Electrospun Polyurethane-Core and Gelatin-Shell Coaxial Fibre Coatings for Miniature Biosensors

Ning Wang, a Krishna Burugapalli, a Shavini Wijesuriya, a Mahshid Yazdi Far, b Wenhui Song, b Francis Moussy, a Yudong Zheng, c Yanxuan Ma, c Zhentao Wu, d Kang Li d

a Brunel Institute for Bioengineering, Brunel University, Uxbridge, London, UK
b Wolfson Centre for Materials Processing, Brunel University, Uxbridge, London, UK
c School of Materials Science & Engineering, University of Science and Technology Beijing, China
d Department of Chemical Engineering, Imperial College, London, UK

*Corresponding author. Tel. +441895266123, Fax. +441895269737, e-mail. wenhui.song@brunel.ac.uk

Running Title: Bioactive coaxial-fibre coatings for biosensors

Abstract

The aim of this study is to introduce bioactivity to the electrospun coating for implantable glucose biosensor. Coaxial fibre membranes having polyurethane (PU) as the core and gelatin (Ge) as the shell were produced using a range of PU concentration while keeping Ge concentration constant in 2,2,2-Trifluoroethanol (TFE) solutions. The core-shell structure, morphology, fibre diameter, pore size, porosity, thickness of the fibre membranes and their corresponding mechanical and surface properties, and glucose diffusion kinetics through the membranes were systematically investigated. The coaxial fibre membranes were electrospun on implantable glucose biosensors for evaluation of their effects on the long-term sensor function in vitro. It was found that the sensitivity and linearity of the sensors could be influenced by the composition, structure and morphology of the coated membranes. 2PU10Ge membranes with small fibre diameter (541 nm) and thick Ge shell (52%) did not affect the sensor sensitivity, but gradually decreased the linearity for the sensor over the time tested. In contrast, 6PU10Ge membranes with large fibre diameter (1133 nm) and thin Ge shell (34%) maintained the sensors’ linearity and stable sensitivity up to test period of 12 weeks and also acted as mass-transport limiting membranes, but significantly decreased the sensor sensitivity.

Keywords: Coaxial polyurethane/gelatin fibrous membrane, implantable biosensor coatings, electrospun coatings, glucose biosensor, glucose diffusion

1. Introduction
Reliable continuous monitoring of physiologically relevant molecules using implantable sensors is a longstanding challenge. As soon as the sensor is implanted in the body, it starts losing sensitivity, which downward drift caused by biofouling and fibrous encapsulation continues rapidly until the sensor fails. Many strategies involving surface modifications or deposition of additional polymeric coatings have been developed to combat the effects of biofouling, fibrous encapsulation and blood vessel regression [1]. Nevertheless, reliable extension of the in vivo sensing lifetime of implantable glucose biosensors is yet to be achieved.

In our research, we apply electrospinning technology towards overcoming the limitations of the traditional strategies to extend the in vivo sensing lifetime of implantable sensors. Recently, we demonstrated that efficacy of electrospun PU (commercial Selectophore™) coatings as mass-transport limiting membranes for miniature coil-type glucose biosensors [2, 3]. Compared to the traditional solvent cast membrane, electrospun PU membrane showed advantages in having tailorable thickness, structure and composition, as well as having minimal effect on sensor sensitivity and function. The membranes were tested for mass-transport limiting, and had subcellular porosity. Hence they are susceptible to biofouling and fibrous encapsulation. An additional tissue engineering layer would be essential to overcome this problem, which again can be achieved using electrospinning technology. Our strategy is to electrospin coaxial fibres having bioactive gelatin as the sheath and PU as the reinforcing core.

The synthetic polymer, PU has the desirable mechanical properties for implantable biomedical device applications, but is relatively inert. In contrast, extracellular matrix derivatives, collagen and gelatin (Ge) are bioactive, but lack the desired mechanical properties. Electrospinning provides an opportunity to produce co-axial fibres with core-shell structure made from different types of synthetic and natural materials that offer combined properties, such as bioactivity and mechanical strength. Recently, electrospun materials due to their higher similarity to native extracellular matrix (ECM) have been of particular interest for tissue engineering [4-7]. Polycaprolactone (PCL)-Ge core-shell electrospun fibres were produced to achieve comparable maximum elongation with PCL [4]. Likewise, PCL-collagen coaxial fibres were reported to have the advantage of resembling the natural ECM compared to the rough collagen coating on the pristine PCL through encouraging cell-matrix interaction [5]. C-PLGA/chitosan membranes were found to have potential in controlling drug release and skin restoration [6]. Similarly, collagen functionalized thermoplastic polyurethane (PU) scaffolds were also developed for soft tissue engineering application [7]. Therefore, it is hypothesised that introduction of a thin layer of natural biopolymer, e.g. gelatin on PU fibres with a controllable diameter and porous structure can be used as
coatings for favourably modifying host responses to implantable biosensors. However, the additional membrane for engineering the tissue responses would in turn also further decrease the pre-implantation sensitivity of the implantable biosensor.

The objective of this study was to optimize the parameters for electrospinning PU core-Ge shell coaxial fibres, characterize them, apply directly on model coil-type implantable glucose biosensors and evaluate their effects on in vitro sensor function. A specialised spinneret, made of concentric tubes connected to two separate fluid sources, such that coaxial fibres can be electrospun, was designed and manufactured. The solution and process parameters for electrospinning coaxial fibres were varied. The gelatin shell was stabilized by crosslinking. The membranes were characterised for morphology, pore sizes, porosity, hydrophilicity, solute diffusion, chemical and mechanical properties. Glucose biosensors were then coated with optimized co-axial fibre membranes and their effects on sensor function evaluated.

2. Materials and Methods

Selectophore™ polyurethane (PU), Gelatin (Ge) from porcine skin (type A), Tetrahydrofunan (THF), N,N-Dimethylformamide (DMF), 2,2,2-Trifluoroethanol (TFE) (≥99.0% (GC)), bovine serum albumin (BSA), glutaraldehyde grade I (50%) (GA), glucose oxidase (GOD) (EC 1.1.3.4, Type X-S, Aspergillus niger, 157,500U/g, Sigma), ATACS 5104/4013 epoxy adhesive, Brij 30, D-(+)-glucose and 0.01M phosphate buffered saline (PBS) tablets were purchased from Sigma–Aldrich–Fluka. Teflon-coated platinum–iridium (9:1 in weight, Ø 0.125mm) and silver wires (Ø 0.125mm) were obtained from World Precision Instruments, Inc. (Sarasota, FL).

2.1 Glucose biosensors

A miniature coil-type implantable glucose biosensor was used as model sensor in this study [2, 8-10]. The amperometric sensor is a two electrode system based on Pt/Ir working electrode with immobilised GOD enzyme and silver/silver chloride (Ag/AgCl) reference electrode. The procedures of sensor manufacture are based on our previous reports [2, 8-10].

