Folic acid-containing nanofibers by simultaneous process for transdermal drug delivery: preparation, characterization, and in vitro biocompatibility

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Abstract

Nanofibers with bioactive agents are good candidates for skin-care applications due to high specific surface area, low density and highly porous structure. In this study, hydrophilic based bioactive nanofibers were produced via an electrospinning and electrospraying simultaneous process. Polyvinyl alcohol (PVA), polyvinyl alcohol-gelatin (PVA-Gel) and polyvinyl alcohol-alginate (PVA-Alg) polymers were used as the matrix material and folic acid (FA) particles were dispersed simultaneously on the surface of these hydrophilic nanofibers. The morphology of the nanofibers (NFs) was uniform and dispersed folic acid particles incorporated into the structure of nanofibers as confirmed by scanning electron microscopy (SEM). Thermal behavior, chemical structure of the composite nanofibers were analyzed/investigated by thermogravimetric analysis (TGA) and Attenuated Total Reflectance-Fourier Transform Infrared Spectroscopy (ATR-FTIR) which showed that no chemical bonding between vitamin and polymers. A controlled release of FA-loaded electrospun fibers were carried out by UV-Vis in vitro study within the 8 hour-period in artificial sweat solutions (acidic media, pH 5.44). The obtained PVA/FA, PVA-Gel/FA and PVA-Alg/FA fibers released 49.6%, 69.55% and 50.88% of the sprayed FA in 8 h, indicating the influence of polymer matrix and polymer-drug interactions, on its release from the polymer matrix. Moreover, biocompatibility of all developed novel NFs was assessed by two different cytotoxicity tests, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay and neutral red uptake (NRU) assay in L929 cell lines. In all cases, it is clearly concluded that these new electrospun fibers had fast-release of the vitamin and the hybrid process is suitable for transdermal patch applications, especially for skin-care products. Moreover, it has been proposed nanofiber with folic acid as a patch may prevent the COVID-19. The results of cytotoxicity assays on L929 cell reveal that all prepared NFs have no or slight cell toxicity. PVA and PVA-Gel with/without FA nanofibers seems more biocompatible than PVA-Alg nanofibers.

1. Introduction

Human skin, the largest organ in the body, has vital functional role by protecting from pollution, ultraviolet radiation and other damage, which cause to skin aging and disorders [1-3]. Skin has well-organized morphological structure and self-renewing barrier property. It comprises of three multilayers structure with different levels of cellular and epidermal differentiations that serves as a protective barrier [4,5]. The outmost layer of the skin is epidermis known as the stratum corneum (SC) contains dead cells (corneocytes) interdispersed within a greasy matrix [1,2,6]. Therefore, SC provides selective permeability and limits hydrophilic rich compounds to penetrate in to the inner layers. Transportation process of many bioactive materials in skin is mainly resulted by passive diffusion [7] in related to Fick's Law [8]. (Ficks Law summarized the absorption model in skin)

Transdermal drug delivery system (TDDS) is defined as pharmaceutical ingredients are transported locally or systemically via skin [1,9,10]. The approach is alternative to oral, intravascular, subcutaneous, and transmucosal routes due to loss of effectiveness and toxicity of drugs (degradation) [11,12]. Besides, TDDS have not cause painful administration and other unwanted side effects on the gastrointestinal tract compared with oral or injection administration, especially. Therefore, this route has not damage to
liver. However, the penetration of drugs via SC has many limitations. It is known that drug molecules penetrate/pass through/ via SC by two routes including intercellular (hydrophilic drugs) and intracellular (hydrophobic drugs) mechanisms. A drug molecule needs some physicochemical properties such as sufficient hydrophobicity, high and sufficient partition coefficient (Log P 1-3), low molecular weight (<500 Da) and the short half life for better penetration [1,13,14].

