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Monitoring drug release from electrospun fibers using an in situ fiber-optics system

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INTRODUCTION

Electrospinning is a highly versatile technique that enables the production of ultrafine fibers based on polymeric solutions. In recent years, the interest in electrospinning has significantly increased as the technique provides promising applications in many scientific fields (1). A special interest has been directed towards applications of electrospun systems for drug delivery (2). Electrospun fibers provide unique characteristics such as high surface-to-volume ratio, and depending on the polymer, tailor-made release kinetics for embedded drugs (2, 3). Additionally, electrospinning provides the ability to incorporate different types of active pharmaceutical ingredients varying in origin, physicochemical properties, and molecular size (4). Conventional dissolution experiments for drug-loaded electrospun fibers are conducted in small vials by immersing a preweighed sample in dissolution medium at physiological pH and temperature. At predetermined time points, defined aliquots are then withdrawn for drug quantification and replaced with fresh dissolution medium (5). However, this dissolution testing procedure is associated with several drawbacks. First, as the method is not automated, manual sampling is required. This creates limitations in terms of the possible overall duration of such experiments, and the accuracy of data obtained from frequent sampling as required for the analysis of rapid drug release processes. Furthermore, as the fiber mats are flexible sheet-like structures, irregular partial folding or creasing of the mats tends to occur when introduced to the aqueous medium. This change of conformation may, in turn, affect release kinetics and, potentially, the reproducibility of acquired data.

As an alternative to conventional dissolution testing involving the withdrawal of medium aliquots for drug ...
quantification, systems based on optical fibers were introduced allowing for continuous in situ sampling directly in the dissolution medium (6). Such systems allow for the direct UV measurement of dissolved drug within samples, and therefore a real-time dissolution profile can be obtained (6). The initial report regarding the application of such systems for the analysis of drug release from tablets was published by Josefson et al. in 1988 (7). Since then, technical progress has paved the way for standardized optical fiber probes with the opportunity for the concentration assessment of dissolved drug within several individual vessels (8, 9). The automation of this process facilitates reproducibility and convenience, while the ability to measure concentration accurately at high frequency allows for improved detection and a better understanding of release kinetics within short time frames. However, even though the suitability of such a method has already been proven, the assessment of release from thin flexible systems like fiber mats provides additional challenges. Due to their low weight and large surface area, the mats may remain suspended within or on the surface of the dissolution medium throughout the course of the experiment rather than sinking to the bottom of the dissolution vessel, thus distorting reliable data acquisition. Furthermore, the aforementioned tendency of such sheet-like structures to assume a partially folded conformation upon contact with liquids is prone to affect the release kinetics of mat-embedded drugs.

In this study, we investigated the suitability of a fully automated dissolution testing system based on fiber optics for in situ monitoring of drug release from electrospun fiber mats in comparison with cast films. To prevent folding of the fiber mats and films in the dissolution medium, we developed an adapter allowing for the attachment of the fiber mats and films to the wall of the dissolution vessels.

MATERIALS AND METHODS

Materials

Poly(vinyl alcohol) (PVA; Mw 85,000–124,000, >99% hydrolyzed) and lysozyme from chicken egg white (~70,000 U/mg) were purchased from Sigma–Aldrich (Steinheim, Germany). Acetic acid was obtained from Fluka (Buchs, Switzerland).

Methods

Fabrication of Electrospun Fiber Mats

PVA was dissolved in purified water (10% w/v) with acetic acid (1% v/v). After adding lysozyme (6% w/w drug/polymer), the solution was spun using an electrospinning setup consisting of a syringe pump system (New Era Pump Systems, Inc., USA) and a high voltage power supply (Acopian, USA). The polymeric solution was pumped through a syringe nozzle with a flow rate of 0.5 mL/h while a high voltage of 20 kV was applied. The formed electrospun fibers were deposited on a metal collector located at a distance of 14 cm from the nozzle. After electrospinning, the lysozyme-PVA fibers were stored in a desiccator for 24 h before further analysis.

Fabrication of Cast Films

Film-casting was performed by pipetting a defined amount of the lysozyme-PVA solution into a Petri dish, followed by homogenous distribution of the solution on the dish surface. The Petri dishes containing the lysozyme-PVA solutions were left at room temperature for at least 24 h to ensure complete solvent evaporation before being stored in the desiccator.

Fourier-Transform Infrared Spectroscopy

An attenuated total reflectance (ATR) unit was used to record Fourier-transform infrared (IR) spectroscopy spectra (Spectrometer 400 ATR–IR, Perkin Elmer, USA). Spectra of raw substances, cast films, and electrospun fibers were collected in the range of 550–4000 cm⁻¹. All spectra were recorded at least in triplicate.

Scanning Electron Microscopy

Samples of the electrospun fibers or cast films were mounted on aluminum stubs using double-sided carbon discs and sputtered with a thin layer of gold prior to analysis (Sputter Coater Q150R ES, Quorum Technologies Ltd., UK). The scanning electron microscopy (SEM) analysis was performed at an accelerating voltage of 5 kV using a Zeiss EVO HD 15 SEM (Carl Zeiss AG, Oberkochen, Germany).