2.2 Electrospinning coaxial PU/Ge fibres

To spin co-axial fibres, a coaxial spinneret was custom-made. It consisted of a stainless steel Tee-Union (1/8” Swagelok, UK), a PTFE union set (1/16”, a PTFE union, two PTFE cones and two PEEK adaptors), and two stainless steel concentric tubes that allow coaxial extrusion of two fluids simultaneously. The inner tube has an inner diameter of 0.508mm and an outer diameter of 0.711mm, while the outer tube
has an inner diameter of 2.88 mm and an outer diameter of 3.0mm. The electrospinning setup utilized for this study was described earlier [2, 3]. The coaxial spinneret was mounted vertically above a grounded steel plate (collector, 16x16 cm²). Two syringe pumps (Fusion 100) were used to pump inner/outer polymer feed solutions in 10 ml plastic syringes (BD). In this study, a fluorinated alcohol, TFE (100%), was used as the solvent for both gelatin (shell) and PU (core). Gelatin was dissolved in TFE under continuous stirring at room temperature for at least 10 h. PU solutions of different concentrations (2, 4, 6, and 8 w/v%) were also prepared in TFE at room temperature, under continuous stirring for 6 h. Dynamic viscosity of the TFE solvent, different PU and Ge solutions were measured using Automated Micro Viscometer (AMVn, Anton Paar). For spinning co-axial fibres, gelatin and PU solutions were separately fed to the outer and the inner needles of co-axial spinneret simultaneously, using two programmed syringe pumps through 1/8” OD and 1/16” OD PTFE tubing with the feed rate of 1.2ml/h and 0.8ml/h respectively. Applied voltage between 11.25-14kV and a tip-to-collector distance of 15cm were used to ensure the formation of a steady coaxial jet with an external meniscus surrounding the inner one ejecting from a stable Taylor Cone for the different PU solution concentrations with Ge concentration kept constant at 10 w/v%. The fibres were spun at ambient room temperature (20±2°C) and humidity (40±5%).

2.3 Crosslinking of gelatin

Gelatin, being water soluble, needs stabilization by crosslinking to maintain the co-axial fibre structure. Two methods were tested for GA crosslinking of the gelatin sheath of the co-axial fibres. First, the coaxial fibre membranes on Al foils were immersed in aqueous glutaraldehyde (25% GA solution diluted with DI water at 1:99 volume), under continuous shaking for 12 hours at room temperature. The membranes were washed in several changes of DI water, dried over night at 40°C and then stored in a vacuum desiccator until further use. Secondly, GA crosslinking of gelatin was achieved by incubating the co-axial fibre membranes on Al foils in a vacuum desiccator with the desiccant replaced by 10 ml of 25% aqueous GA solution (in a petri dish) for 3 days at room temperature. The samples were then transferred to a power-assisted vacuum desiccator for removal of excess GA and stored in desiccator until further use.

2.4 Electrospinning co-axial fibres directly on sensor surface

A dynamic collector configuration reported in our previous study [2], was used for electrospinning co-axial fibres directly on sensor surface, wherein the sensor was inserted in 0.5 inch stainless steel needle (23G blunt-tip) and the needle fixed at the end of a custom-made rotator. A rotation speed between 660 – 690
rpm, obtained by setting both voltage and current constant at 5V and 0.11A respectively, was chosen to obtain random orientation of the electrospun fibres.

Two types of Ge-PU coaxial fibrous membranes (ESC) were electrospun both on sensors (Pt-GOD) coated with epoxy-PU (EPU) semi-permeable membrane (Pt-GOD-EPU-ESC) and those without (Pt-GOD-ESC) to study the ability of coaxial-fibre membranes as mass-transport limiting membranes. The designations and electrospinning conditions for co-axial fibre membranes spun directly on the sensor surface are summarised in Table 1. GA crosslinking of Ge in the co-axial fibres was achieved by incubating the sensors in vacuum desiccator saturated with GA fumes as described above (section 2.3). Six sensors per electrospinning coating configuration (Pt-GOD-EPU-ESC or Pt-GOD-ESC) were tested for in vitro functional efficacy and six sensors without any electrospun coatings (Pt-GOD-EPU) used as controls.

2.5 Characterization of electrospun membranes

2.5.1 Infrared spectroscopy

An ATR-FTIR spectrophotometer (PerkinElmer Inc.) was used to verify the core-shell fibre structure of electrospun coaxial fibres. Each spectrum, acquired in transmittance mode, was an average of 128 scans at a resolution of 4 cm\(^{-1}\).

2.5.2 Core-shell structure of the fibres and morphology of the membranes

Transmission electron microscope (TEM, HITACHI H-600) was used to examine its coaxial structure, with an accelerating voltage of 100kV. The samples for TEM observations were prepared by collecting the nano-fibres onto carbon-coated Cu grids. The electrospun membranes were also sputter coated for 30 sec with gold using an AGAR high-resolution sputter-coater and observed under a field emission scanning electron microscope (FESEM, Zeiss Supra 35VP) in SE mode for morphology.

2.5.3 Fibre diameter and membrane thickness

The fibre diameters were measured on SEM images using a user friendly application developed in-house using Matlab for length measurements. A total of 160 measurements were made on 8 different SEM images, each representing a non-overlapping random field of view for each electrospun membrane configuration. To obtain the fine cross section images for the electrospun membranes (both sheets and on sensors), the membranes were snap-frozen in liquid nitrogen, and then cut using a scalpel. The
resulting samples were processed for SEM and oriented appropriately to obtain image of cross-sections of the membranes. The above-mentioned software for diameter measurements was also used to measure the thicknesses of the membrane using SEM images of their cross-sections. The thickness of the electrospun membranes were also measured using a digital micrometer having a resolution of 0.001mm. The membranes were sandwiched between two slides and their thickness determined by subtracting the glass slides’ thickness.

2.5.4 Pore size and Porosity

The pore size for the different membranes was measured using extrusion porosimetry (also called bubble point measurement) as reported earlier in details [3, 11]. The range of pore sizes (radius $\alpha$) was calculated using the Young-Laplace Eq. 1:

$$\alpha = \frac{2 \gamma_{st} \cos \theta}{\Delta P} \quad \text{(Eq. 1)}$$

where $\Delta P$ is the differential pressure, $\gamma_{st}$ the surface tension of the wetting liquid and $\theta$ the wetting angle, which for a completely wetted membrane is 1 [11].