During the last decades, nanofibers have become excellent candidate for transdermal drug release applications due to high porosity, large specific surface area and small size of electrospun fibers [10,15]. There have been many attempts about pharmaceutical ingredients incorporated into nanofibers for usage of transdermal applications. In this regard, Galkina et al. (2013) reported cellulose NFs – titania nanocomposites grafted with three different drugs can be potential transdermal patch [16]. Gencturk et al. (2017) fabricated polyurethane/hydroxypropyl cellulose electrospun nanofibers with Donepezil (DNP) hydrochloride and in their study, controlled release of DNP from the fibers within 6 h was observed [17]. With the similar strategy, Esenturk et al. (2020) have investigated in vitro drug release behavior of voriconazole (VCZ) loaded hydrophilic PVA/SA electrospun NFs [18]. The obtained NFs indicated sustainable release behavior with 8 h. Also in a similar study reported by Shen et al. (2014) that prazosin Hydrochloride-loaded Poly(vinyl alcohol) fibres were prepared by conventional electrospinning, the resulting materials released drug with sustain behavior over the 24 h [19]. Kataria et al. (2014) developed transdermal patch based on ciprofloxacin loaded poly vinyl alcohol (PVA) and sodium alginate (NaAlg) electrospun composite nanofiber for wound healing [20]. These NFs showed cumulative drug release reached to about 99 % after 24 h. In another study, Tran et al. (2015) investigated release behavior of Ibuprofen from NFs. PCL, pNIPAM and pNIPAM/PCL were used as carriers of the drug for thermo-responsive transdermal delivery systems [21]. The release behavior of the fibers significantly changed with 4-hour period depending on the temperature. Song et al (2016) developed an alternative patch for the use of Daidzein in oral delivery applications [22]. The Daidzein-loaded PLGA fibers showed around 65.38% with sustained released behavior after 72 h.

Furthermore, vitamin loaded NFs have received increasing attention as transdermal patches. Within this aim, Madhaiyan et al. (2013) synthesized vitamin B12-loaded polycaprolactone NFs and the resulting fibers indicated gradually vitamin release behavior in 48-hour period [23]. A slow release of the vitamin E was observed from silk fibroin NFs over 72 h to utilize for skin care applications [24]. In another study, electrospun fibers made of gelatin was loaded with the vitamin A and E and investigated as wound healing patch [25]. The in vitro test results displayed the fibers have sustained release behavior for more than 60 hours.

Previous studies mainly focused on drug-loaded NFs fabricated by conventional electrospinning. Apart from electrospinning, there is one approach used in drug release applications which is called electrospraying. Electrospraying is sister technology of electrospinning to fabricate nanomaterials with < 1 μm diameter [26]. In electrospinning, polymer solution concentration is sufficient to develop chain formation in the capillary by using high voltage potential while the polymer solution concentration is too low and droplets is spraying from the capillary in electrospraying [27]. However, nanoparticles
micro/nanoparticle dispersions are processed with electrospraying due to low viscosity. There are many studies on nanofibers produced together with electrospinning and electrospraying process. However, to the best of our knowledge, there is no report on the transdermal drug delivery of nanofibers produced with simultaneous electrospinning and electrospraying process.

In the current study, electrospinning and electrospraying were performed simultaneously to fabricate the electrospun PVA, PVA-Gel and PVA-Alg fibrous mats. Folic acid utilized as model drug. By varying the polymer matrix type, the properties of as-spun/FA NFs were investigated in terms of morphology, thermal and chemical structure (polymer-polymer and polymer-drug compatibility). In vitro drug release from the NFs was assessed. The biocompatibility of the nanofibers was also evaluated through by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) and neutral red uptake (NRU) assay the L929 cell lines (mouse fibroblasts) as recommended in ISO10993-5 standard.

2. Materials And Methods

2.1. Materials

The Polyvinyl alcohol (PVA) (density 0.4-0.6 g/cm³, purity 87.8 %) was purchased from ZAG Chemical (Turkey). Gelatine from bovine skin (225 Bloom, type B), Alginate and sodium bicarbonate (NaHCO₃) (99.7 % purity) and acetic acid (CH₃COOH) with 80 % purity obtained from Sigma Aldrich Chemical Company (USA) and ethanol (99.9% purity, Tekkim). Folic acid (C₁₉H₁₉N₇O₆) for Biochemistry (98-102 % purity) was supplied from ChemSolute Company (Germany). To prepare the artificial sweat solutions, L-Histidine monohydrochloride monohydrate (C₆H₉O₂N₃.HCl.H₂O) (>99% purity), sodium chloride (>99.5 purity) and sodium dihydrogen phosphate dihydrate (NaH₂PO₄.2H₂O) were purchased from Sigma Aldrich Chemical Company (USA). The chemicals used in the MTT and NRU assay tests: RPMI (Roswell Park Memorial Institute) 1640 medium, fetal bovine serum (FBS), penicillin-streptomycin, neutral red (NR) (3-amino-7-dimethyl-amino-2- methylphenazine hydrochloride), 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), trypan blue, ethanol from Sigma; dimethyl sulfoxide (DMSO) from Merck; Triton X-100, trypsin–EDTA, Dulbecco's phosphate buffered saline (DPBS) from Gibco. Distilled water was used in the experiments and all of the reagents were used without any purification.