In Vitro Drug Release

In vitro drug release experiments were performed using an in situ fiber-optic monitoring system, the µDISS Profiler, supported by Au PRO software (version 5.1.0.0, Pion Inc., Woburn, MA, USA). The fiber-optic probes were operated with tips of 20-mm path length. Prior to the experiments, each channel was calibrated with its own standard curve prepared from multiple additions of a fresh solution of lysozyme in phosphate buffered saline (PBS) buffer. Punches of electrospun fiber mats and cast films were prepared (16-mm diameter), and drug release experiments were carried out in 15 mL PBS pH 7.4 at 37 °C with magnetic stirring at 100 rpm. UV absorption (280 nm) was measured at a time interval of 10 sec for the first 5 min, followed by measurements at 30-sec intervals for the following 10 min. After this initial period, measurements were performed every 60 sec. Release experiments were performed at least in triplicate.
RESULTS AND DISCUSSION
Fabrication and Characterization of Electrospun Fibers and Cast Films
To investigate the applicability of an in situ fiber-optic monitoring system for testing drug release from electrospun fibers, lysozyme was used as a model drug and encapsulated within electrospun fibers based on PVA. For comparison, cast films based on the same solution used for electrospinning were prepared. Despite the identical composition of both polymeric systems, a clear difference in the visual appearance could be noticed. The electrospun fiber mats exhibited an opaque white color (Figure 1A) that can be attributed to the ultrafine substructure of the fiber network (Figure 1B) (10). In contrast, the cast films appeared transparent with a smooth surface (Figure 1C,D). Both samples were prepared as circular punches with the same diameter and therefore the same apparent surface area. However, the ultrafine substructure of the electrospun fibers (individual fiber diameters approximately 500 nm) increases the actual surface area of these samples, which can be expected to have an impact on the release kinetics of embedded drugs.

To elucidate any potential chemical interactions as a result of sample fabrication, ATR–IR spectra of pure compounds, fiber mats, and cast films were acquired. When comparing the ATR–IR spectra of the pure compounds to the spectra of the drug-loaded systems, neither the appearance of novel peaks nor the extinction of initial peaks (indicative of a chemical interaction as a result of sample preparation or instability) occurred (Figure 2). Further, comparison of the spectra of films and fiber mats does not reveal any differences due to the respective fabrication technique. In addition, the incorporation of lysozyme into the fibers as well as into the films can be detected by the amide I peak (1650 cm$^{-1}$).

Figure 1. (A) Surface morphology of the electrospun fibers, (B) SEM image of the electrospun fibers, (C) surface morphology of a cast film, (D) SEM image of a cast film.

Figure 2. ATR–IR spectra of pure compounds, electrospun fiber mats, and cast films.
Conventional Release Experiments with Electrospun Fibers

Conventional drug release experiments are performed by immersing the drug-loaded dosage form in buffer and quantifying the concentration of the released drug at predetermined time points. Although thin polymeric dosage forms such as electrospun fiber mats, films, or transdermal patches can potentially be investigated using this method, the thin and flexible structure of these dosage forms make them susceptible to folding upon immersion in the release medium as illustrated in Figure 3. This folding process is likely to decrease the surface area of the investigated sample, leading to inaccurate assessments of the release kinetics.

This phenomenon necessitates the preservation of the flat structure of the sample during the release experiment for improved accuracy and reproducibility. Indeed, dissolution testing for transdermal patches as described in the pharmacopeia (11) requires the fixation of the sample by using a strip of double-sided adhesive tape or an extraction cell. The same concept has been presented for polymeric films (12, 13). For electrospun fibers, however, drug release experiments are generally conducted by simply immersing the fiber mats in buffer without any fixation of the sample (5), even though there are few reports where the issue of sample folding was addressed. For instance, a modified Finn chamber was used for investigating drug release from electrospun fibers, hence keeping the sample from folding during the release experiments (14). In the present study, a novel flexible adapter was designed. The aim was to keep the sample fixed in a certain position during the release study, thus preventing migration and folding.

The designed adapter was composed of commercially available silicone and a nylon-based net (Figure 4). The mechanical flexibility of silicone allowed for convenient use, while the net provided a suitable support and, at the same time, unrestricted diffusion of the release medium and released drug. Prior to the release experiments, interactions of net material and drug were experimentally excluded (data not shown).

In Vitro Drug Release with the Fiber-Optic System

Release experiments in PBS pH 7.4 were performed with the electrospun fiber mats as well as with the cast films. The samples were mounted using the designed adapter, which assured holding the sample flat in position and prevented undesired sample folding (Figure 4). During the experiment, the dissolution vials were tightly sealed to prevent volume change due to medium evaporation. Each fiber-optic channel (corresponding to a separate dissolution vessel) was calibrated with its own standard curve prior to experimentation, and a very good linearity was achieved as shown in the representative calibration curve (Figure 5A). The theoretical drug content in the fiber mats and films was calculated based on the solid mass of lysozyme compared with the solid mass of the polymer. Thus, solvent evaporation upon electrospinning (or film casting) did not affect the lysozyme/polymer mass ratio. These calculations were successfully proofed by drug content analysis after destroying fiber mats and films. Complete drug release from the electrospun fibers as well as from the cast films was observed after approximately 20 min, which can be expected due to the hydrophilic nature of the drug and the polymeric fiber material. The results of the release experiments are shown in Figure 5B, and the data are represented as mean ± SD (n = 3).
The frequent sampling intervals provided a deeper insight into the release process from the electrospun fibers as well as from the cast films. In this context, we can see that lysozyme was released faster from the electrospun fibers than from the cast films (Figure 5B). This can be attributed to the higher surface area provided by the ultrafine fibrous structure, which facilitated the release of lysozyme into the dissolution medium. Although the overall difference in the release kinetics can be considered marginal, the identification of this occurrence could be achieved successfully.

CONCLUSION

Fully automated fiber-optic systems provide many advantages for the dissolution testing of electrospun fibers as well as from the cast films. In this context, we can see that lysozyme was released faster from the electrospun fibers than from the cast films (Figure 5B). This can be attributed to the higher surface area provided by the ultrafine fibrous structure, which facilitated the release of lysozyme into the dissolution medium. Although the overall difference in the release kinetics can be considered marginal, the identification of this occurrence could be achieved successfully.

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