The porosity of the membranes was also determined using gravimetry as described earlier using the following equations:

$$\rho_{\text{app}} = \frac{m}{d \times A} \quad \text{(Eq. 2)}$$

$$\varepsilon = 1 - \left( \frac{\rho_{\text{app}}}{\rho_b} \right) \quad \text{(Eq. 3)}$$

where $m =$ the mass of the membrane (g), $d =$ the thickness of the membrane (cm), $A =$ the area of nano-fibrous mat (cm$^2$), $\rho_b =$ the bulk density of materials (g/cm$^3$). $\rho_b$ was estimated using by Eq. 4:

$$\rho_b = \frac{\rho_{PU} \cdot r^2 + \rho_{Ge} (R^2-r^2)}{R^2} \quad \text{(Eq. 4)}$$

Where $r$ is PU core radius, $R$ the total fibre radius (measured using SEM), and $\rho_{PU}$ and $\rho_{Ge}$ are the bulk densities of Selectophore™ PU (1.04g/cm$^3$ as reported by manufacturer) and crosslinked Ge respectively (1.369g/cm$^3$) [12].
2.5.5 Uniaxial tensile testing

A 40min electrospinning time was used to obtain membranes thick enough for easy handling for mechanical testing. For the tensile tests, the membranes were first cut into 50 mm long and 10 mm wide strips following a method reported earlier [13]. The thickness of the membrane was measured using a digital micrometer having a resolution of 0.001mm. For the tensile testing, the ten mm wide strips were soaked overnight in distilled water. The wet strips were mounted onto an Instron tester (Model 5542) fitted with automatic clamps (30mm apart). Preload of 0.01 N was used to precondition the samples followed by the test to failure using a 10N static load cell and the test speed of 10 mm/min at room temperature. Load/extension data was logged using a computer equipped with Instron's Bluehill® Lite software. The ultimate tensile strength (UTS) (F_{max}/original cross-sectional area, N/mm^2) and modulus of elasticity (from the relationship \( \sigma \) E, where \( \sigma \) is the stress, \( \varepsilon \) is the strain, and E is the modulus of elasticity, N/mm^2) and strain at UTS were then determined.

2.5.6 Contact angle measurements

The contact angles for a drop of distilled water on electrospun membranes were measured using a contact angle instrument (OCA15+, Data-physics, Germany) at room temperature. A single drop of 1μL DI water was dropped on the surface of a flat 10 x 10 mm membrane using a syringe perpendicular and image captured in <1s after the water droplet became stable on the surface. This process was repeated four times on each membrane. The contact angles were then measured using the instrument's SCA20 software.

2.5.7 Diffusion test

The effects of fibre diameter, thickness and porosity of electrospun Ge-PU coaxial membranes on their permeability to glucose were tested using a biodialyser (singled-sided biodialyser with magnet, 1ml, Sigma-Aldrich) in a beaker as a two-component diffusion chamber. For diffusion test, the donor solution chamber (A) of the biodialyser was filled with 1ml of glucose solution in PBS (pH 7.4 at 37°C) and the wet membrane was mounted and secured with the treaded cap ring exposing a 113.14 mm² membrane area for diffusion. The assembly was immersed and rotated in 49 ml of receiving PBS (chamber B) [14, 15]. Pre-calibrated amperometric glucose sensors made in our lab were immersed in the receiving PBS to continuously log the changes in glucose concentration (as described in Section 2.6). The donor chamber
solution concentration was chosen such that the eventual equilibrium glucose concentration of receiver solution reaches 30 mM. More details of the test can be found in paper [3].

The effective diffusion coefficient was determined from the time dependent glucose concentration assuming that a quasi-steady-state concentration condition within the membrane. Based on the assumptions, combining the diffusion equation with Fick’s law and mass balance conditions between the two chambers A and B given Eq. 5 [16, 17]:

$$\frac{(C_B - C_A)}{(C_{B0} - C_{A0})} = e^{-\frac{t}{\tau}}$$  \hspace{1cm} (Eq. 5), with

$$\tau = \left(\frac{d}{D_{eff}S}\right) \times \left(\frac{(V_A V_B)}{(V_A + V_B)}\right)$$

where $C_A$, $C_B$ and $V_A$, $V_B$ are the concentration and volume of the chamber A and B, d is the membrane thickness, S is the membrane surface area, t is time and $D_{eff}$ is the effective diffusion constant for the membrane. The mean relaxation time $\tau$ for each membranes with different thickness was calculated by linear regression of $\ln\left(\frac{(C_B - C_A)}{(C_{B0} - C_{A0})}\right)$ vs t from experimentally measured values of tracer concentration of chamber B at different times, and then $D_{eff}$ could be calculated from the mean relaxation time $\tau$.

2.6 Sensor function testing

2.6.1 Basic Test

Sensor function was tested by amperometric measurements of glucose in PBS using Apollo 4000 Amperometric Analyser (World Precision Instruments Inc., Sarasota, FL) at 0.7 V versus Ag/AgCl reference electrode. The buffer solution (PBS) was continuously stirred to ensure mixing of glucose in solution. Calibration plots for the sensors were obtained by measuring the current while increasing the glucose concentration from 0–30 mM (stepwise). The response time was calculated as 90% of the maximum response time after increasing the glucose concentration from 5 to 15 mM. The sensitivity (S) of each sensor was calculated using Eq. 6:

$$S = \left(\frac{I_{15mM} - I_{5mM}}{I_{15mM}}\right) / 10$$ \hspace{1cm} (Eq. 6)

where $I_{15mM}$ and $I_{5mM}$ are the steady state currents for 15 and 5 mM glucose concentration respectively. All experiments were carried out at room temperature.

2.6.2 Efficacy of electrospun membrane coatings on sensor function and longevity
The performance of the sensors was studied by the same intermittent measurements of sensor response sensitivity and linearity as described in our recent paper [2]. We monitored long-term performance of first generation glucose biosensor by tracing of sensitivity rather than the response currents in order to avoid the latter being affected by the background current or the accumulated $H_2O_2$ in the enzyme layer [10, 18, 19]. Briefly, the sensor function test on each sensor was repeated several times up to 84 days to test their longevity. The sensors (Pt-GOD-EPU-ESC, Pt-GOD-ESC and Pt-GOD-EPU) were calibrated at 1, 3 and 7 days before and tested on 1, 3, 7, 14, 21, 28, 35, 42, 56, 70, and 84 days after applying the electrospun coatings. The control sensors, Pt-GOD-EPU sensors without ESC were also processed and tested similar to those with ESC. Between the tests, sensors were stored in PBS at 37°C. The storage PBS was replaced with fresh PBS every 2 to 5 days.

2.7 Statistical analysis

Statistical analyses were carried out using statistical software (SPSS v.15). Statistical variances between groups were determined by one-way analysis of variance (ANOVA). Tukey's test was used for post hoc evaluation of differences between groups. A P value of <0.05 was considered to be statistically significant. Unless otherwise mentioned, all data presented is expressed as mean ± standard deviation.

