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Bovine Serum Albumin Adsorbed PGA-co-PDL Nanocarriers for Vaccine Delivery via Dry Powder Inhalation

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ABSTRACT

Purpose Dry powder vaccine delivery via the pulmonary route has gained significant attention as an alternate route to parenteral delivery. In this study, we investigated bovine serum albumin (BSA) adsorbed poly(glycerol adipate-co- ω -pentadecalactone), PGA-co-PDL polymeric nanoparticles (NPs) within L-leucine (L-leu) microcarriers for dry powder inhalation.

Methods NPs were prepared by oil-in-water single emulsion-solvent evaporation and particle size optimised using Taguchi's design of experiment. BSA was adsorbed onto NPs at different ratios at room temperature. The NPs were spray-dried in aqueous suspension of L-leu (1:1.5) using a Büchi-290 mini-spray dryer. The resultant nanocomposite microparticles (NCMPs) were characterised for toxicity (MTT assay), aerosolization (Next Generation Impactor), in vitro release study and BSA was characterized using SDS-PAGE and CD respectively.

Results NPs of size 128.50 ± 6.57 nm, PDI 0.07 ± 0.03 suitable for targeting lung dendritic cells were produced. BSA adsorption for 1 h resulted in 10.23 ± 1.87 μ g of protein per mg of NPs. Spray-drying with L-leu resulted in NCMPs with $42.35 \pm 3.17\%$ yield. In vitro release study at 37°C showed an initial burst release of $30.15 \pm 2.33\%$ with $95.15 \pm 1.08\%$ over 48 h. Aerosolization studies indicated fine particle fraction (FPF%) < 4.46 μm as $76.95 \pm 5.61\%$ and mass median aerodynamic diameter (MMAD) of 1.21 ± 0.67 μm . The cell viability was $87.01 \pm 14.11\%$ (A549 cell line) and $106.04 \pm 21.14\%$ (16HBE14o-cell line) with L-leu based NCMPs at 1.25 mg/ml concentration after 24 h treatment. The SDS-PAGE and CD confirmed the primary and secondary structure of the released BSA.

Conclusions The results suggest that PGA-co-PDL/L-leu NCMPs may be a promising carrier for pulmonary vaccine delivery due to excellent BSA adsorption and aerosolization behaviour.

KEY WORDS dry powder inhalation · nanoparticles · pulmonary delivery · spray drying · vaccines

ABBREVIATIONS

APCs	Antigen presenting cells
BSA	Bovine serum albumin
DCs	Dendritic cells
DoE	Design of experiment
LN	Lymph node
NPs	Nanoparticles
NCMPs	Nanocomposite microparticles
PGA-co-PDL	Poly(glycerol adipate-co- ω -pentadecalactone)
PLA	Poly(lactide or poly-L-lactic acid)
PLGA	Poly(lactic-co-glycolic-acid)
PVA	Poly(vinyl alcohol)
SD	Spray-drying

INTRODUCTION

Vaccination refers to induction of an immune response using antigens coupled with adjuvants for generating a protective immunity against plausible future infections (1,2). Traditional vaccines are often administered *via* the parenteral route requiring infrastructure such as cold-chain, sterilized water for reconstitution of dry powder vaccines and trained medical personnel. Lack of these facilities in low and middle income countries (LMICs) is leading to many eligible children and adults not getting vaccinated (3). Moreover, the majority of the potential vaccines in development employ purified subunits or recombinant proteins that are often poorly immunogenic thus needing adjuvants and effective delivery systems to generate an optimal immune response (1,2). To address these issues, particulate delivery systems and non-invasive routes of delivery are being investigated. The pulmonary route has gained significant attention for delivery of vaccines as it is one of the main entry portals for pathogens, and can address some of the challenges such as invasiveness, cold-chain requirement, and stability of the antigen by delivering the antigen as a dry powder (3).

Biodegradable polymeric nanoparticles (NPs) have gained significant attention and are largely being explored as delivery vehicles for delivery of peptides, proteins, antigens, DNA *etc.* (3–5). These polymers offer controlled or sustained drug release, biocompatibility with surrounding cells and tissues, degrade into low molecular weight non-toxic products and act as adjuvants helping in generating cellular and humoral immune responses (1,3,6). In this current investigation we aim to use poly(glycerol adipate-co- ω -pentadecalactone), PGA-co-PDL, a biodegradable polyester polymer, that has extensively been studied by our group for delivery of both small molecule and model drugs (dexamethasone phosphate, ibuprofen, sodium fluorescein), and large molecule drugs (α -chymotrypsin, DNase I) (7–10).

In these biodegradable polymeric nanoparticulate formulations, the vaccine antigens (*i.e.* proteins, peptides *etc.*) are either adsorbed onto the surface or encapsulated within the particles (3). Encapsulated antigens are protected by polymeric NPs and their release can be modified by tailoring the properties of the polymers. Adsorbed antigen, however, offers enhanced stability and activity over the encapsulated antigen by avoiding contact with organic solvents employed during particle preparation steps (11–13).

Recent strategies for effective vaccine delivery have been to target the dendritic cells (DCs), the true professional antigen presenting cells (APCs) (14). DCs have the exceptional ability to internalize, and in lymph nodes (LNs) they process and present antigens through major histocompatibility complex (MHC) class I and II pathways thereby activating naïve T- cells resulting in induction of a strong immune response (3,14,15). A study conducted by Manolova *et al.* indicates the importance of particle size in determining the uptake by DCs, where it was shown that upon intracutaneous injection of polystyrene beads of varying sizes, large particles (500– 2,000 nm) associated with DCs from the site of injection whereas small particles (20–200 nm) drained freely to the LNs and were present in LN resident DCs (15). In addition, Kim *et al.* have shown that uptake of 200 nm sized NPs by bone marrow DCs to be more than that of 30 nm sized NPs (16). Furthermore, Foged *et al.* has shown that particle size of 500 nm or below were preferred and have shown fast and efficient uptake by human DCs derived from blood (17). The above literature suggests that smaller particles of 200 to 500 nm could effectively be up taken by DCs and thus generates a stronger immune response compared to vaccine alone. However, these studies cannot be directly compared to lung DCs but owing to the lack of information on the effect of NP size on uptake by lung DCs the same can be assumed.

In this study, the Taguchi L18 orthogonal array design of experiment (DoE) was used to optimize the formulation parameters to achieve NPs (~150 nm) for targeting the lung DCs. The literature suggests that factors such as molecular weight (MW) of the polymer, organic solvent, aqueous phase, sonication time and stirrer speed have an influence on the size of the resultant NPs (18–20) and these were evaluated using the experimental design. As a dry powder, the nanosized particles cannot be directly used for inhalation as their size is too small and it is expected that majority of the inhaled dose will be exhaled depositing very minimal doses in the lung (21). Thus, these NPs are formulated into nanocomposite microparticles

(NCMPs) using additives such as lactose (22), L-leucine (L-leu) (22,23), trehalose (24), mannitol (24) by various manufacturing techniques such as freeze drying, spray drying (SD), spray-freeze drying or super-critical fluid technologies (3,25). The NCMPs in the size range of 1 to 5 μm in diameter are reported to be deposited in the respirable airways and periphery of the lung (26). The additives used to form NCMPs dissolve upon encountering the respiratory environment thereby releasing the NPs (27).

