Effect of parameters on the morphology and fibre diameters of edible electrospun chitosan-cellulose acetate-gelatin hybrid nanofibres

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Abstract: Electrospinning is the favorite process to fabricate fibres with diameter in the range nanoscale through the action of electric field. In this study, 3-7% chitosan, 18.0% cellulose acetate and 30.0% gelatin solution in aqueous 80% acetic acid solution were blended at the volume ratio of 4:1:5 have been successfully electrospun. The effect of processing parameters and the concentration of the polymer solution on the morphology and diameter of electrospun were investigated. The morphology and diameter of electrospun fibres were observed by scanning electron microscope. The diameters of chitosan-cellulose acetate-gelatin nanofibres ranging from 78.94 to 421.05 nm. The results showed that the fibre diameters increased when the solution concentration and flow rate were increased, whereas the fibre diameters decreased when the applied voltage and distance between tip to collector were increased. The conditions of the solution concentration 18.8 %wt, applied voltage at 23 kV, flow rate at 11.67 µL/min and collector distance at 10 cm were selected to prepare the desirable electrospun nanofibres for applications and the further research.

Keyword: Electrospinning, electrospun nanofibre, edible electrospun

1 Introduction

Nanofibres was a nanomaterial that was tiny fibre with diameter in the range of 1-100 nanometre. It has a variety technique to prepared nanofibres such as drawing, template synthesis, phase separation, self-assembly and electrospinning [1-3]. However, electrospinning was the most attractive process to fabricate nanofibres due to that the apparatus was uncomplicate and flexible for the raw material selection. The basis apparatus of electrospinning consists of (i) high voltage power supply, (ii) syringe with a blunt ended stainless needle, (iii) syringe pump and (iv) ground collector [4,5]. In the electrosprining process, polymer solution is taken in the syringe with needle then the positive electrode which has a high voltage power supply was connected to a needle while the negative electrode was connected to a ground collector. When the high voltage was applied, the droplet shape at the end of needle will be changed from a semicircle into a conical shape as known “Taylor’s cone” due to the charge - charge repulsion [5-8]. During the jet transportation cross the high voltage electric field, the whipping instability motion and stretching of fibre will be observed. The solvent will be evaporated. Thin and dry fibres on the collector are obtained [8,9]. These fibres were called “electrospun nanofibres”. Electrospun nanofibres presented the outstanding properties such as high surface area to volume or mass ratio, light weight, small inter-fibrous pore size with high porosity [5,7,10]. They were used for several applications such as filtration membranes, nanofiber composites, wound dressing, tissue scaffolding, drug delivery systems and food packaging [7,11].

Morphology and fibre diameters of electrospun depended on various parameters that including solution properties, process parameters, and ambient parameters [2,5,12]. Understanding the parameters will help to fabricate the desirable electrospun nanofibers. Chitosan, cellulose acetate and gelatin are the natural polymers that were selected in this study due to there are edible, biocompatible, biodegradable, eco-friendly and low toxicity. Dhandayuthapani et al. [1] and Haider et al. [13] successfully fabricated electrospun nanofibre mats from chitosan-gelatin blend for applications in the biomedical field. In addition, several reports demonstrated that cellulose acetate -gelatin blend could use for electrospun nanofibre mats preparation as well [14,15]. However, as far as we have known that the availability data for the electrospun nanofibre from the blend of chitosan-cellulose acetate-gelatin hybrid was
never presented. The aim of this study was to investigate the effect parameter including of concentration of polymer solution, applied voltage, flow rate and distance between tip to collector on the electrospinning process. The morphologies of electrospun including diameters and diameter distribution were determined. The outcomes of this study may give the optimum condition for fabricate uniform electrospun fibres with desirable properties for applications in food sector.