3 Results and discussion

3.1 Process optimization for electrospinning PU-GE co-axial fibres

The solubility of Ge in a variety of organic solvents is very poor and a highly polar TFE was essential to obtain an electrospinnable solution of Ge. At the same time, the incompatibility of Ge with other solvents meant that it was also essential to prepare the co-electrospinning PU solution in TFE. However, at concentrations >8% (wt/v) PU solution in TFE was saturated and too viscous for electrospinning (Table 2). Hence for this study, PU concentration in TFE was varied from 2 to 8%, while that for Ge kept constant at 10%.

TFE being highly volatile, special care was needed to maintain a stable Taylor cone throughout the electrospinning process. The process was eventually optimized for obtaining coaxial fibres with the following combination of parameters: electrical voltage in a range of 11-15kV, a working distance (between spinneret and collector) of 15cm, and flow rates of inner (PU) and outer (Ge) solution at 0.8ml/h and 1.2ml/h respectively. The ambient temperature (20±2°C) and humidity (40±5%) were not controlled. The formation of the co-axial fibre structure was first ascertained using TEM as illustrated in Fig 1a.
Thereafter, to make Ge insoluble and maintain the co-axial fibre structure (Fig 1b), it was crosslinked with GA using two methods. Firstly, freshly electrospun co-axial fibre membranes were immersed in GA solution, which disrupted the co-axial fibre structure (Fig 1c). Secondly, the membranes were incubated in a GA saturated air for 3 days, which preserved the co-axial fibre structure while making the Ge sheath insoluble in water (Fig 1d). Similar result was also reported earlier [20]. Four co-axial fibre compositions, designated as 2PU10Ge, 4PU10Ge, 6PU10Ge and 8PU10Ge were further characterised in this study.

3.2 Core-shell fibre structure

To ascertain the formation of core-shell fibre structure and any interactions between Ge and PU at their interface, ATR-FTIR spectra (Fig 2a) were recorded for the different coaxial fibres without crosslinking the Ge shell, while having their feed components – Ge powder and PU pellets as controls. Ge showed its typical amide bands at 1634 cm\(^{-1}\) (amide I), 1531 cm\(^{-1}\) (amide II) and 1233 cm\(^{-1}\) (amide III) corresponding to C=O stretching, the coupling of N–H bending and C–N stretching vibrations, and N–H bending respectively pertaining to the triple helical structure of gelatin [21]. The typical PU peaks included the C–H stretching vibrations between 2825 and 2946 cm\(^{-1}\), 1731 cm\(^{-1}\) the free urethane carbonyl peak, 1106 cm\(^{-1}\) the soft segment ether peak and 1033 cm\(^{-1}\) the hard segment ether peak. The spectra for co-axial fibres showed the unique peaks of both PU and Ge indicating the presence of both PU and Ge in the co-axial fibre membranes. The broad absorption band centred at ~3288 cm\(^{-1}\) was found in the FTIR spectra of all PU/GE coaxial fibres, which can be attributed to overlapping peaks of the N–H and O–H stretching vibration. Moreover, no shift in characteristic peaks of either PU or Ge was observed for any of the electrospun coaxial fibre membranes, suggesting that there may be no obvious interaction between PU and GE. This is consistent with the observation on electrospun Ge/PU blended nano-fibres reported earlier [22].

FTIR spectra were also recorded for co-axial fibre membranes from which the Ge shell was washed off using DI water at 40°C for 1 week with water changes every 24h. The spectra for all membranes were similar to that of pure PU, with the exception of 2PU10Ge samples, which showed a prominent peak, at about 1633 cm\(^{-1}\) unique to pure Ge (Fig 2b). Moreover, the morphology of 2PU10Ge showed spreading of the polymer between the fibres (Fig 2c-f). Although, not spreading out as observed with 2PU10GE, the fibres of the other membranes appeared to fuse with neighbouring fibres, which could be indicative of some degree of blending of PU and Ge at the interface between the PU core and Ge shell during
electrospinning. When the Ge is washed off, the residual PU in the transition layer on the surface of PU core could cause neighbouring fibres to fuse.

To visualise the core-shell structure of the co-axial fibres under SEM, following crosslinking of Ge-shell with GA vapour, the PU core was dissolved using THF. The resulting hollow fibres for the different membranes are shown in Fig. 3. The tubular structure was prominent for all membranes. However, for 4PU10Ge membranes some of the fibres did not form complete tubes (Ge-shell) (Fig. 3b), which could be due to an observed experimental anomaly, wherein the fast evaporation of solvent causes solidification of gelatin on the tip of the nozzle either blocking or significantly slowing the shell fluid flow. The amount of the shell liquid in Taylor cone thus decreases to a point where the viscous drag applied by the sheath solution could be insufficient to confine the core solution within the Taylor cone. We observed that the evaporation of the solvent (TFE) could be slowed to ensure formation of proper co-axial fibre by saturating the air around the spinneret with the solvent or by increasing ambient humidity.

3.3 Morphology, diameter and composition of coaxial fibres

The morphology and fibre diameter distributions for 2PU10Ge, 4PU10Ge, 6PU10Ge and 8PU10Ge are presented in Fig. 4. For the 2PU10Ge membranes occasional spindle shaped beads formed (Fig. 4a), which could be due to the disparity in solvent content in the core PU solution compared to the shell Ge solution. Due to the low viscosity of the core PU solution (~33.28 x 10^{-4} pa/s, Table 2), droplets could form while the polymer solution accelerates to the collector resulting in bead formation, similar to that reported earlier [23]. For 4PU10Ge, 6PU10Ge and 8PU10Ge, the viscosity of core PU solution (Table 2) was sufficient to result in seamless coaxial fibres without beads (Fig. 4b to d). Further, their narrow fibre diameter distributions indicated a stable electrospinning process (Fig. 4e to h). The average fibre diameter increased significantly with increasing feed solution concentration of the core PU solution, which phenomenon is widely reported [24-26]. Fibre diameters were submicron only for 2PU10Ge membranes.

The diameter of PU-core was measured on SEM images (Fig 2C-F) of co-axial membranes whose Ge-shells were washed off and subtracted from that of the co-axial fibres with GA crosslinked Ge-shells (Fig 4) to obtain the total thickness Ge-Shell. The results are presented in Table 3 and Fig S1. The data for 2PU10Ge could not be obtained because the thin PU fibres failed to retain their integrity as well as the presence of residual Ge (Fig 2b). However, for the rest of the membranes that had intact fibres, the increase in concentration of feed PU solution significantly increased the diameter of the PU-core in the coaxial fibre, while the thickness of its Ge-shell showed a gradual decrease (Table 3). The ratio of the
diameter of PU core to the total thickness of Ge shell on the co-axial fibres further reiterates the increasing volume of PU core with increasing feed PU solution concentration (Table 3). Similar increase in the diameter of PCL-core with a parallel decrease in Ge-shell thickness was also reported by Zhang et al., who explained the decrease in the shell thickness to be due to the same mass of the shell layer distributed over a larger core [27].