In this project, we aim to produce PGA-co-PDL NPs of optimum size to be effectively taken up by the DCs, surface adsorb a model protein, bovine serum albumin (BSA) and formulate into nanocomposite microparticles (NCMPs) using L-leu as a carrier for delivery via dry powder inhalation.

MATERIALS AND METHODS

Materials

Dichloromethane (DCM) was purchased from BDH, laboratory supplies, UK. Novozyme 435 (a lipase from *Candida antarctica* immobilized on a microporous acrylic resin) was purchased from Biocatalytics, USA. Acetonitrile (HPLC grade), albumin tagged with fluorescein isothiocyanate (FITC-BSA), bovine serum albumin (BSA, MW 67 KDa), phosphate buffered saline (PBS, pH 7.4) tablets, poly(vinyl alcohol) (PVA, MW 9–10 KDa, 80%), trifluoroacetic acid (TFA, HPLC grade), RPMI-1640 medium with L-glutamine and NaHCO_3 , thiazolyl blue tetrazolium bromide (MTT), tween 80[®] and ω -pentadecalactone were obtained from Sigma-Aldrich, UK. L-leucine (L-leu) was purchased from BioUltra, Sigma, UK. Seventy-five cm^2 /tissue culture flasks with vented cap, 96-well flat bottom plates, acetone, antibiotic/antimycotic solution (100X), dimethyl sulfoxide (DMSO) were purchased from Fisher Scientific, UK. Divinyladipate was obtained from Fluorochem, UK. Fetal calf serum (FCS) heat inactivated was purchased from Biosera UK. poly(glycerol adipate-co- ω -pentadecalactone) (PGA-co-PDL), MW of 14.7, 24.0 KDa was synthesized in our laboratory at LJMU and micro BCA[™] protein assay kit was purchased from Thermo Scientific, UK. A549 cell line was purchased from ATCC. 16HBE14o- cells were obtained from Dr Dieter Gruenert from the California Pacific Medical Center, University of California San Francisco, USA.

Polymer Synthesis

The PGA-co-PDL polymer of MW of 14.7, 24.0 KDa was synthesized in our laboratory via enzyme catalyzed co-polymerization of three monomers as described by Thompson *et al.* (28). The synthesized linear polyester was characterized by gel permeation chromatography, GPC (Viscotek TDA Model 300 using OmniSEC3 operating software), pre-calibrated with polystyrene standards (polystyrene standards kit, Supelco, USA), and ¹H-NMR spectroscopy (Bruker AVANCE 300 MHz, Inverse probe with B-ACS 60, Auto sampler with gradient chemming) as described by Thompson *et al.* (28).

Preparation of Nanoparticles

PGA-co-PDL NPs were fabricated using a modified oil-in-water (o/w) single emulsion solvent evaporation method (29). Briefly, 200 mg PGA-co-PDL polymer (MW 14.7 KDa) (and Nile Red, NR 0.5 mg for characterization of protein adsorption onto the surface of PGA-co-PDL via confocal microscopy) was dissolved in 2 ml DCM and probe sonicated (20 μm amplitude) upon addition to 5 ml of 10%w/v poly(vinyl alcohol) (PVA) (1st aqueous solution) for 2 min to obtain an emulsion. This whole process was performed using ice. This was immediately added drop wise to 20 ml of 2nd aqueous solution (0.75%w/v PVA) under magnetic stirring at a speed of 500 RPM. The whole mixture

was left stirring at room temperature for 3 h to facilitate the evaporation of DCM. The particle size, PDI and zeta-potential were then characterised as mentioned in “Nanoparticle Characterization” Section. The NP suspensions were collected by centrifugation (78,000 g, 40 min, 4°C) and surface adsorbed with protein as indicated in “Protein Adsorption” Section.

Taguchi Design of Experiment (DoE)

In order to evaluate the influence of formulation parameters and minimize the number of experiments, Taguchi DoE being appropriate to study large number of factors, was employed through Minitab® 16 Statistical Software. Seven factors, namely, polymer MW, organic solvent, internal aqueous phase concentration and volume, sonication time, stirrer speed and external aqueous phase concentration were evaluated by constructing and using L18 orthogonal array design with 1 factor, MW, at 2 levels and remaining 6 factors at 3 levels (Table I).

The design was applied to identify the significant factors that would affect the size of PGA-co-PDL NPs. Optimum conditions were indicated by high signal-to-noise (S/N) ratios, where signal factor (S) is the outcome, the particle size, and noise factors (N) are parameters such as humidity, temperature, experience of the experimenter *etc.* A greater S/N ratio corresponds to minimum variance of the outcome, the particle size *i.e.* a better performance. In other words, the experimental parameter having the least variability is the optimum condition (30). The optimization of size was carried out using the Taguchi’s ‘smaller-is-better’ criterion *i.e.* to get the outcome, the particle size, to an ideal target of zero or as small as possible.

Table I Taguchi’s Experimental Design L18 for Producing PGA-co-PDL Nanoparticles

Levels	Units	1	2	3
A—MW of Polymer	KDa	14.7	24.0	—
B—Org Sol (DCM)	ml	1	1.5	2
C—Aq. Vol (PVA)	ml	3	4	5
D—1 st Aq. Conc (PVA)	%w/v	2.5	5	10
E—Sonication Time	min	1	2	5
F—Stirrer Speed	RPM	400	500	600
G—2 nd Aq. Conc (PVA)	%w/v	0.5	0.75	1

Protein Adsorption

NP suspension (equivalent to 10 mg *i.e.* 1.25 ml of suspension) was centrifuged (78,000 g, 40 min, 4°C) and the resultant pellet was resuspended in vials containing 4 ml of BSA (or FITC-BSA for characterization of protein adsorption onto the surface of PGA-co-PDL *via* confocal microscopy), for 1 h, at different ratios of 100: 4, 100: 10 and 100: 20 (NP: BSA) corresponding to 100, 250 and 500 µg/ml BSA concentrations, respectively. The resulting suspension (for 100: 20) was left rotating for 30 min, 1, 2 and 24 h at 20 RPM on a HulaMixer™ Sample Mixer (Life Technologies, Invitrogen, UK). After respective time points, the protein adsorbed NP suspensions were centrifuged and the supernatant analysed for protein content using micro BCA protein assay kit. The amount of BSA adsorbed per milligram of NPs (*n*=3) was calculated using Eq. 1:

Adsorption (μg per mg NPs)

$$= \frac{(\text{Initial protein conc} - \text{Supernatant protein conc})}{\text{Amount of NPs}} \quad (1)$$

The particle size and PDI were then characterised as mentioned in “Nanoparticle Characterization” Section.

Nanoparticle Characterization

Particle size, poly dispersity index (PDI, a number between 0 and 1 describing the homogeneity of the sample) and zeta- potential were measured by laser diffraction using a laser particle size analyser (Zetasizer Nano ZS, Malvern Instruments Ltd, UK). For NPs suspension, an aliquot of 100 μl was diluted with 5 ml of deionized water and for NP suspensions with and without BSA adsorption, 2 mg of NPs were resuspended in 5 mL of deionized water, loaded into a cuvette and the measurements were recorded at 25°C ($n=3$).