2 Materials and methods

Chitosan powder (degree of deacetylation 95.04%) and gelatin powder (250 bloom) were purchased from Thai Food and Chemical Co., Ltd (Thailand). Cellulose Acetate (CA, average Mn ~ 30,000) was purchased from Aldrich Co. Ltd (USA). Glacial acetic acid (>99%) was purchased from Aldrich Co. Ltd (New Zealand). Chitosan, CA and gelatin were separately dissolved in 80% acetic acid (v/v) under continuous stirring at room temperature until completely dissolved. Then, the Chitosan-Gelatin-Cellulose acetate were prepared at the volume ratio of 4:1:5, respectively. The concentration of Chitosan-CA-Gelatin in mixed polymer were 18.0, 18.8 and 19.6% wt with the constant concentration of Gelatin and CA. The morphology and diameters of the electrospun nanofibres was investigated using scanning electron microscopy (SEM JSM-7800F, JEOL, MA, USA). Statistical analyses were performed by ANOVA with 3 replicates. Differentiate analysis among the data was analyzed using Duncan at p-value <0.05.

Electrospinning set up and process

The mixed Chitosan-CA-Gelatin solution was loaded in 10 mL plastic syringe connected with 18 gauges metal needle. The flow rate (11.67–25.00 μL/min.) was controlled by a syringe pump (New Era NE-300, USA). A high voltage DC power supply (20–26 kV) was applied at distance of 8-12 cm. The electrospinning was performed at room temperature (27±2°C) and all electrospun nanofibres were dried overnight to remove residual solvent.

3 Results and discussion

Effect of solution concentration

The solution of Chitosan in acetic acid was varied from 8.0%, 9.0%, 10.0%, 11.0% and 12.0% wt at 25 kV, the distance collector was 8 cm and flow rate was 25.00 μL/min. It was found that no fibre formation and only droplets were collected as shown in Fig. 1. Average droplet’s size was 1.042 - 43.235 μm. The diameter and roughness were increased by increasing Chitosan concentration. Gelatin electrospun fibre mats were obtained by the 30.0%wt gelatin at 23 kV, flow rate was 11.67 μL/min. The electrospun fibre mats were only obtained by 18.0%wt CA at 25 kV, flow rate was 50.00 μL/min. and the distance was 10 cm (Fig. 2A and 2B). The diameter of Gelatin fibres ranged from 384.61-2564.10 nm and average diameter was 1061.53±500.25 nm. On the other hand, Oraby et al. [16] reported that no fibres at 30.0% wt Gelatin concentration was investigated, the fibres were first appeared at 40.0% wt Gelatin (69-138 nm) and increased to when Gelatin concentration was increased.

Fig. 1. SEM images (100x) of electrospun drops at different Chitosan concentration (A) 8.0%, (B) 9.0%, (C) 10.0%, (D) 11.0% and (E) 12.0% wt

Fig. 2. SEM images of electrospun fibre mats (A) Gelatin electrospun; (B) CA electrospun (2,000x (A) and 2,500x (B))

Whereas Haider et al. [13] showed that at below 15.0%wt Gelatin solution, the fibres were not detected and the smooth fibres electrospun nanofibres was firstly detected at 15.0%wt with average diameter of 149 nm at 20 kV and the electrical field distance at 10 cm. The increasing of the solution concentration could be resulted to the viscosity properties, especially the entanglement properties of polymer. At the high concentration, above entanglement concentration, the nanoscaled fibres would be no obtained. However, the flat fibres and the ribbon-type fibre were achieved [12, 17]. For the diameter of CA fibres ranged from 47.61-142.85 nm and the average diameter was 85.14±24.05 nm. Moreover, beads were observed on the fibres. On the contrary, Han et al. [18] have reported that CA at higher concentrations than 17.0 wt% in acetic acid could be electrospun, long uniform nanofibres without beads were observed.

Fig. 3. SEM images (10,000x) of electrospun nanofibres and diameters distribution for the various solution (A) 18.0wt% (B) 18.5wt% and (C) 19.6wt%
Fig. 3 showed the Chitosan-CA-Gelatin nanofibres and diameter distribution obtained at concentration of 18.0, 18.8 and 19.6 %wt in 80% acetic acid. All of solution concentrations of Chitosan-CA-Gelatin could be electrospun. The diameter of Chitosan-CA-Gelatin nanofibres ranged from 78.94-421.05 nm and the average diameter were 148.42±41.12, 152.32±14.48 and 248.42±72.02 nm, respectively. The fibres diameter were increased continuously by increasing polymer concentration from 18wt% to 19.6wt% (Fig. 4).