2.2. Fabrication of Hybrid Nanofibers

The hydrophilic based polymer/FA nanofibers were fabricated by combining electrospinning and electrospraying process (INOVENSO Nanospinner24, Turkey). PVA powders were dissolved in pure water at 90°C till a 10 % (w/w) homogeneous PVA solution was obtained. Gelatin was dissolved in pure water and acetic acid binary-solvent systems (7:3 w/w) solution to obtain a 20 % (w/w) gelatine solution by stirring for 2 hours at ambient conditions. Alginate powders were also dissolved in water to obtain a 2 % (w/w) alginate solution. PVA-alginate (4:1 v/v) and PVA-gelatin solutions (3:2 v/v) were mixed at room temperature for overnight. In our previous experiments, it was determined optimum folic acid (FA) concentration in electrospraying system. 22 mg FA was dispersed in 10 ml (2:1 v/v) pure water and
alcohol solution for each polymer system via ultrasonic homogenizer (Bandelin/Sonoplus HD3200). Each polymer solution (PVA, PVA-gelatin and PVA-alginate) and FA particle dispersion were transferred into two separate plastic syringes (10 mL) and put side to side on two syringe pumps. The plastic syringes were attached to two stainless steel nozzle. The PVA, PVA-gelatin and PVA-alginate solutions and FA dispersion were dispensed via nozzle which was vertical to the collecting plate. The feed rates were performed at 2 mL/h and 1.5 mL/h for polymers and FA dispersion, respectively. Neat PVA, PVA-gelatine and PVA-alginate solutions were prepared as a control (Fig. 1).

2.3. Nanofiber Characterization

2.3.1. Scanning electron microscopy (SEM)

Morphology of all nanofibers was observed with Carl Zeiss/Gemini 300 Scanning Electron Microscope (SEM) (ZEISS Ltd., Germany). All samples were coated with gold for 20 minutes before analysis. Fiber diameters were measured using Image J, version 1.520 software.

2.3.2. Thermal Analysis (TGA)

TA /SDT650 TGA(USA) were used for the thermal analysis. TGA analyses were performed under nitrogen atmosphere with 20°C min⁻¹ heating rate and 30°C - 600°C temperature range and applied oxygen atmosphere with 20°C min⁻¹ heating rate 600°C-900°C temperature.

2.3.3. Fourier transform infrared spectroscopy (FT-IR)

FTIR data were obtained with a Thermo Nicolet iS50 FTIR (USA) spectrometer with a ATR (Attenuated Total Reflectance) adapter (Pike, USA) in the range of 4000-500 cm⁻¹ recorded with 16 scans at 4 cm⁻¹ resolutions.

2.3.4. In vitro Release Study

The pH of skin surface ranges from 4.2 and 5.6 [28]. Furthermore, solubility of folic acid as a drug is maximum at pH 5 to 6 [29]. Therefore, artificial sweat solutions were prepared according to ISO 105-E04:2013 method [30]. The vitamin-release behavior of FA sprayed resulting fibers were studied in acidic sweat solutions at pH 5.44 by total immersion method [23]. 12-30 cm² of the nanofibers were put into sealed glass tubes with each containing 50 mL of acidic sweat solution, separately. Then, they were placed in shaking incubator at 37.5 °C with stirring 120 rpm in order to apply the release profile of the folic acid. Samples of 3 ml was removed at the specified time intervals with the sweat solution and the corresponding absorbance value was determined in a UV spectrophotometer (Scinco/NEOYSY 2000) at λ max = 282 nm, which was the characteristic peak of folic acid. The drug concentration was obtained from the calibration curve of the model vitamin prepared with a folic acid solution of known concentrations in acidic solutions (pH 5.44). The calibration curve was found to be Y = 0,0486X + (-0,0402) (R² = 0,99992), where X is the concentration of FA (mg/L) and Y is the solution absorbance at
282 nm (linear range of 0.5–25 mg/L) (Fig. 2). The amount of released drug was determined using UV–
Vis spectroscopy.