Nanocomposite Microparticles

Spray-drying was employed to incorporate the NPs into nano-composite microparticles (NCMPs) using L-leu as a carrier, and at nanoparticles-to-carrier ratio of 1:1.5w/w. Blank NPs, BSA-loaded NPs or FITC-BSA-loaded NR NPs were dispersed in 20 ml water with L-leu dissolved and spray-dried using a Büchi B-290 mini spray-dryer (Büchi Labortechnik, Flawil, Switzerland) with a nozzle atomizer, and nozzle orifice diameter of 0.7 mm. The SD was performed at a feed rate 10%, an atomizing air flow of 400 L/h, aspirator capacity of 100% and an inlet temperature of 100°C (outlet temperature approximately 45–47°C). The dry particles (PGA-co-PDL/L-leu NCMPs) were separated from the air stream using a high-performance cyclone (Büchi Labortechnik), and the dry particles were collected and stored in desiccator until further use.

Nanocomposite Microparticles Characterization

Yield

The dry powder yield was determined as the difference in the weight of the sample vial before and after product collection. The weight difference was compared to the initial total dry mass and the yield in % (w/w) was calculated ($n=3$).

Particle Size and Morphology

To confirm the recovery of NPs from NCMPs with an appropriate size range for cellular uptake, particle size and PDI of NPs following re-dispersion of blank and loaded NCMPs in water were measured. The measurements were recorded as mentioned in “Nanoparticle Characterization” Section, where 5 mg of NCMPs were dispersed in 2 ml of deionized water then loaded into a cuvette and the measurements were recorded at 25°C ($n=3$).

Spray dried PGA-co-PDL/L-Leu NCMPs samples were mounted on aluminium stubs (pin stubs, 13 mm) layered with a sticky conductive carbon tab and coated with palladium (10–15 nm) using

a sputter coater (EmiTech K 550X Gold Sputter Coater, 25 mA for 3 min). The particles were then visualized using scanning electron microscopy (FEI Quanta™ 200 ESEM, Holland).

Confocal Laser Scanning Microscopy

FITC-BSA-loaded NR NPs spray-dried into NCMPs were observed under confocal microscope to visualise the adsorption of BSA onto the NPs. Briefly, a Zeiss 510 Meta laser scanning microscope mounted on a Axiovert 200 M BP computer-controlled inverted microscope was used to obtain the confocal images. A few milligrams of spray-dried NCMPs were placed in a single well of 8-well chambered (Fisher Scientific, UK) and imaged by excitation at a wavelength of 488 nm (green channel for FITC-BSA), 543 nm (red channel for Nile Red NPs) and a Plan Neofluar 63×/0.30 numerical aperture (NA) objective lens. Image analysis was carried out using the Zeiss LSM software.

Protein Quantification by HPLC

An HPLC method was developed to quantify the amount of BSA present in NCMPs. The chromatographic conditions were as follows: HPLC system Agilent 1100 series (Santa Clara, CA, USA) equipped with a column (Aeris 3.6 µm C4 200A Wide Pore 4.6 mm i.d. × 150 mm length), security cartridge of the same material (Phenomenex, UK) and soft-ware for data processing; mobile phase was composed of (A) 0.1% TFA in water and (B) 0.1% TFA in acetonitrile with a gradient flow of A/B from 80:20 to 35:65 in 25 min, post-time 6 min; flow rate of 0.8 ml/min; injection volume of 100 µl; run temperature 40°C; UV detection at 214 nm and BSA retention time of 14.4 min. BSA calibration curve was prepared by accurate dilution of a previously prepared stock solution (1 mg/ml) in HPLC water and PBS (pH 7.4) to obtain the following concentrations: 0.5, 1, 2.5, 5, 10, 25, 50, 100 and 200 µg/ml of BSA (n=9, R²=0.999). All solutions used in the process were filtered using 0.45 µm filters prior to use. Limit of Detection (LOD) in water—1.98 µg/ml, PBS—1.48 µg/ml and Limit of Quantification (LOQ) in water—3.24 µg/ml, PBS—3.20 µg/ml.

In Vitro Release Studies

BSA adsorbed PGA-co-PDL/L-leu NCMPs (20 mg) were transferred into Eppendorf's and dispersed in 2 ml of PBS, pH 7.4. The samples were incubated at 37°C and left rotating for 48 h at 20 RPM on a HulaMixer™ Sample Mixer (Life Technologies, Invitrogen, UK). At pre-determined time intervals up to 48 h, the samples were centrifuged (accuSpin Micro 17, Fisher Scientific, UK) at 17,000 g for 30 min and 1 ml of the supernatant removed and replaced with fresh medium. The supernatant was analysed using the HPLC method as mentioned above. Each experiment was repeated in triplicate and the result was the mean value of three different samples (n=3). The percentage cumulative BSA released was calculated using Eq. 2:

$$\% \text{ Cumulative BSA released} = \frac{\text{Cumulative BSA released}}{\text{BSA loaded}} \times 100 \quad (2)$$

Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis (SDS-PAGE)

The primary structure of BSA released from the NCMPs after SD was determined by SDS-PAGE. SDS-PAGE was performed on CVS10D omniPAGE vertical gel electrophoresis system (Geneflow Limited, UK) with 9% stacking gel prepared using ProtoGel stacking buffer (Geneflow Limited, UK) containing 0.4% of SDS. Protein molecular weight markers in the range 10–220 KDa (Geneflow

Limited, UK) and BSA were used as control. The protein loading buffer blue (2X) (0.5 M Tris-HCl (pH 6.8), 4.4% (w/v) SDS, 20% (v/v) Glycerol, 2% (v/v) 2-mercaptoethanol and bromophenolblue in distilled/deionised water) was added to the samples in 1:1 (v/v) buffer-to-sample ratio. After loading the samples (25 µl sample per well), the gel was run for approximately 2.5 h at a voltage of 100 V with Tris-Glycine-SDS PAGE buffer (10X) (Geneflow Limited, UK) containing 0.25 M Tris base, 1.92 M glycine and 1% (w/v) SDS. The gel was stained with colloidal coomassie blue and then destained in distilled water overnight. An image of the gel was scanned on a gel scanner (GS-700 Imaging Densitometer, Bio-Rad) equipped with Quantity One software.

Circular Dichroism (CD)

The secondary structure of standard BSA (control), BSA supernatant (after 1 h adsorption followed by centrifugation) and BSA released from NPs after 48 h was determined by measuring circular dichroism spectra. All CD experiments were performed using a J-815 spectropolarimeter (Jasco, UK) at 20°C as previously described (31). Five scans per sample using a 10 mm path-length cell were performed over a wavelength range 260 to 180 nm at a data pitch of 0.5 nm, band width of 1 nm and a scan speed 50 nm min⁻¹. Far-UV CD spectra were collated for standard BSA, supernatant BSA in HPLC grade water, BSA released in PBS for 48 h. For all spectra, the baseline acquired in the absence of sample was subtracted (32). The secondary structure of the samples was estimated using the CDSSTR method (33) from the DichroWeb server (33–35).

Activity

The activity of BSA was investigated using 4-nitrophenyl acetate esterase substrate (NPA, Sigma Aldrich, UK) as described by Abbate et al. (36). Briefly, 1.2 ml of released BSA sample (50 µg/ml) in PBS was added to freshly prepared NPA solution (15 µl of a 5 mM solution in ACN) and incubated for 1 h using HulaMixer™ Sample Mixer. Thereafter, the solution was transferred to a plastic cuvette and absorbance measured at 405 nm. For positive control, standard BSA (50 µg/ml) was treated exactly the same as released sample whereas for negative control, PBS buffer alone was treated as sample. The relative residual esterolytic activity of the samples was calculated as the ratio of absorbance between the released BSA/standard BSA, with the esterolytic activity obtained for standard BSA considered to be 100%.