![Fig. 4 The relation between the fibre diameters and polymer concentration](image)

However, the diameters at 18.0 and 18.8wt% solution concentration were different without significant ($p>0.05$). The smooth, non-bead and continuously fibres were obtained. At 19.6 wt% solution concentration, the junction and sticky were observed due to the higher viscosity and surface tension of polymer jets [19]. In contrast, the diameter fibres were significant increased ($p<0.05$). Polymer concentration and viscosity are related factor, when one increased, the other would be increased that resulted to the size of diameter fibres. At low concentration, polymeric micro or nano-particle will be obtained, the suitable concentration, smooth, uniform and continuous nanofibres was presented [5,10,12,16].

Effect of applied voltage
Voltage involved the force in the electrical field. Applied voltage resulted to the formation of Taylor cone according to the repulsive force in an opposite direction of the surface tension. The overcome state resulted on the initiation of fine jet from the cone and flying to collector [5,6,12]. In this experiment, effect of applied voltage on fibre morphology at 20, 23 and 26 kV were investigated. The average fibres diameters at 20, 23 and 26 kV were 170.21±42.15, 152.32±41.48 and 106.95±35.39 nm respectively. Smooth, non-bead and continuous fibres were obtained from all treatments. In addition, fibres diameters were decreased while applied voltage was increased (Fig. 5 and 6), it was in accordance with the previous reports [12,18]. However, the beads of PVA/alginate was also detected [7]. Moreover, the electrospun fibres with bigger beads were obtained at 45 kV. It was investigated that when applied voltage was increased, number of junctions and fibres diameters decreased while the size of beads increased. Effect of increasing voltage could be that was increased the density of charge in polymer solution, then the Taylor cone was formed, stretched and formation of smaller fibres due to the higher repulsive forces. Moreover, the more increasing of voltage, the more electric field strengths between needle and ground collector also increased. The polymer jet could fly to collector in short time that results in increasing of bead formation and/or larger fiber diameters [5,7,12,19].

![Fig. 5 SEM images (10,000x) of electrospun nanofibres and diameters distribution at different applied voltage (A) 20kV (B) 23kV and (C) 26kV](image)

![Fig. 6 The relation between the fibre diameters and applied voltage](image)

Effect of flow rate
The average fibres diameter at flow rates of 11.67 µL/min, 16.67 µL/min, and 25.00 µL/min were 152.32±41.48, 154.21±45.40 and 165.68±54.75 nm respectively. As show as Fig. 7 and 8. Smooth, non-bead and continuous fibres were obtained.

![Fig. 7 SEM images (10,000x) of electrospun nanofibres and diameter distribution at the different flow rate 3(A) 11.67µL/min (B) 16.67 µL/min and (C) 25.00 µL/min](image)

The flow rate influence to the initiate droplet shape. At high flow rates, bigger droplet was formed, and the flying ability across electrical field was decreased. The
thick fibres were formed on the collector. Therefore, the lower flow rate was more recommend [8,12,21].

In this research, diameters at flow rates 11.67 μL/min, and 16.67 μL/min were different without significant (p>0.05).

**Effect of distance**
The distance showed a direct effect on the jet flying time and solvent evaporation time. At the short distance, bead formation, thick and wet fibres could be obtained, whereas at the longer, small bead, smooth and continuous fibres with fibres diameters decreasing could be obtained owing to the flying time of jet and completely evaporated [2,12,19]. The average diameter of electrospun Chitosan-CA-Gelatin fibres at distance 8 cm, 10 cm, and 12 cm were 158±48.84, 152.32±41.48 and 146.74±40.22 nm respectively.

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It could be indicated that fibres diameters decreases when distance was increased (Fig. 9 and 10). Increasing the distance could give time for the fluid jet to stretch and for the solvent to evaporate completely [19,20].

**4 Conclusion**
Formation of Chitosan-CA-Gelatin was successfully obtained from the mixed of those at the ratio of 4:1:5. According to the study, it can be indicated that the morphology of Chitosan-CA-Gelatin was depended on polymer concentration, applied voltage, polymer’s flow rate and the distance between tip to collector. The optimum conditions were 18.8 % wt of polymer concentration, 23 kV, 11.67 μL/ min and 10 cm in distance.

This study was provided by the Thammasat University scholarship and National Research Council of Thailand. Scanning Electron Microscope was supported by TU-ARC.

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