### 2.3.5. In vitro biocompatibility

**Preparation of nanofiber extract solutions for cytotoxicity assays**

In this study in order to determine the cytotoxic effects of novel NFs prepared in different compositions, extracts of all samples were prepared among the methods recommended by the UNI EN ISO 10993-12:2009 regulation [31] and also suitable for the nature and shape of NFs as biomaterials. Before performing the extraction procedure, equal sizes of NF patch samples (3 cm²) were cut and each sample was sterilized by UV light for 1 hour in order to keep the structural properties of the nanofibers intact. The extraction procedure was carried out in sterile tubes containing 5ml of culture medium (99% RPMI 1640 and 1% Penicillin-Streptomycin, without serum in order to prevent protein interaction). All samples were kept in this medium at 37 °C for 30 minutes and it was observed that they were completely dissolved during this time period. Equal sample of an aluminum foil was also used simultaneously in the experiments as a reference base support material of NFs and treated as a sample and also extraction medium without sample was used as control and culture medium containing 1% Triton X-100 was used as a positive control.

**Cell culture**

All experimental parts of cell culture and cytotoxicity assays were conducted at the cell culture laboratory of Bioengineering Department of Bursa Technical University according to the good cell culture practices - Coecke et al. (2005) [32]. The L929 cell lines (mouse fibroblasts) were used for cytotoxicity assays as a reference cell line for cytotoxicity testing of medical devices and materials, according to the UNI EN ISO 10993-5 rule:2009 [33]. L929 cells (kindly obtained from Yeditepe University, Department of Genetics and Bioengineering) were seeded in 75cm² culture flask containing RPMI 1640 supplemented with 10% FBS and 1% penicillin streptomycin. Cells were grown in an incubator at 37°C in an atmosphere supplemented with 5% CO₂ and monitored daily by using an inverted microscope with phase contrast attachment (Olympus CKX41). Subcultures were performed when an 80% of confluence was observed. Following disaggregation with trypsin/EDTA and resuspension of cells in medium, a total of 5x10⁴ cells/well were plated in 96 well tissue-culture plates.

**Determination of the cytotoxicity of nanofibers by 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay**

MTT assay was performed by the method of Mosmann (1983) [34] with the modifications of Hansen et al. (1989) [35] and Kuz’ma et al. (2012) [36]. After 24 h incubation of cells seeded in 96 well tissue-culture plates, cells were exposed to the different concentrations of nanofiber extract solutions in medium for 24 h at 37 °C in 5% CO₂ in air. After exposure, the medium was aspirated and MTT (5 mg/ml of stock in PBS) was added (10 μl/well in 100 μl of cell suspension), and cells were incubated for an additional 4 h
with MTT dye. At the end of incubation period, the dye was carefully taken out and 100 µl of DMSO was added to each well. The absorbance of the solution in each well was measured in a microplate reader at 570 nm. Results were expressed as the mean percentage of cell growth from three independent experiments.

**Determination of the cytotoxicity of nanofibers by neutral red uptake (NRU) assay**

The cytotoxicity of nanofibers was performed in L929 cells by NRU assay following the protocols described by Virgilio et al. (2004) [37] and Saquib et al. (2012) [38]. After 24 h incubation of L929 cells in 96 well plates, 200 µl of all sample solutions were added. The cells were incubated for 24 h at 37 °C in 5% CO2 in air, then the medium was aspirated. The cells were washed twice with PBS and incubated for an additional 3 h in the medium supplemented with NR (50 µg/ml). After the medium was discarded, the cells were rinsed five times with warm PBS (pH 7.4) to remove the non incorporated excess dye and 200 µl of destain solution (50% ethanol, 1% acetic acid, and 49% distilled water) was added to each well to fix the cells and bring the NR into solution. The plates were shaken for 20 min, and the absorbance of the solution in each well was measured in a microplate reader at 540 nm and compared with wells containing untreated cells. Results were expressed as the mean percentage of cell growth from three independent experiments.