In Vitro Aerosolization Studies

The Next Generation Impactor (NGI) was employed to assess the aerosol performance of spray-dried NCMPs. The BSA adsorbed NCMPs were weighed (4 capsules each corresponding to 12.5 mg spray-dried, powder equivalent to 5 mg of NPs) and manually loaded into the hydroxypropyl methylcellulose, HPMC, capsule (size 3), and placed in a Cyclohaler® (Teva pharma). The samples were drawn through the induction port into the NGI using a pump (Copley Scientific, Nottingham, UK) operated at a flow rate of 60 L/min for 4 s. The plates were coated with 1% tween 80: acetone solution and samples collected using a known volume of distilled water, and left on a roller-shaker for 48 h for the BSA to be released from NCMPs. The samples were centrifuged using an ultracentrifuge (as mentioned in “Preparation of Nanoparticles” Section) and the supernatants analysed using HPLC method as mentioned above to determine the amount of BSA deposited. The Fine Particle Fraction (FPF, %) was determined as the fraction of emitted dose deposited in the NGI with $d_{ae} < 4.46 \mu\text{m}$, the mass median

aerodynamic diameter (MMAD) was calculated from log-probability analysis, and the fine particle dose (FPD) was expressed as the mass of drug deposited in the NGI with $d_{ae} < 4.46 \mu\text{m}$ ($n=3$).

Cell Viability Study

The *in vitro* cytotoxicity of the empty PGA-co-PDL/L-leu NCMPs was evaluated using the MTT assay. The adenocarcinomic human alveolar basal epithelial cell line, A549 (passage no. 32) or 16HBE14o- cells (passage no. 32) were seeded in 100 μl (2.5×10^5 cells/ml) of RPMI-1640 medium supplemented with 10% foetal calf serum/1% Antibiotic/Antimycotic solution (complete medium) in 96-well plates and placed in an incubator at 37°C for 24 h supplemented with 5% CO₂. Then, 100 μl of freshly prepared NCMP dispersions in complete medium were added to the wells to an appropriate concentration (0–2.5 mg/ml) ($n=3$), and 10% dimethyl sulfoxide (DMSO) as a positive control. The formulations were assayed for toxicity over 24 h of incubation, followed by the addition of 40 μl of a 5 mg/ml MTT solution in PBS to each well. After 2 h of incubation at 37°C, the culture medium was gently removed and replaced by 100 μl of dimethyl sulfoxide in order to dissolve the formazan crystals. The absorbance of the solubilised dye, which correlates with the number of living cells, was measured at 570 nm using a plate reader (Molecular Devices, SpectraMAX 190). The percentage of viable cells in each well was calculated as the absorbance ratio between nanoparticle-treated and untreated control cells.

Statistical Analysis

All statistical analysis was performed using Minitab® 16 Statistical Software. One-way analysis of variance (ANOVA) using Minitab® 16 Statistical Software with the Tukey's comparison was employed for comparing the formulations with each other. Statistically significant differences were assumed when $p < 0.05$. All values are expressed as their mean $\pm$ standard deviation.

RESULTS

Polymer Synthesis

The synthesized PGA-co-PDL co-polymer (monomer ratio, 1:1:1) was a white powder with a molecular weight of 14.7,24.0 KDa for 6 h, 24 h as determined by the GPC. The integration pattern of the co-polymer was confirmed by ¹H- NMR spectra, (δ H CDCl₃, 300 MHz): 1.34 (s, 22 H, H-g), 1.65 (m, 8 H, H-e, e', h), 2.32 (m, 6 H, H-d, d', i), 4.05 (q)-4.18 (m) (6 H, H-a, b, c, f), 5.2 (s, H, H-j).

Taguchi Design of Experiment (DoE)

The Taguchi design was applied in this study to identify the significant factors that would influence the size of PGA-co- PDL NPs. Considering seven factors (1 factor at 2 levels and 6 factors at 3 levels) to be investigated, non-usage of an experimental design would have resulted in $2 \times 3^6 = 1,458$ individual experiments which would be an arduous task and inefficient.

The Taguchi L18 orthogonal array design resulted in 18 runs to be performed to yield the optimum conditions for each factor in achieving the smallest PGA-co-PDL NP size. Table II illustrates the structure of the L18 orthogonal array, the corresponding results and S/N ratios. The results obtained from 18 runs indicated particle sizes ranging from 138.7 ± 6.4 nm (run 3) to 459.4 ± 69.5 nm (run 11). Figure 1 shows the mean S/N graph of the particle size for each parameter level. The parameter with the largest range and corresponding rank (indicates the relative importance compared to other parameters) was considered as the critical factor affecting that particle size.

Analysis of the particle sizes of 18 runs using the Taguchi's 'smaller-is-better' criterion in Minitab® 16 Statistical Software, the optimum conditions inferred from the range, rank and the S/N response graph were A1B3C3D3E2F2G2. The optimum formulation made using these conditions yielded NPs with a size of 128.50 ± 6.57 nm lower than the minimum size of 138.7 ± 6.4 nm obtained using run 3, PDI of 0.07 ± 0.03 and zeta-potential of -10.2 ± 3.75 mV.

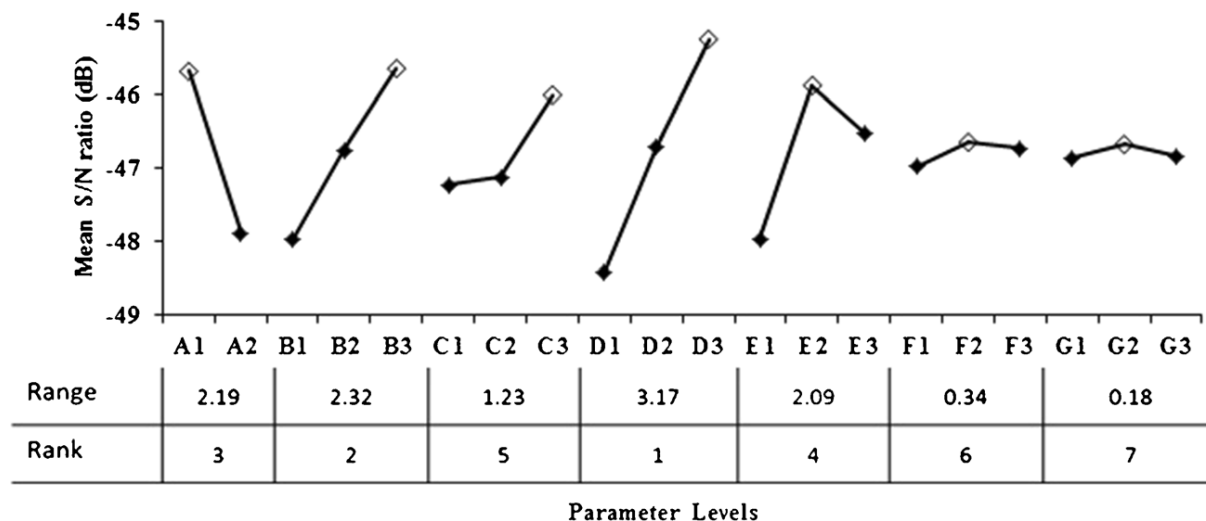


Fig. 1 Mean S/N graph for particle size response. Letters (A–E) denote the experimental parameters and numeric values denote the parameter levels (white diamond indicates maximum S/N value) (Mean $\pm$ SD, n=6); Note: A—MW of Polymer, B—Org Sol (DCM), C—Aq. Vol (PVA), D—1st Aq. conc (PVA), E— Sonication time, F—Stirrer Speed and G—2nd Aq. conc (PVA).