3.4 Pore size and porosity

To study the effect of thickness on pore sizes, two electrospinning times, 10 and 40 min were chosen for spinning coaxial fibre membranes. The pore size distributions are presented in Fig. 5. For membranes electrospun for 10 minutes, an increasing pore size was observed with increasing fibre diameter (Table 4 and Fig. 5a), ranging from 920.70 nm for 2PU10Ge-10’, 1964.34 nm for 4PU10Ge-10’, 2624.23 nm for 6PU10Ge-10’ through to 3051.74 nm for 8PU10Ge-10’. Furthermore the pore size distribution was narrow and sharp for 2PU10Ge-10’, which became broader for other coaxial fibre membranes with increasing fibre diameters (Fig. 5). The membranes having smaller fibre diameters have been widely reported to have smaller pore sizes and narrower pore size distribution [3, 28-30]. When the thickness of the coaxial fibre membranes was increased by increasing the electrospinning time to 40 min, the pore sizes again increased with increasing fibre diameter (Table 4 and Fig. 5b), with the exception of 8PU10Ge membranes which had pore sizes smaller than 4PU10Ge and 6PU10Ge (Fig. 5b). However, compared to membranes electrospun for 10 min, all the thicker membranes showed smaller pore sizes: 333.85, 1444.32, 1535.61 and 1028.91 nm in radius respectively for 2PU10Ge-40’, 4PU10Ge-40’, 6PU10Ge-40’ and 8PU10Ge-40’ (Fig. 5b). Such influence of electrospinning duration on pore size was also observed by Chiu et al., who fabricated electrospun polyacrylonitrile ion-exchange membranes [31]. They reported a sharp decrease in average pore diameter in the first 1 to 3h electrospinning time following which the pore sizes stabilized. The decrease in pore sizes with increasing membrane thickness could be due to the tighter packing of fibres induced by the increasing weight of the fibres being continuously deposited. The notably lower porosity of the thickest 8PU10GE-40’ membranes (Fig. 5b) can also be attributed to the densely accumulated fibres leading to smaller pore sizes similar to that observed by Soliman et al. [30]. The denser fibre packing causing lower porosity was also supported by the gravimetry based porosity estimations (Table 4).

Membrane thickness, material bulk density, fibre packing density and pore volume estimations were calculated and summarised in Table 3 for the different electrospun coaxial fibre membranes electrospun
for 10 and 40 min. As the core diameter in 2PU10Ge could not be detected, its corresponding porosity was not calculated. No significant change in the overall thickness was observed when the coaxial fibre membranes (4PU10Ge to 8PU10Ge) were electrospun for 10 min. In addition, they had apparent densities (or fibre packing density) in the range of 0.21-0.28 g/cm³, which change was again not statistically significant, in spite of increasing feed PU solution concentration. Therefore, the comparable resultant pore volumes can primarily be ascribed to their corresponding similar bulk density of the composite PU-Ge fibres (Table 3). All the coaxial fibre membranes electrospun for 10 min had pore-volumes greater than 60%, which could be useful for tissue engineering application requiring cellular infiltration to the bulk of porous scaffolds [32].

However, when the electrospinning time was increased, inconsistent membrane thickness, fibre packing densities and pore volumes were observed (Table 3). The thickness of the membranes decrease from 2PU10Ge to 6PU10Ge, and that of 8PU10Ge was higher than all the other membranes. The fibre packing densities also followed the trend observed with thickness measurements, but the fibre packing density for 8PU10Ge was comparable to that observed for 6PU10Ge. The pore volume estimations revealed a steady trend similar to that observed with membranes electrospun for 10 min. However, the pore volumes of coaxial fibre membranes electrospun for 40 min was significantly lower than those electrospun for 10 min. Overall, the insignificant variations in pore volumes with decreasing PU core volumes from 8PU10Ge to 4PU10Ge can be due to the cross-interference of the decreasing mechanical support in the inner layer and cross-linking process. Although the porosity (pore volume) would decrease with the shrinking of Ge shell due to GA crosslinking as demonstrated for hydrogels in general [33-36], core PU with good mechanical properties can moderate the deformation during crosslinking process. Similar observation was also reported by Zhao et al., wherein crosslinking of Ge shell layer resulted in negligible effects on porosity of coaxial fibre membranes having semi-crystalline hydrophobic PCL core [36]. To sum up, the pore sizes for PU-GE core-shell fibrous membranes are mainly dependent on fibre diameter, while the pore volume was dependent on the dielectric properties of the electrospinning solution. Electrospinning duration was shown to have a decreasing effect on pore sizes and volume for the electrospinning times tested in this study.

3.5 Contact angle

Addition of hydrophilic gelatin to the co-axial fibre structure was expected to increase the hydrophilicity of the electrospun membranes. The surface hydrophilicity for the coaxial fibres was so quick that for
4PU10Ge, 6PU10Ge and 8PU10Ge it was not possible to measure contact angles. 2PU10Ge was the only coaxial fibre configuration for which contact angle could be measured (101°, Fig S2). The rapid wetting of coaxial fibres can be attributed both to hydrophilic surface chemistry and surface roughness [37, 38]. The smooth and essentially non-porous surfaces of GA crosslinked Ge (Fig S3) and pure PU films had contact angles of 64° and 86° respectively [3]. Increasing surface roughness for the electrospun pure PU membranes showed increasing hydrophobicity that can be attributed to the hydrophobic air pockets in the pores [3]. In contrast, the increased roughness of co-axial fibre membranes made their surface highly hydrophilic, with an exception of 2PU10Ge. For 2PU10Ge a contact angle of 101° was observed, which was comparable to the pure PU fibre membranes (8PU, 104°) of similar submicron porosity. This may be reminiscent of “lotus effect” in that the intrinsic hydrophobicity of a surface can be enhanced by being textured with different length scale of roughness. The average pore radius of about 333nm observed with 2PU10Ge (Table 4), could be responsible for preventing the water droplet from wetting the nanostructured space between the nano-fibres. The macro-pores on the surface of 4PU10Ge, 6PU10Ge and 8PU10Ge, could disrupt the integrity of water drop and thus, essentially accelerate its absorption into the macro pore network through capillary action.

