of poly \varepsilon-caprolactone (PCL) and polylactide (PLA) Blending is a low cost, easy implement process for modification of polymer mixture. Mixing several immiscible polymer with different physical and chemical properties has possiblity to improve the overall constituents' performance. Nowadays, the blending material of poly ε -caprolactone (PCL) and polylactide (PLA) shows a wide application in many areas. PLA has excellent biocompatibility, biodegradability and dynamic property, which shows broad prospects for tissue engineering development and other areas. But pure PLA has some drawbacks, including poor toughness, low 33 degradation rate. Moreover, the crystallinity of PLA is low. These drawbacks severely restrain the application of PLA 34 material. Researchers have attempted several polymer to make blends with PLA. PHB, PCL, PP and other material which has good performance has been attempted in recent years. Physical property improvment has been found in some researchers' blending experiments. PCL The blend of PCL 37 and PLA shows high degradation rate, better tensile strength, which is the properties originated from PLA, while PCL with much slower degradation rate and better toughness. The blend of PCL and PLA is a promising material which can meet the requirement of environment and physical conditions. Qiaolian Lv et al.[1] used a injection molding blend of PCL and PLA. They got a maximum $\sigma = 29.8 \pm 0.9$ MPa and $E = 922.5 \pm 9.8$ Mpa. For electrospun process, Pisani et al.[2] got a maximum $E = 49.10 \pm 0.12$ Mpa. 42 The physical property of PCL-PLA blend depends on the supermolecular structure, which is dominated by the 43 crystallization condition. The melting point (T_m) and glass transient temperature (T_g) of PCL are far lower than those of 44 PLA, and the crystallization temperature (T_c) of PCL is even lower than the Tg of PLA. Someone's research manifests

The goal of this research work to improve the physical properties of the blend. Stretching of the electrospun mat

that the presence of minor PCL phase favors cold crystallization of PLA in their blend system because PCL is in its

molten state during PLA crystallization, reducing system viscosity as a result, or acts as additional substrates.

- 49 induces the inner fibers to align in one direction. The unmolten PLA fibers offer crystallization loci for PCL phase.
- one the other hand, the molten PCL phase favors cold crystallization of PLA fibers. The present works lack this kind
- research method. Does this process improve the performance of blends? This question interests the author. Therefore, in
- 52 this work electrospinning, stretching and hot pressing experiments are carried. The some morphology, microstructure,
- 53 dynamic and thermal properties are characterized.

2 Experiment

55 2.1 Sample Preparation

- 56 Firstly, electrospinning process was carried out. PLA was supplied by PCL was supplied by 20 wt% PCL solution and
- 57 20 wt% PLA solution were prepared. 2 nozzles containing PCL solution and 1 nozzle containing PLA solution were
- used in the electrospinning process, so the mass ratio of PLA: PCL was 33:67. The electrospinning voltage was set
- ₅₉ at 8 keV. The humidity of electrospun environment was 40%, the temperature was 25 °C. The electrospun mats were
- 60 made as standard size samples.
- On the next stage, the electrospun mat was stretched by a mechanical tester. The electrospinning mats were
- stretched to a series of elongation ratios. The velocity of the tensile process was 4mm/minute, which is a low speed
- to avoid the fracture of the samples. 25%, 50%, 75%, 100% and 125% elongation ratio mats were made. After that
- the mats' double edges were fixed by heat-resistant tape in order to keep the elongation status since the mats had a
- rebound trend. Subsequently, the extended mats were put between two foils which sticked to 300 °C-resistant film.
- The mats were deposited in a heat oven which kept an 80 °C environment lasting 1 hour for PCL's melting. After 1
- 67 hour, the mats were transfered into another heat oven which kept a 30 °C environment lasting 30 minutes for isothermal
- 68 crystallization.
- Finally, the sample was tailored into a dimension of 10mm × 5mm, and the thickness was recorded by a film thick-
- ness gauge. Abudant standard sample with same length and width were prepared for the following characterizations.