3. Results And Discussion

3.1. Morphology of electrospun nanofibers

The morphology of the resulting electrospun fibers was investigated with SEM and presented in Fig. 3, respectively. When comparing with SEM images of PVA and PVA/FA fibers, neat PVA sample has intensive beads on a string structure in the fiber (Fig. 3A&B). However, applying to electrospray procedure of FA on the fibers enhanced the formation of beadless morphology. With the adding of FA, the average diameter of fiber with beads decreased from about 532 nm (neat PVA) to about 291 nm (PVA/FA) (Fig. 4A&B). Folic acid molecules connected PVA fibers showed a little rough, randomly interconnected structure with FA clusters formed. Unfortunately, folic acid clusters were not seen too much on surface of the PVA fibers and this situation can be explained by using the same solvent in both electrohydrodynamic process and thus folic acid molecules attached into PVA fibers. The addition of gelatin into PVA solution also provided thin and homogenous fiber structure (Fig. 3C). PVA/Gel fibers with thin and regular structure were also obtained previously by Lihn et al. [39]. Furthermore, PVA-Gel NFs had a predominant fiber diameter of 80-120 nm; whereas, PVA-Gel/FA fibers had a fairly even distribution ranging from 60 to 220 nm. (Fig. 4 C&D). Folic acid molecules caused to form of thicker fiber structure and destroyed the homogenous fiber morphology. There were 2D flat fiber forms in some regions. In addition, folic acid clusters on the gelatin fibers were seen clearly. Fig. 3E displays the blend of PVA and alginate fibers. The diameter of fiber increases by adding alginate into PVA solution. Therefore the resulting blend sample has shown slightly interconnected fiber morphology. Although there are FA beads on the PVA-alginate fibers, the structure destroyed seriously during electrospraying of FA. The reason of this is similar to
PVA/FA fibers related to same solvent for two hydrodynamic process. As shown in Fig. 4E and F, nano-scale PVA-Alg and PVA-Alg/FA fibers were produced with diameters ranging from 150 to 550 nm and 50 to 350 nm, respectively.

As a result, SEM images revealed FA clusters have both partially deposited on the surfaces of hydrophilic fibers and integrated into the fiber structure. Moreover, morphology PVA fibers with beads enhanced with their blends.

3.2. Thermal Properties

Fig. 5 illustrates TGA and DTG thermograms for neat PVA, PVA blends and FA loaded electrospun PVA fibers. All electrospun fibers show weight loss in the range of 30-100°C related to moisture out of fibers. The amount of weight loss was different for all fibers. The maximum weight loss was seen at the PVA-Gel/FA mixture and the minimum was at the PVA/FA mixture. The amount of the weight losses were 9.28 % and 4.15 % for PVA-Gel/FA and PVA/FA , respectively. Apart from the weight loss related to moisture out three steps of weight loss were observed for the neat PVA fiber. The major weight losses of 43% taken place in the temperature range from 210°C to 350°C due to dehydration of PVA [40,41]. The second step is dominated by chain scission about 450°C to 600°C [42] and the amount of weight loss was 8.18 %. The last step was related to burning of the pyrolysis product which was formed during analysis in the N\textsubscript{2} atmosphere. The weight loss of 3.77 % from the last step. It is found that FA loaded PVA fibers started to decompose at about 150°C with 45% weight loss and continue to 300°C. The following weight loss of 13% from 350°C to 460°C was attributed to both elimination reaction of PVA and loss of pterin and then p-aminobenzoic acid units in folic acid. The last step was related to burning of the pyrolysis product which was formed during analysis in the N\textsubscript{2} atmosphere. The amount of weight loss was 8.83 % in this step.

The first weight loss of almost 67 % apart from moisture out was seen between 280-450°C for PVA-Gel fibers. The second weight loss in the amount of 6.49 % was occurred in the region of 450-550 °C. The last weight loss was about 11.68 % related to pyrolysis product which was formed during analysis in the N\textsubscript{2} atmosphere. The weight loss of blend fiber had a decrease at 280°C and continue to 500°C with 78% weight loss which was mainly due to the cleavage of C=N bond is concerned with the presence of protein molecules in gelatin [43].

Thermal behavior of the PVA-Gel/FA was slightly different from the thermogram of the PVA-Gel. The difference was only at the region of the 150-250°C temperature. There was slightly weight loss of the PVA-Gel/FA (about 3.34 %). The other difference was occurred after the temperature of the 450°C due to the FA content. The decomposition speed was decrease as a consequence of C=N group.