3.6 Tensile mechanical properties of electrospun coaxial fibre membranes

The tensile mechanical properties of water swollen electrospun coaxial fibre membranes are summarised in Table 4 and Fig S3. Electrospun pure PU fibre (8PU) and Ge fibre [22] membranes are also listed as controls. 8PU membranes were highly flexible and elastic, while. On the other hand, Ge fibre membranes was reported to be brittle and weak, especially when wet [22]. The co-axial fibre membranes showed intermediate tensile properties between that observed for 8PU and Ge fibre membrane controls. Among the coaxial fibre membranes, the tensile properties improved with increasing core PU content (Table 4) with the exception of 2PU10Ge, which had significantly higher Young’s modulus and lower strain at break. The higher elasticity of 2PU10Ge membranes can be attributed to higher density of fibres per unit volume as well as higher degree of inter-fibre contacts and crosslinking. The lower strain at break for 2PU10Ge can also be due to its larger content of crosslinked Ge (≥52%). The other co-axial fibre membranes became more flexible with a significant increase of strain at breaks with increasing diameter of PU core. Similar results were also reported earlier [7, 39, 40]. Furthermore, the failure of coaxial fibres, containing Ge shell, under tensile load is said to first start with cracks in the hydrated Ge, which is then translated to the core [4, 22, 39]. This presumption was verified by Zhao et al. [36] using SEM showing
the cross section of crosslinked Ge coated PCL fibres after fracture. Therefore, the mechanical properties of the co-axial fibrous membranes are resultant of the synergetic effect of core-shell structure.

### 3.7 Effective diffusion coefficient

The permeability of the coaxial fibre membranes 2PU10Ge and 6PU10Ge of varying thicknesses to glucose was tested using biodialysers. The glucose diffusion followed an initial linear increase that plateaued off to a constant when the diffusion rate attained equilibrium (Fig 6A & B). The slope of the linear increase typically decreased with increasing membrane thickness, which was statistically significant for 2PU10Ge-10’ membrane. The data was also fitted in Eq 5 to calculate relaxation times and effective diffusion constants for the membranes, which results are presented in Fig 6C and Table 5. The $D_{eff}$ for PU-Ge coaxial fibre membranes ranged from $9.5\pm0.47 \times 10^{-5}$ to $5.84\pm0.44 \times 10^{-4}$ mm$^2$/s, which in general were higher compared to that we reported for pure PU fibre membranes ($0.429\pm0.45 \times 10^{-5}$ to $4.88\pm0.44 \times 10^{-4}$ mm$^2$/s) depending on their fibre composition and thickness [3]. The comparatively higher flux rates for PU-Ge fibres can be attributed to the higher pore volumes (about 78 to 82%, Table 3) and the hydrophilic Ge surface of the co-axial fibres, which was maximum for 2PU10Ge-2.5’ membranes, even closer to that of glucose in water $6.73 \times 10^{-4}$ mm$^2$/s [41]. However, with increasing membrane thickness a decrease in $D_{eff}$ with a concomitant increase in average relaxation time ($\tau$) was observed (Fig. 6C and Table 5) consistent with that observed with pure PU fibre membranes [3].

### 3.8 Efficacy of electrospun coaxial fibre membranes as coatings for implantable coil-type glucose biosensors

Among 4PU10Ge, 6PU10Ge and 8PU10Ge, the variation in fibre diameters was narrow ranging from 1.04 to 1.22 µm (Table 3). Coaxial fibre structure was not consistent for 4PU10Ge, while the pore sizes distribution for 8PU10Ge was broader than both 4PU10Ge and 6PU10Ge (Fig. 5a). Hence, the 6PU10Ge membrane, having consistent coaxial fibre structure and pore size, was chosen as coating for glucose biosensors. However, for studying the effects of fibre diameter of coaxial fibre membranes on sensor function, 2PU10Ge membrane that has about half the average fibre diameter observed for 6PU10Ge, was also chosen. Effectively, the effects of coaxial fibre membranes on glucose biosensor function were evaluated as a function of fibre diameter and Ge content. 2PU10Ge membrane had an average fibre diameter of 540.61±90.83 and ~51.8% Ge while that for 6PU10Ge was 1152.93±128.77 and ~34.5% respectively. The two types of membranes were electrospun directly on glucose biosensors using the
dynamic collector, wherein the biosensor was rotated at about 660 to 690 rpm in the electrospinning field. The parameters are listed in Table 1.

Each of the coaxial fibre membranes, were tested on sensors with (Pt-GOD-EPU) and without (Pt-GOD) EPU mass-transport limiting membrane. The results were compared with that of Pt-GOD and Pt-GOD-EPU sensors. The sensor function was tested at regular intervals starting one week before applying coatings, to 84 days (12 weeks) after coating. The sensitivity and linearity results at each tested time point were normalized to that at day 7, before applying coatings.

The effect of 2PU10Ge coatings on glucose biosensor function is illustrated in Fig. 7. The sensitivity profiles for all sensors before and after coating with membranes were similar to that shown by Pt-GOD sensors indicating no obvious effects of either EPU or 2PU10Ge membranes on sensor sensitivity (Fig. 7A). However, the linearity ($R^2$) for the detection range of 2 to 30 mM glucose was only improved for EPU membrane, indicating that the 2PU10Ge membranes did not function as a durable mass transport limiting membrane over the period of time tested (Fig. 7B). Thus, as summarised in Fig 7C, 2PU10Ge did not affect sensor sensitivity, but also did not function as a mass-transport limiting membrane.

Different from 2PU10Ge coatings, 6PU10Ge caused a reduction in sensor sensitivity, but extended the linear detection range for glucose biosensors (Fig 8). The sensitivity profiles for sensors coated with 6PU10Ge were significantly lower than that of Pt-GOD and Pt-GOD-EPU sensors (Fig 8A). However, both EPU and 6PU10Ge membranes extended the linear detection range for Pt-GOD to cover the physiologically relevant detection range of 2 to 30 mM glucose, demonstrating that 6PU10Ge membranes function as a mass transport limiting membrane (Fig 8B). The trends in % change in sensitivity and linearity as a function of sensor coating composition further reiterate the above observations (Fig 8C).