Table II Structure of Taguchi's L_{18} Orthogonal Array, Corresponding Particle Size and S/N Ratios (Mean $\pm$ SD, $n=6$)

Runs	Parameters							ParticleSize (nm)	PDI	S/NRatio (dB)
	A	B	C	D	E	F	G			
Run 1	1	1	1	1	1	1	1	315.5 $\pm$ 6.90	0.151 $\pm$ 0.05	-49.979
Run 2	1	1	2	2	2	2	2	186.5 $\pm$ 4.30	0.097 $\pm$ 0.03	-45.412
Run 3	1	1	3	3	3	3	3	138.7 $\pm$ 6.40	0.093 $\pm$ 0.01	-42.843
Run 4	1	2	1	1	2	2	3	210.1 $\pm$ 18.7	0.116 $\pm$ 0.04	-46.449
Run 5	1	2	2	2	3	3	1	208.7 $\pm$ 49.9	0.123 $\pm$ 0.06	-46.389
Run 6	1	2	3	3	1	1	2	182.0 $\pm$ 3.20	0.075 $\pm$ 0.04	-45.199
Run 7	1	3	1	2	1	3	2	192.9 $\pm$ 9.30	0.077 $\pm$ 0.04	-45.705
Run 8	1	3	2	3	2	1	3	149.3 $\pm$ 2.50	0.075 $\pm$ 0.01	-43.481
Run 9	1	3	3	1	3	2	1	192.9 $\pm$ 23.0	0.050 $\pm$ 0.03	-45.704
Run 10	2	1	1	3	3	2	2	269.1 $\pm$ 68.9	0.205 $\pm$ 0.04	-48.598
Run 11	2	1	2	1	1	3	3	459.4 $\pm$ 69.5	0.233 $\pm$ 0.02	-53.243
Run 12	2	1	3	2	2	1	1	242.5 $\pm$ 19.1	0.188 $\pm$ 0.06	-47.694
Run 13	2	2	1	2	3	1	3	253.2 $\pm$ 47.3	0.155 $\pm$ 0.10	-48.069
Run 14	2	2	2	3	1	2	1	217.3 $\pm$ 18.9	0.116 $\pm$ 0.01	-46.742
Run 15	2	2	3	1	2	3	2	240.5 $\pm$ 35.1	0.133 $\pm$ 0.05	-47.622
Run 16	2	3	1	3	2	3	1	169.3 $\pm$ 7.60	0.144 $\pm$ 0.04	-44.573
Run 17	2	3	2	1	3	1	2	235.9 $\pm$ 29.6	0.119 $\pm$ 0.08	-47.453
Run 18	2	3	3	2	1	2	3	221.7 $\pm$ 11.0	0.150 $\pm$ 0.04	-46.915

A—MW of Polymer, B—Org Sol (DCM), C—Aq. Vol (PVA), D—1st Aq. conc (PVA), E—Sonication time, F—Stirrer Speed and G—2nd Aq. conc (PVA)

Protein Adsorption

Figure 2a shows the amount of BSA adsorbed per mg of NPs for different concentrations of BSA loaded. The average adsorption of BSA, μg per mg of NPs, increased significantly from 100: 4 (NP: BSA) loading concentration (4.75 ± 0.39), 100: 10 (6.59 ± 1.28) to 100: 20 (10.23 ± 1.87) ($p < 0.05$, ANOVA/Tukey's comparison). Figure 2b shows the amount of BSA adsorbed in μg per mg of NPs at different time points for 100: 20 (NP: BSA) loading concentration. The average adsorption increased significantly from 30 min (1.84 ± 0.82) to 1 h (10.23 ± 1.87) ($p < 0.05$, ANOVA/Tukey's comparison) with no significant difference beyond 1 h compared to that of 2 h (8.76 ± 0.34) and 24 h (8.95 ± 0.39) ($p > 0.05$, ANOVA/Tukey's comparison) indicating maximum adsorption at 1 h. Table III lists the particle size and PDI of PGA-co-PDL NPs with and without BSA adsorption. As seen, there is a significant increase ($p < 0.05$, ANOVA/Tukey's comparison) in size which is attributed to the adsorption of BSA onto NPs as confirmed using confocal microscopy ("Confocal Laser Scanning Microscopy" Section).

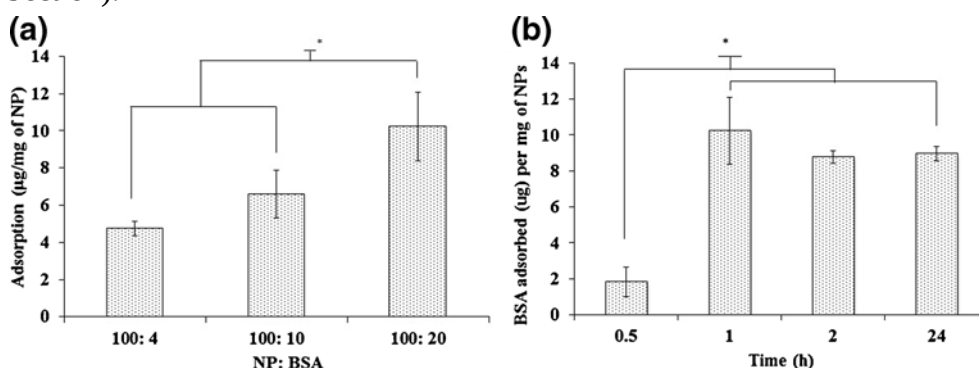


Fig. 2 Amount of BSA adsorbed per mg of NPs at different (a) BSA loading concentrations (NP: BSA) and (b) time points for 100: 20 (NP: BSA) BSA loading concentration, * is $p < 0.05$, ANOVA/Tukey's comparison (Mean $\pm$ SD, $n=3$).

Table III Particle Size and PDI of PGA-co-PDL Nanoparticles Without and with BSA Adsorption

	NP suspension	Without BSA adsorption	With BSA adsorption
Particle Size (nm)	128.50 ± 6.57 ^a	203.9 ± 2.55 ^{b*}	299.03 ± 32.02 ^{c*}
PDI	0.070 ± 0.030	0.205 ± 0.007	0.322 ± 0.060

* $p < 0.05$, ANOVA/Tukey's comparison

^a NPs characterised immediately after preparation without centrifugation, ^b NPs characterised after centrifugation but without adsorption of BSA, ^c NPs characterised after centrifugation and BSA adsorption.

Nanocomposite Microparticles Characterization

Yield

A reasonable yield of SD, 40.36 ± 1.80% for the empty PGA-co-PDL NCMPs and 42.35 ± 3.17% for the BSA adsorbed PGA-co-PDL/L-leu NCMPs was obtained.

Particle Size and Morphology

The size of NPs after recovery from spray-dried blank NCMPs in water was 210.03 ± 15.57 nm and PDI 0.355 ± 0.067 and for that of BSA loaded NCMPs was 282.46 ± 2.17 nm and PDI 0.36 ± 0.008, which is in the range of 200 to 500 nm for uptake by dendritic cells (DCs) (16–18). The shape and surface texture of NCMPs were investigated using scanning electron microscopy (Fig. 3). Photomicrographs of NCMPs showed irregular and corrugated microparticles.