The lowest decomposition temperature was occurred for PVA/Alg blend nanofibers about 170°C [44]. The first decomposition weight loss of 68 % from the fiber is related to dehydration of PVA. The second decomposition step temperature range of about 350-450°C, corresponding to the degradation of alginate with weight loss of 7.52. The third decomposition step the temperature ranges of about 450-600°C with
weight loss of 6.21 % from the fiber was related to PVA chain scission as the other PVA samples. The loss of the moisture from PVA-Alg/FA was less than the PVA-Alg. Furthermore, as seen in Fig. 5, PVA-Alg fibers showed faster decomposition than PVA-Alg/FA after 450°C. There is no residue for all NFs.

3.3. FT-IR Analysis

The chemical composition of the all electrospun fibers and folic acid were determined through FT-IR analysis. As seen in Fig. 6, the spectrum of folic acid has a number of characteristic peaks at 3590, 3496, 3330, 2925, 2840, 1694, 1650, 1605, 1487 and 1405 cm⁻¹. The band between 3600-3300 cm⁻¹ is associated with (-OH) stretching bands of glutamic acid moiety and -NH group of pterin ring. The band at 1650 cm⁻¹ belongs to the (-C=O) stretching of (-CONH₂). The another characteristic IR absorption peaks at 1605, 1694 and 1487 cm⁻¹ is due to the N-H bending vibration of (-CONH) group, (-C=O) amide stretching of the a-carboxyl group and absorption band of phenyl ring respectively [45-47]. The frequencies for the neat PVA are indicated as follows: 3284 cm⁻¹ for the stretching vibration peak of its (-OH) groups, 2943 and 2910 cm⁻¹ for the stretching vibration of -CH₂ group, 1242, 1088, 1023 and 945 cm⁻¹ (C=O) stretching vibration, respectively [41,48,49]. On the other hand, for the PVA-Gel fibers the band appears at 3290 cm⁻¹ for the (-OH) group. The bands appear at 2938 and 2913 cm⁻¹ belongs to -CH₂ stretching vibration. The characteristic absorption of gelatin peaks show mainly to the peptide bonds (-CONH) with the amide I–III vibrations. The peak is 1643 cm⁻¹ (amide-I) is related to -C=O stretching vibration whereas the peak at 1535 cm⁻¹ (amide-II) is due to N-H bending and C-H stretching vibration. 1243 cm⁻¹ for the (amide-III) peak was occurred. In addition, 1435 cm⁻¹ (-CH₂ bending), 1374 cm⁻¹ (C-H wagging), 1088 cm⁻¹ (-C-O-C) and 837 cm⁻¹ (C-C) stretching [50,51].

In literature, the characteristic peaks of alginate assigned 3297 cm⁻¹ (-OH stretching), 2940 cm⁻¹ (-CH₂ stretching), 1612 and 1424 cm⁻¹ asymmetric and symmetric (–COO) stretching and also 1096 cm⁻¹ (C-O) stretching vibrations, respectively [52]. The electrospun PVA-Alg and PVA fibers were same absorption peaks approximately. However, the absorption peaks at 3284 cm⁻¹ (-OH stretching) belongs to PVA that shifted to the 3297 cm⁻¹ [53,54].

It is worth mentioning that the spectrums of NFs with drug (Fig. 6) were similar the spectrum to neat NFs, indicating the absence of any chemical reaction between polymer and drug. This is explained by -OH groups in polymer structures shielded the characteristic peaks of drug molecules between 3300-3600 cm⁻¹, corresponding to penetration of drug into the fibers. Additionally, the characteristic peaks of drug molecules could not seen after 1700 cm⁻¹ due to the low intensity.

3.4. UV-Vis Spectroscopy- in vitro Study

UV-Vis spectrometry was used to evaluate the release profile of FA from the nanofibers, which helped to reveal the structure-function relationship of the electrospun FA-loaded fibers in artificial sweat solution
(pH 5.44, 37°C) during the time course of 480 min. The cumulative FA release rate profiles, in the sweat solution FA from the fiber samples, are plotted in Fig. 7.