3.9 Long-term stability of sensor function for glucose biosensors coated with coaxial fibre membranes

The sensitivity of sensors coated with both 2PU10Ge and 6PU10Ge were stable throughout the study for 12 weeks. However, a decrease in linearity for 2PU10Ge coated Pt-GOD and Pt-GOD-EPU sensors was observed. To identify the cause for this undesirable decrease in linearity, the morphology of the sensors after completion of the study was assessed under SEM. As shown in Fig. 9A&B, the integrity of the fibro-porous membrane structure was disrupted for 2PU10Ge forming a PU fibre-reinforced Ge composite film. On the other hand, the fibro-porous structure of 6PU10Ge membranes was intact even after 12 weeks of immersion in PBS pH 7.4 at 37°C although neighbouring fibres fused (Fig 9C&D).
4 Conclusions

The solvent, solution concentration and process parameters for electrospinning Selectophore™ PU-core and Ge-Shell coaxial fibre membranes were optimized. PU-Ge coaxial fibres of varying composition and structure were prepared and characterized. With increasing PU feed solution concentration, an increase in the diameter of PU core, with concomitant reduction in Ge shell thickness was demonstrated. Accordingly, the fibres also inherited intermediate mechanical properties of their constituents - PU and Ge. The higher the Ge-shell content the more brittle, while the higher the PU-core content the higher was the strength and the elongation at break for the coaxial fibre membranes. The Ge-shell ensured higher hydrophilicity and flux-rates for glucose transport across the PU-Ge coaxial fibre membranes. The effects of the coaxial fibre membranes on in vitro function of electrochemical glucose biosensors were also evaluated. Higher Ge content in the coaxial fibre resulted in maintenance of sensitivity similar to that of control sensor without the coaxial fibre coating, but failed to function as a mass transport limiting membrane and also lost the integrity of its fibro-porous structure. In contrast, a thinner Ge-shell layer on PU-core resulted in lowered sensitivity, but the coaxial membrane functioned as a mass-transport limiting membrane, while maintaining the integrity of its fibro-porous structure till the end of the study period of 12 weeks. Thus, the electrospun PU-Ge coaxial-fibre membranes having significantly high pore volumes, interconnected porosities and tailorable mechanical properties, permeability and surface chemistry compared to conventional solvent cast membranes can find applications as tissue engineering coatings for biosensors requiring analyte exchange and other implantable biomedical devices. Such biomimetic coatings are anticipated to play an important role in engineering tissue responses to the implanted biosensors, which pre-clinical functional efficacy studies will be reported in our next paper.

5. Acknowledgements

This research is supported by Brunel University, the Royal Society research grant (RG100129), the Royal Society-NSFC international joint project grant (JP101064) and the National Institute of Health (NIH/NIBiB, grant R01EB001640).

6. References


Huang Z-M, Zhang YZ, Ramakrishna S, Lim CT. Electrospinning and mechanical characterization of gelatin nanofibers. Polymer 2004;45:5361.


Table 1. Electrospinning conditions used for spinning coaxial PU-Ge fibres directly on biosensor surface (n=6)

<table>
<thead>
<tr>
<th>Designation</th>
<th>Concentration (w/v %)</th>
<th>Voltage (kV)</th>
<th>Feed Rate (ml/h)</th>
<th>Distance (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt-GOD-EPU-6PU10Ge</td>
<td>6 10</td>
<td>13.25</td>
<td>0.8 1.2</td>
<td>15</td>
</tr>
<tr>
<td>Pt-GOD-6PU10Ge</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt-GOD-EPU-2PU10Ge</td>
<td>2 10</td>
<td>14</td>
<td>0.8 1.2</td>
<td>15</td>
</tr>
<tr>
<td>Pt-GOD-2PU10Ge</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Table 2.** Dynamic viscosity of the different PU and Ge solutions used for electrospinning co-axial fibre membranes, * viscosity was higher than measuring limit for the instrument.

<table>
<thead>
<tr>
<th>Solvent/Solution</th>
<th>Concentration – wt/v (wt/wt) (%)</th>
<th>Dynamic Viscosity x 10^-4 (Pa/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TFE</td>
<td>100</td>
<td>7.684±0.008</td>
</tr>
<tr>
<td>PU 2 (1.5)</td>
<td></td>
<td>332.763±1.038</td>
</tr>
<tr>
<td>PU 4 (2.9)</td>
<td></td>
<td>1373.714±0.968</td>
</tr>
<tr>
<td>PU 6 (4.3)</td>
<td></td>
<td>3764.733±6.055</td>
</tr>
<tr>
<td>PU 8 (5.8)</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Ge 10 (6.8)</td>
<td></td>
<td>797.122±1.006</td>
</tr>
</tbody>
</table>
Table 3 Porosity estimations for electrospun coaxial fibre membranes based on gravimetry, calculated using Eq.s (2) and (3). n=3.

<table>
<thead>
<tr>
<th>Electrospun time, min</th>
<th>2PU10Ge</th>
<th>4PU10Ge</th>
<th>6PU10Ge</th>
<th>8PU10Ge</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibre Diameter (μm)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>0.54±0.06</td>
<td>1.04±0.10</td>
<td>1.15±0.13</td>
<td>1.22±0.11</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>1.34</td>
<td>1.34</td>
<td>1.32</td>
</tr>
<tr>
<td>Fibre Bulk density (g/cm³)</td>
<td>-</td>
<td>1.34</td>
<td>1.34</td>
<td>1.32</td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>920.7</td>
<td>1964.34</td>
<td>2624.23</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>333.85</td>
<td>1444.32</td>
<td>1535.61</td>
</tr>
<tr>
<td>Pore size in radius (nm)</td>
<td>10</td>
<td></td>
<td>21.6±4.0</td>
<td>23.4±2.6</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>56.8±4.9</td>
<td>40.3±5.7</td>
<td>36.0±5.4</td>
</tr>
<tr>
<td>Thickness (μm)</td>
<td>10</td>
<td>21.6±4.0</td>
<td>23.4±2.6</td>
<td>25.1±5.2</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>56.8±4.9</td>
<td>40.3±5.7</td>
<td>36.0±5.4</td>
</tr>
<tr>
<td>Fibre packing density (g/cm³)</td>
<td>10</td>
<td>0.285±0.068</td>
<td>0.276±0.009</td>
<td>0.257±0.0066</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>0.595±0.045</td>
<td>0.539±0.057</td>
<td>0.474±0.017</td>
</tr>
<tr>
<td>Porosity (%)</td>
<td>10</td>
<td></td>
<td>79.2±0.63</td>
<td>80.79±4.93</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td></td>
<td>59.78±4.29</td>
<td>64.51±1.3</td>
</tr>
</tbody>
</table>
Table 4. Composition, dimension, porosity and mechanical properties of the membranes (wet).