2.2 Morphology and microstructure characterizations

72 2.2.1 Polarized Optical Microscope (POM)

- POM is an effective facility to observe the microstructure of polymer material. With Maltese cross phenomenon in bire-
- frigence polymer crystall, the amrphous and crystallization zone can be clearly distinguished. A Linkam heating stage
- vas used to handle the electrospinning films with the same heat treatment process as the experiment in subsection 2.1.
- 76 The melting and the cold crystallization process were observed and recorded by a Leica POM.

77 2.2.2 Small Angle X-ray Scattering(SAXS)

- 78 Small Angle X-ray Scattering(SAXS) experiment was carried on beamline BL19U2, Shanghai Synchrotron Radiation
- ₇₉ Facility. A Pilatus 1M detector(981 × 1043 pixels with a pixel size of 172mm) was performed to collect the SAXS
- pattern. The X-ray wavelength was 0.103 nm, and the sample to detector distance was set at 2131 mm. Considering the
- scale of X-ray pattern on BL19U2 is much larger than the diameter of PLA fibers inside the samples. It was essential
- to focus the X-ray pattern. A beryllium compound reflective lens(CRLs) was deployed to get a $5\mu m$ diameter X-ray
- pattern. Besides a portable POM was placed on the light path in order to determine location on blends characterized by
- 84 the SAXS method.

85 2.2.3 Wide Angle X-ray Scattering(WAXS)

86 2.3 Dynamic Property Characterization

- 87 The dynamic properties of the neat PCL and its blends with various annealing histories were determined by an In-
- stron Mechanical Tester (ASTM D638) at a crosshead speed of 50 mm·min⁻¹) at 25 °C using the dog-bone shaped
- specimens. Strength and modulus values reported here represent an average of the results for tests run on 6 specimens.

Table 1. Crystallinity of components in PCL/PLA blends

Sample	Crystallinity (%) of PCL phase	Crystallinity (%) of PLA phase	Melting point(°C) of PCL phase
0%	22.1	-	58.7
25%	20.8	-	56.6
50%	21.1	-	58.6
75%	23.2	-	59.1
100%	29.3	-	63.0
125%	29.0	-	58.5

 χ_c calculated using Δ_m^c of PCL of 139.5(J·g⁻¹)

2.4 Thermal Property Characterization

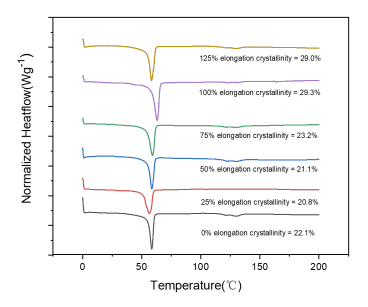
2.4.1 Thermogravimetric Analysis (TGA)

- TGA measurements were carried out in a Mettler Toledo TGA 2 thermal analyzer. The experiments were performed
- under nitrogen atmosphere (flow rate of 50 mL \cdot min⁻¹). Each sample was heated from 0 °C to 600 °C at 10 °C \cdot min⁻¹.
- The initial degradation temperatures (T_0) were determined at 5% mass loss, whereas temperatures at the maximum
- 95 degradation rate (*Tmax*) were calculated from the first derivative of the TGA curves (DTG).

96 2.4.2 Differential Scanning Calorimetry (DSC)

- DSC experiments were performed in a Mettler Toledo DSC 3+ under nitrogen atmosphere (flow rate of $50 \text{ mL} \cdot \text{min}^{-1}$).
- Sample weights of 2 mg were sealed in aluminum pans and heated from 0 °C to 200 °C at 10 °C · min⁻¹. The degree
- of crystallinity(χ_c) was calculated through Equation 1

$$\chi_c = 100\% \times \left[\frac{\Delta H_m - \Delta_{cc}}{\Delta H_m^c}\right] \frac{1}{W_{PCL}} \tag{1}$$



3 Result and Analysis

01 3.1 Crystallization behavior

4 Conclusion

5 Acknowlagement

References

- [1] Lv, Q. *et al.* Crystallization of poly(e-caprolactone) in its immiscible blend with polylactide: insight into the role of annealing histories. *RSC Advances* **6**, 37721–37730 (2016).
- [2] Pisani, S. *et al.* Design of copolymer pla-pcl electrospun matrix for biomedical applications. *Reactive and Func-*tional Polymers **124**, 77–89 (2018).