The release profile shows that the percentage cumulative FA releases are 49.6 %, 69.55 % and 50.88 % for PVA/FA, PVA-Gel/FA and PVA-Alg/FA, respectively after 8 h. As seen in Fig. 7, all fibers exhibited initial burst release profile at an early stage of the analysis. Especially, in PVA/FA fiber at an early stage of the analysis within first 5 min, more than half of the FA was released from the fiber. In PVA-Gel/FA fiber within first 30 min about 65 % FA release whereas in PVA-Alg/FA composite fibers 49 % FA release. The rapid release of FA from the composite fibers is related to many factors, such as hydrophilic nature of matrix (PVA, gel and alg), high surface area of fibers that supports to increase in wettability [55,56]. Illangakoon et al. have reported hydrophilic PVP with paracetamol/caffeine NFs release almost all caffeine and paracetamol within the first 6 min [57]. In this study, the burst release behavior of all fibers might be related to the existence of hydrophilic based polymer matrix and drug as well. Furthermore, drug molecules deposited on the fiber surface also cause the initial release due to the hybrid process. On the other hand, burst release is a phenomenon which may be preferred to obtain quick results in dermal applications [58]. Kataria et al. also prepared ciprofloxacin loaded transdermal patches with fast release. In this study, the resulting PVA and PVA-Alg with ciprofloxacin fibers have sustained and controlled release profile and reached maximum drug release curve in the first 7 hours [20].

It is clearly seen PVA-Gel/FA had the slowest behavior in fibers for 30 min. From the SEM images, the average fiber diameter of PVA/FA, PVA-Gel/FA and PVA-Alg/FA are 291, 145 and 158 nm. Moreover, the sample has the highest drug release rate compared with the other samples in total time. This is explained as the thickness of the fibers increase, the pathway drug diffusion of will increase [59].

The maximum drug was released from the fibers within sustained behaviour until 45 min. From the Fig. 7 showed diffusion based drug release occurred for 30 min, then followed by a constant drug release till 8 h due to erosion of polymer matrix. Similar UV-Vis results were reported by Arthani et al. [53]. However, the release rate in PVA-Gel/FA fiber is partially slower than others for the first 30 min. due to the interaction hydrogen bonding between PVA, gelatin and folic acid. PVA-Alg/FA fiber also indicated similar situation, but unfortunately the low amount of alginate in the fiber limit the formation of too many interaction of hydrogen bonding (Fig. 6). Consequently, PVA-Gel/FA could be used as an efficient drug delivery system for beauty mask purpose with 8 h period.

Recently, there have been a variety approaches about COVID-19 pandemia. In this regard, inhibition of furin enzyme activity provide to limit of viral and bacterial growth [60]. Sheybani et al. have claimed that folic acid prevents furin activation. It has been simulated foid acid could be a potential drug in early respiratory diseases caused by COVID-19[61]. Results of the study indicates folic acid molecules interacted with active sites of furin due to formation of hydrogen bonds. Therefore, we also proposed nanofiber with folic acid as a patch may prevent the COVID-19.

3.5. Cytotoxic effects of nanofibers by MTT assay
The cytotoxic effects of PVA/FA, PVA-Gel/FA and PVA-Alg/FA nanofibers extract solutions were evaluated by MTT assay in L929 cells. To determine and compare the cytotoxicity, extract solutions of all NFs and also a negative control group without any chemicals, aluminum foil as a planch material and a positive control group containing 1% Triton x-100 were prepared. Eight wells of each sample were repeated in 96 well plates. The average absorbance values and standard deviation values of living cells were calculated by averaging all the data obtained. In addition, cell viability in the control group was considered as 100%, and living cell percentages were determined for all samples compared to the control. Results of MTT test are shown in Fig. 8. According to the results of MTT test applied after 24 h of treatment in L929 cells, the groups treated with nanofiber extracts caused some decreases in cell viability in all groups compared to the negative control group. However, since viability did not fall below 70%, it was not evaluated as cytotoxic by MTT assay.

3.6. Cytotoxic effects of nanofibers by NRU assay

The cytotoxic effect of the extract solutions of prepared NFs were evaluated by NRU assay in L929 cells. Results of NRU test are shown in Fig. 9. According to the results of NRU test applied after 24 h of treatment in L929 cells, the groups treated with nanofiber extracts caused decreases in cell viability in all groups compared to the negative control group. Except the PVA-Gel NF cell viability decreased under 70%, but not decreased under 50%. PVA-Alg NFs caused more cytotoxicity in NRU assay. On the other hand, L929 cells were treated with a high concentrations of nanofiber extracts in this study to investigate exact cytotoxicity.