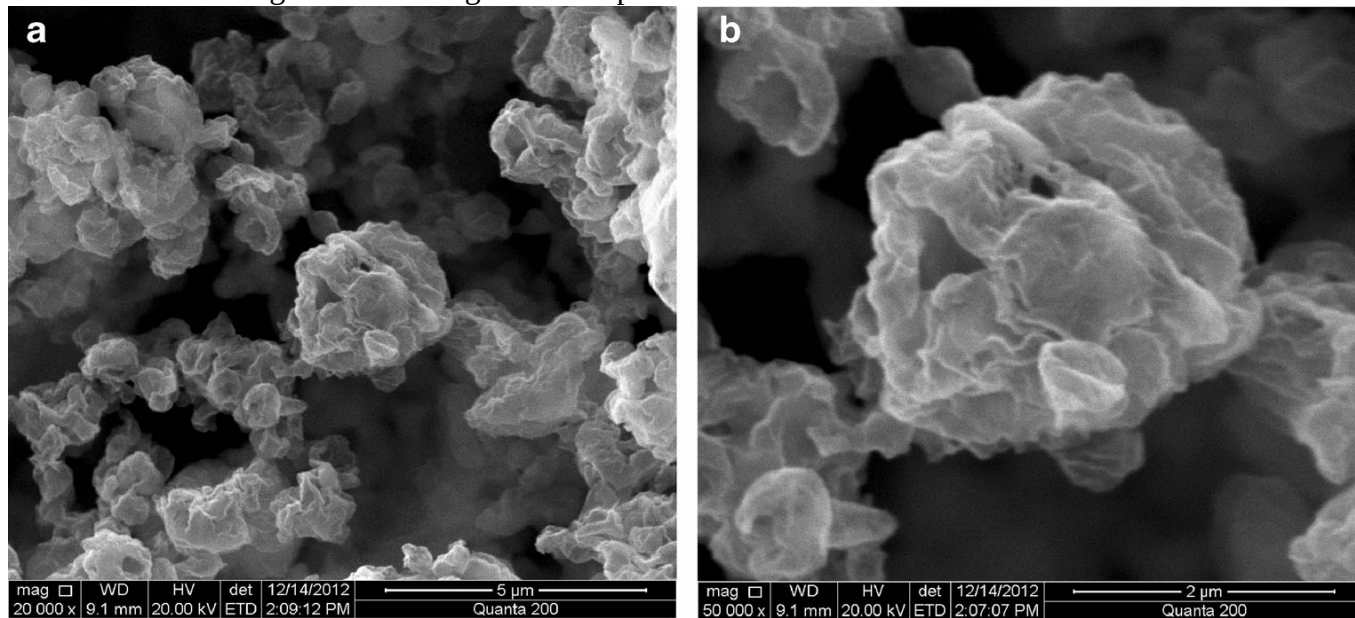


Fig. 3 SEM pictures of PGA-co-PDL/L-Leu Nanocomposite Microparticles (a) 5 μm and (b) 2 μm.

Confocal Laser Scanning Microscopy

CLSM was used to observe the interaction of BSA with NPs. The microscopic images in Fig. 4a (split view) and 4b (orthogonal view) shows the spray-dried NCMPs containing the fluorescent NPs (red, labelled using NR dye) adsorbed with FITC-BSA (green). The image shows that FITC-BSA was evidently only present where the NPs were present, indicating their association. Moreover, the increase in size observed after adsorption also confirms the adsorption of BSA onto PGA-co-PDL NPs (Table

III).

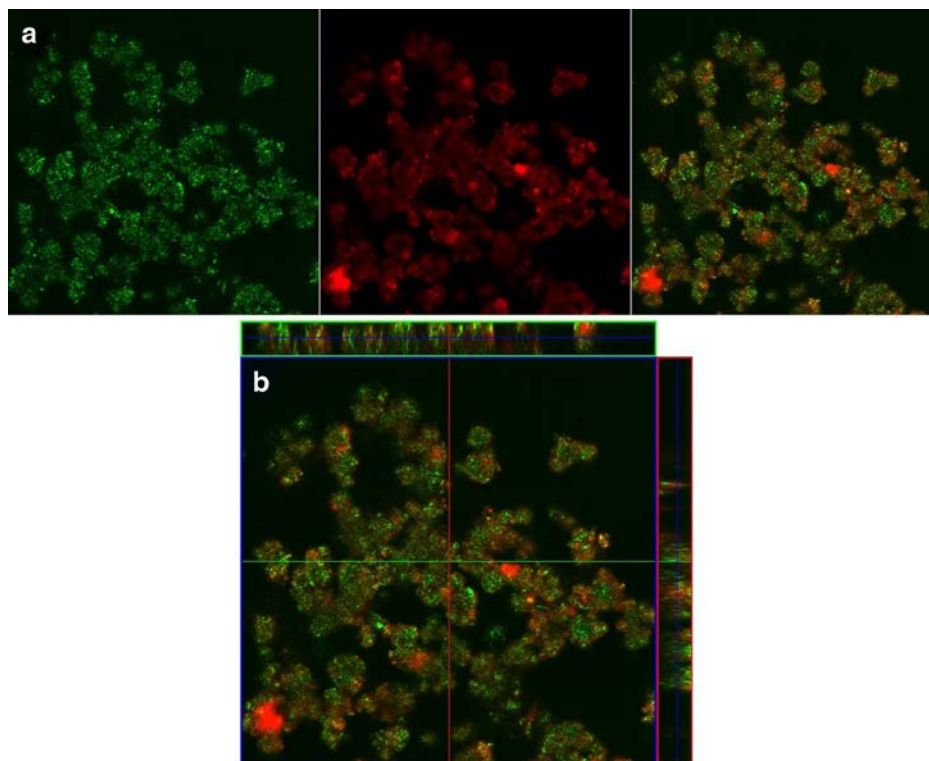


Fig. 4 Confocal microscopic image of spray-dried microparticles containing the fluorescent nanoparticles (red, labelled using Nile red dye) adsorbed with FITC-BSA (green) (a) Split view and (b) Orthogonal view.

In Vitro Release Studies

In vitro release studies were performed on NCMPs and reported as cumulative percentage BSA released over time (Fig. 5). An initial burst release of $30.15 \pm 2.33\%$ (BSA) was observed followed by continuous release up to 5 h, with BSA release of $86.07 \pm 0.95\%$. After this time period, a slow continuous release of BSA was observed with release of $95.15 \pm 1.08\%$ over 48 h, indicating an excellent release profile for the PGA-co- PDL/L-leu NCMPs.