<table>
<thead>
<tr>
<th></th>
<th>Ge B[22]</th>
<th>2PU10Ge</th>
<th>4PU10Ge</th>
<th>6PU10Ge</th>
<th>8PU10Ge</th>
<th>8PU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean fibre diameter (nm)</td>
<td>496±62</td>
<td>541±91</td>
<td>1045±97</td>
<td>1153±129</td>
<td>1216±112</td>
<td>348±87</td>
</tr>
<tr>
<td>Mean Core (PU) diameter (nm)</td>
<td>-</td>
<td>-</td>
<td>610±74</td>
<td>738±97</td>
<td>900±107</td>
<td>-</td>
</tr>
<tr>
<td>Approx. PU content (%)</td>
<td>0</td>
<td>48.2</td>
<td>58.5</td>
<td>65.5</td>
<td>73.4</td>
<td>100</td>
</tr>
<tr>
<td>Approx. Porosity (%)</td>
<td>-</td>
<td>-</td>
<td>57.1±4.6</td>
<td>63.3±1.3</td>
<td>60.6±4.7</td>
<td>44.2±2.5</td>
</tr>
<tr>
<td>Thickness (μm)</td>
<td>-</td>
<td>37.8±3.4</td>
<td>44.2±13.9</td>
<td>44.1±4.0</td>
<td>79.1±7.4</td>
<td>45.0±8.5</td>
</tr>
<tr>
<td>Young's Modulus 0-20% (MPa)</td>
<td>1.4±0.1</td>
<td>2.25±0.6</td>
<td>0.90±0.05</td>
<td>0.93±0.14</td>
<td>1.22±0.17</td>
<td>2.3±0.9</td>
</tr>
<tr>
<td>Ultimate Tensile Strength (MPa)</td>
<td>0.6±0</td>
<td>1.92±0.6</td>
<td>1.49±0.24</td>
<td>1.83±0.21</td>
<td>2.16±0.46</td>
<td>5.8±2.1</td>
</tr>
<tr>
<td>Strain at Break (%)</td>
<td>78±50</td>
<td>81±35</td>
<td>125±10</td>
<td>133±7</td>
<td>136±17</td>
<td>191±61</td>
</tr>
</tbody>
</table>
Table 5. The thickness of electrospun membranes used for diffusion tests and their corresponding
average relaxation time, \( \tau \), and effective diffusion coefficient \( (D_{\text{eff}}) \) as a function of electrospinning time.
n=5, \( p<0.05 \) between * and **. (n=5)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean of Thickness (µm)</th>
<th>mean relaxation time, ( \tau ) (min)</th>
<th>Effective diffusion coefficient, ( D_{\text{eff}} ) (mm²/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2PU10Ge-2.5'</td>
<td>17.99</td>
<td>5.08±0.67</td>
<td>5.84±0.82 ( \times 10^4 )</td>
</tr>
<tr>
<td>2PU10Ge-5'</td>
<td>20.53</td>
<td>7.92±2.27</td>
<td>4.51±1.52 ( \times 10^4 )</td>
</tr>
<tr>
<td>2PU10Ge-10'</td>
<td>25.61</td>
<td>43.99±2.22</td>
<td>9.5±0.47 ( \times 10^3 )</td>
</tr>
<tr>
<td>6PU10Ge-2.5'</td>
<td>16.03</td>
<td>8.01±0.54</td>
<td>3.27±0.23 ( \times 10^4 )</td>
</tr>
<tr>
<td>6PU10Ge-5'</td>
<td>17.43</td>
<td>21.10±3.85</td>
<td>1.38±0.28 ( \times 10^4 )</td>
</tr>
<tr>
<td>6PU10Ge-10'</td>
<td>20.23</td>
<td>29.24±8.39</td>
<td>1.18±0.29 ( \times 10^4 )</td>
</tr>
</tbody>
</table>
Figure 1: Morphology of 8PU10Ge co-axial fibre membranes as seen under TEM (A & B) and SEM (C to E). A) As-spun PU fibre, B) As-spun co-axial fibre showing PU-core and Ge-Shell, C) discreet as-spun fibres, D) membranes crosslinked by immersion in GA solution showing disruption of fibro-porous structure, and E) membranes crosslinked using GA vapour that maintained their fibro-porous structure, but with fibres crosslinked with neighbouring fibres at contact points.
Figure 2 FTIR spectra (A & B) and SEM images (C to F) of electrospun co-axial fibre membranes with non-crosslinked Ge shell (A) and with Ge shell washed off (B to F). Morphology: C) 2PU10Ge, D) 4PU10Ge, E) 6PU10Ge and F) 8PU10Ge.
Figure 2 (Continued)
Figure 3. SEM images showing the hollow fibres (with PU core removed by dissolving in THF) for A) 2PU10Ge, B) 4PU10Ge, C) 6PU10Ge and D) 8PU10Ge electrospun co-axial fibre membranes.
Figure 4. SEM images (A to D) and fibre diameter distribution histograms (E to H) for the different crosslinked co-axial fibre membranes 2PU10Ge (A & E), 4PU10Ge (B & F), 6PU10Ge (C & G) and 8PU10Ge (D & H) respectively. Average fibre diameters for each membrane were significantly different from all other membranes (p<0.05).
Figure 5. Pore size distributions for the co-axial fibre membranes 2PU10Ge, 4PU10Ge, 6PU10Ge and 8PU10Ge electrospun for A) 10 and A) 40 min.
Figure 6. Glucose diffusion across a) 2PU10Ge and b) 6PU10Ge membranes as a function of time, and thickness (electrospinning times of 2.5, 5 and 10 min). (c) the trends of the effective diffusion coefficient calculated from the diffusion measurement as a function of time, and thickness (electrospinning times of 2.5, 5 and 10 min). Data is represented as Mean ± SE of mean, n=5.
Figure 7. Effect of electrospun 2PU10Ge coaxial fibre membranes on the \textit{in vitro} function coil-type implantable glucose biosensor: % change in A) sensitivity and B) linearity normalised to that at day 7 before applying coating(s) as a function of time; and C) % change in sensitivity and linearity plotted as a function of sensor coating configurations.
Figure 8. Effect of electrospun 6PU10Ge coaxial fibre membranes on the *in vitro* function coil-type implantable glucose biosensor: a & b % change in sensitivity and linearity normalised to that at day 7 before applying coating(s) as a function of time; and c) the % change in sensitivity and linearity plotted as a function of sensor coating configurations. Each of * and ** indicate statistical difference from all other groups.
Figure 9. The morphology of coaxial fibre membranes after 12 weeks of immersion in PBS (pH 7.4) and intermittent sensor function tests of 2PU10Ge (A & B) and 6PU10Ge (C & D).
**Figure S1.** Cross-sectional dimensions for coaxial fibres electrospun on a flat plate collector as a function of increasing PU (core) feed solution concentrations (% wt/v), while maintaining the Ge (shell) feed solution concentration. For each of total fibre diameter, PU core diameter, total Ge shell thickness and PU core diameter to total Ge shell thickness ratios, statistical differences were observed between each of the membrane configurations (±SD, n=160) (p<0.05).
Figure S2. The contact angle for A) GTA crosslinked Ge film and B) 2PU10Ge coaxial fibre membrane. The surface wetting for 4PU10Ge, 6PU10Ge and 8PU10Ge membranes was too quick to take any contact angle measurements.
Figure S3. Plots of a) Typical stress-strain curves based on uniaxial tensile tests for the different electrospun co-axial fibre membranes compared to that of 8PU and b) tensile properties of the electrospun coaxial-fibre membranes, n=4. 8PU membranes were used as controls. (Mean±SE).