In the NRU method, which is based on the measurement principle of lysosomal activity, cell viability is shown by a different mechanism than MTT assay (based on the measurement of mitochondrial activity as metabolic activity). It is thought that the nanofiber extract solutions may cause toxic effects by interacting with the cell membrane or enter the cell through the membrane and accumulating in organelles and cytoplasm.

Conclusions

In this study, we successfully produced fast-dissolving drug delivery systems derived from hydrophilic composite fibers by simultaneous electrospraying and electrospinning method. FA was used as model drug and the resulting composite fibers consists of FA clusters entrapped in PVA, PVA-Gel and PVA-Alg NFs. The SEM images showed the composite fibers possessed relatively uniform with average fiber diameters between 145-291 nm and most of the FA clusters have been integrated into the fibers rather than deposit on the fiber surface due to polymers and folic acid solutions are hydrophilic based nature. Moreover, as FA is added fiber morphology is formed with bead-free for PVA/FA fiber. The FTIR spectra results demonstrated the apparent slight shifts of some peaks, corresponding to physical interactions of FA with fibers. The TGA results pointed out that by the incorporation of FA into NFs, the degradation rate was increased slightly. However, FA was stable at high temperature. The in vitro release test clearly confirmed that the obtained PVA/FA, PVA-Gel/FA and PVA-Alg/FA NFs could release the FA in a sustained
manner with initial burst release for the 8 hour-period. The observation of fast dissolving of all fibrous structures with FA in 30 min is directly related to two reasons: the strong hydrophilic nature of PVA, PVA-Gel and PVA-Alg NFs and FA clusters deposited on fiber surfaces due to electrospraying process.

The biocompatibility test results based on the cytotoxicity methods adapted from the ISO10993-5 standards indicated that PVA/Gel/FA, PVA-Alg/FA electrospun nanofibrous patches revealed no cell toxicity on cultured fibroblasts in MTT assay but slight cell toxicity in NRU assay. In this study, L929 cells were treated with high concentrations of nanofiber extracts to investigate exact cytotoxicity due to the nature of certain biodegradable/leachable and extractable biomaterials. As indicated in biocompatibility test protocols the quantity exposed to used cell lines is dependent to the interface area, the volume of extraction, pH, temperature, time and many other factors. Results of this study and similar studies in the literature also indicated that the cytotoxic effect the different PVA nanofibers are based on the components and content amounts. Our results reveal that PVA and PVA-Gel with/without FA nanofibers seems more biocompatible than PVA-Alg nanofibers. However, further biocompatibility tests should be carried out in different conditions, concentrations and different cell lines according to potential use for dermal or other biomedical applications as scaffolds. Moreover, the new nanofibers can also be used as an effective patch against the COVID-19 thanks to folic acid molecules.

**Declarations**

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**Conflict of interest**

The authors declare no conflict of interest.

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Figures

Figure 1

Schematic illustration of hybrid process of nanofibers.
Figure 2

Calibration curve of FA at maximum 282 nm in acidic sweat solution pH 5.44.
Figure 3

SEM micrographs of A.PVA, B.PVA/FA, C.PVA-Gel, D.PVA-Gel/FA, E.PVA-Alg and F.PVA-Alg/FA nanofibers. (3kX).
Figure 4

The average fiber diameters of A. PVA, B. PVA/FA, C. PVA-Gel, D. PVA-Gel/FA, E. PVA-Alg and F. PVA-Alg/FA fibers.
Figure 5

TGA and DTG thermograms of nanofibers.
Figure 6

FT-IR spectra of nanofibers.
Figure 7

FA release profiles of nanofibers.
Figure 8

Effects of PVA nanofiber extract solutions on cell viability of L929 cells by MTT assay. Results were expressed as the mean percentage of cell growth inhibition from 3 independent experiments. Cell viability was plotted as percent of control (assuming data obtained from the absence of nanofiber as 100%).
Figure 9

Effects of PVA nanofiber extract solutions on cell viability of L929 cells by NRU assay. Results were expressed as the mean percentage of cell growth inhibition from 3 independent experiments. Cell viability was plotted as percent of control (assuming data obtained from the absence of nanofiber as 100%).