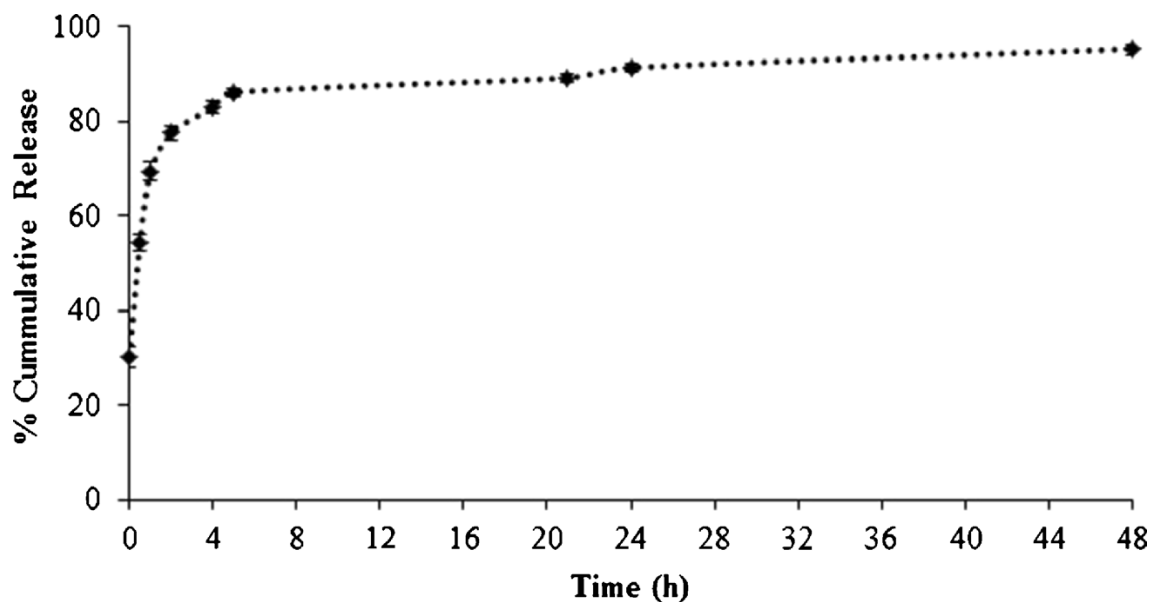


Fig. 5 In vitro release profiles for BSA adsorbed PGA-co-PDL/L-leu NCMPs in phosphate buffer

saline, pH 7.4 (Mean $\pm$ SD, n=3).

Protein Stability (SDS-PAGE and CD) and Activity

The primary structure of BSA released from the NCMPs was investigated using SDS-PAGE analysis. Figure 6 reveals identical bands for the standard BSA and desorbed BSA from NCMPs without any newly noticeable bands of high and low molecular BSA.

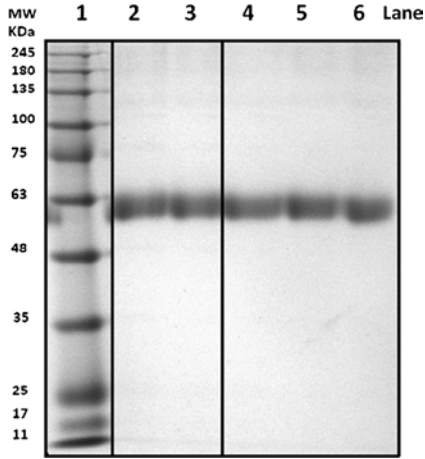


Fig. 6 SDS-PAGE of Lane 1: molecular weight standards, broad range (Bio-Rad Laboratories, Hercules CA, USA), Lane 2, 3: BSA standards, Lane 4, 5, 6: Desorbed BSA from PGA-co-PDL/L-leu NCMPs after 24 h.

The secondary structure analysis was performed using CD spectral data. Figure 7a and b shows the structure of standard BSA, BSA supernatant and BSA released. In Fig 7a, the CD spectra show minima at 221–222 and 209–210 nm and maximum at about 195 nm for both samples, which is characteristic of α -helical structure. Further structural analysis showed that the predominant structure of the protein was helical displaying 51 and 62.5% helicity respectively (Table IV). Moreover, the experimental data obtained for the standard BSA are in good agreement with previous reports (37). Figure 7b shows that BSA released displayed double minima at 208 and 222 nm and further spectra analysis indicated this sample adopted a reduced level helical conformation (*circa* 36% helical) (Table IV). Comparing the CD results of BSA released with that of standard BSA, the content of α -helix decreases by 15%, the β -sheet content increases by 8.9%, the turns content increases by 1%, and the random coils' content increases by 3%, respectively.

The residual esterolytic activity of the released BSA sample was calculated to be $77.73\pm3.19\%$ relative to standard BSA.

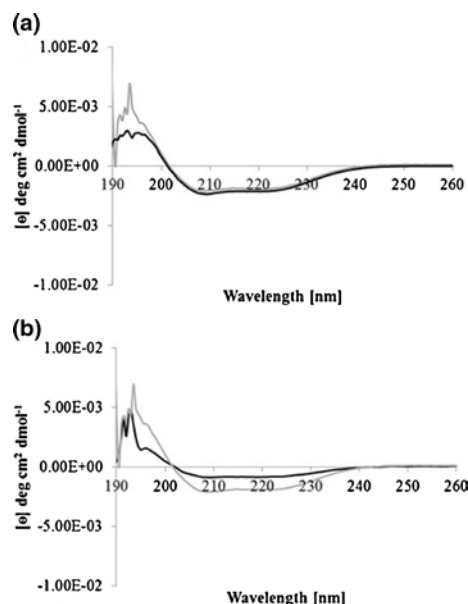


Fig. 7 CD spectra of (a) standard BSA (grey) and supernatant BSA (black) and (b) standard BSA (grey) and BSA released (black).

Table IV The Percentages of the Secondary Structures of Standard, Supernatant and Released BSA Samples

Sample	Helix	Strand	Turns	Unordered
Standard BSA	51 ± 0.007	21.1 ± 0.07	6.0 ± 0.01	18 ± 0.007
Supernatant BSA	62.5 ± 0.035	22.0 ± 0.021	5.5 ± 0.05	9.5 ± 0.06
Released BSA	36.0 ± 0	30.0 ± 0	7.0 ± 0	21 ± 0.007

The content and level of secondary structure elements in the peptide was calculated from spectral data using the DichroWeb server software as described in “Materials and Methods” Section.

In Vitro Aerosolization Studies

The deposition data obtained from spray-dried formulations displayed a FPD of $112.87 \pm 33.64 \mu\text{g}$, FPF of $76.95 \pm 5.61\%$ and MMAD of $1.21 \pm 0.67 \mu\text{m}$. This suggest that the BSA adsorbed PGA-co-PDL/L-leu NCMPs were capable of delivering efficient BSA to the lungs, and are expected to deposit the majority of the emitted dose to the bronchial-alveolar region of the lungs (3).

Cell Viability Study

The non-adsorbed PGA-co-PDL NCMPs appear to be well tolerated by both the cell lines, with a cell viability of $87.01 \pm 14.11\%$ (A549 cell line) and $106.04 \pm 21.14\%$ (16HBE14o- cell line) (Fig. 8) at 1.25 mg/ml concentration after 24 h exposure indicating a good toxicity profile without any significant difference in cell viability between particle loadings. This provides an indication about the feasibility of using PGA-co-PDL polymers as safe carriers for pulmonary drug delivery.

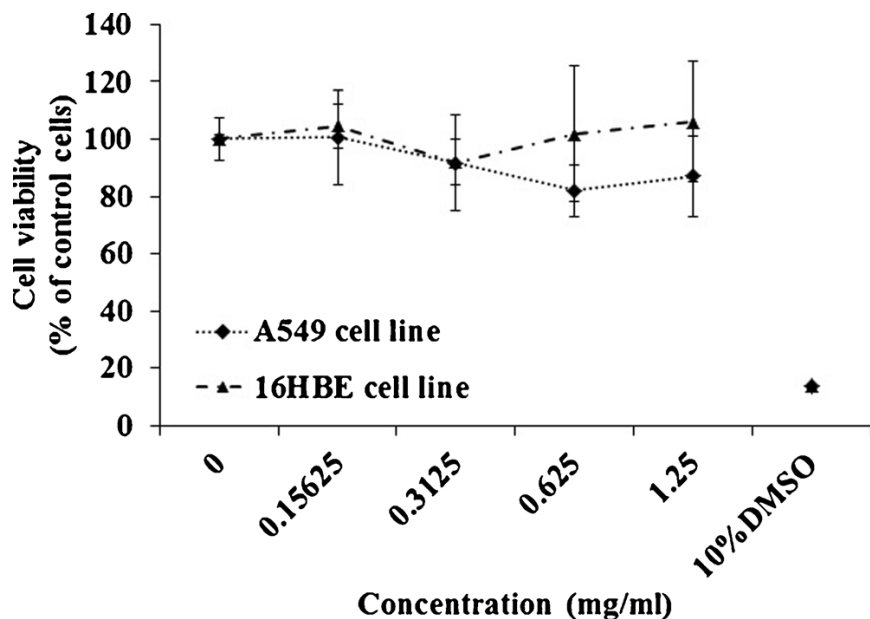


Fig. 8 A549 & 16HBE14o- cell viability measured by MTT assay after 24 h exposure to PGA-co-PDL NCMPs (Mean±SD, n=3).

DISCUSSION

Nanoparticle Preparation and Characterization

The PGA-co-PDL NPs were prepared using a modified oil-in water(o/w) single emulsion solvent evaporation method (29). The results of 18 runs, suggested by Taguchi's L18 orthogonal array, resulted in NPs of size<150 nm. However, this increases to about 200–300 nm after centrifugation and BSA adsorption. This according to the literature suggests an effective uptake by DCs (15–17). The effects of each factor are discussed in detail below:

Factor D, PVA concentration (range=3.17, rank=1), is the most important factor affecting the particle size. The S/N ratios at three levels indicated that particle size almost linearly decreased with increase in surfactant concentration from 2.5 to 10%w/v (S/N ratio, $r^2=0.997$). The particle size decreases because at lower concentrations there is inadequate amount of surfactant to cover all the surfaces of PGA-co-PDL NPs (38). The uncovered NPs then tend to aggregate until a point where there is adequate amount of surfactant to cover the total surface area of the aggregated NPs, and form a stable system leading to larger particles. However, with an increase in surfactant concentration it was possible to efficiently cover all the surfaces of NPs thereby stabilizing the system avoiding aggregation and resulting in smaller PGA-co-PDL NPs (38). This effect of decrease in particle size with an increase in surfactant concentration, PVA, was also observed by Mitra and Lin (39).

The S/N ratios of factor A, molecular weight of the polymer (range=2.19, rank=3), at two levels, suggested a directly proportional relationship with MW of polymer, *i.e.* the particle size decreases with a decrease in the MW of the polymer. This can also be evident from the lower particle size measurements observed using 14.7 KDa MW polymer (runs 1–9) relative to 24 KDa MW polymer (runs 10–18). As the MW of polymer increases, the viscosity of the polymeric solution also increases, thereby imposing difficulty in breaking them into smaller emulsion droplets when compared to lower MW polymer requiring lower efficiency to breakdown under similar conditions. This increase in size of the

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particles has also been observed by others and is reported to be associated with high MW polymers (5,40,41).

The S/N ratios of factor B, volume of organic solvent (DCM, range=2.32, rank=2), at three levels indicated that particle size almost linearly decreased with increase in volume from 1 to 2 ml (S/N ratio, $r^2=0.999$). This decrease in particle size is attributed to the decrease in viscosities of the polymer solution (keeping the amount and MW of polymer constant). This makes it easier to break into smaller emulsion droplets resulting in a decreased particle size as explained above. This effect could also be observed with factor C, volume of 1st aqueous phase (range=1.23, rank=5), where a decrease in volume increased the viscosity thereby resulting in an increase in particle size.

The S/N ratios for factor E, sonication time (range=2.09, rank=4), did not follow any particular trend; however, the 2nd level was found to be the optimum for achieving smaller particle size. The S/N ratios for parameters F, stirrer speed (the speed at which the magnetic bar was rotating for evaporation of DCM in the external phase) and G, 2nd PVA concentration have a low range of 0.34 and 0.18 respectively indicating that they have a minimal influence over the size of NPs produced. Therefore, the optimum conditions inferred result in NPs of size suitable for cellular uptake into DC as established in the literature (15–17,42,43).

The adsorption of BSA onto NPs is expected to be mainly driven by hydrophobic, electrostatic (ionic) interactions and hydrogen bonding (44). However, in this study as the NPs, evident from zeta-potential values, are negatively charged and BSA in water is also highly negatively charged (45) suggesting that the electrostatic interactions are minimal and that the adsorption process is dominated by the hydrophobic interactions and hydrogen bonding. The BSA adsorption onto NPs increased with an increase in NP: BSA ratio from 100: 4 to 100: 20, which was expected as the amount of BSA available for adsorption increased. Figure 2b suggests that the surface of NPs was saturated with BSA after 1 h suggesting maximum adsorption with 100: 20 (NP: BSA) BSA loading concentration.

Nanocomposite Microparticles Characterization

NCMPs were produced by SD using L-leu as a carrier and a dispersibility enhancer. The SEM pictures (Fig. 3) show irregular or wrinkled surface which is due to an excessive build-up of vapour pressure during water evaporation in the SD process and occurs with hydrophobic amino acids, such as L-Leu (8,46,47).

The release profile shows more than 90% of the BSA released within 48 h this is because of weaker hydrophobic interactions between BSA and NPs compared to the strong ionic interactions. Moreover, the identical bands observed for BSA standard and desorbed BSA from NCMPs suggests that protein has maintained its primary structure and was neither degraded nor affected by the adsorption and SD procedure.

The secondary structure of BSA in the formulation was analysed using CD spectroscopy, a valuable technique in analysing the protein structure (31). The BSA released samples confirms the presence of α -helix and β -sheets though decreased compared to standard BSA. However, in protein secondary structure, it is believed that the β -sheet structure is sometimes observed as a special α -helix only with two amino acid residues through stretching resulting from the breakage of hydrogen bond (37,48).

It is established that BSA possesses an enzyme-like activity with the ability to hydrolyse substrates

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such as p-nitrophenyl esters (36,49,50). In this study, the released BSA sample retained approximately 77% of relative residual esterolytic activity compared to standard BSA. A reduction in BSA activity to 60% was also observed by Abbate *et al.* when released from biohybrid hydrogels (36). The adsorption and desorption process of BSA could have influenced the structure (evident from a decreased helicity determined by CD) and thus the activity. However, the retention of 77% ester hydrolysis activity would encourage the exploration of the delivery system for further usage.

The FPF value suggests an excellent aerosolization performance and deep lung deposition profile. The surface activity of the relatively strong hydrophobic alkyl side chain of L-leu accumulating at the particle surface during SD reduces the surface free energy of the dry powder and cohesive inter-particulate interactions and this might be a plausible explanation for the enhanced dispersibility (8,23). In addition, the dispersibility enhancing property of L-leu resulting from its corrugated surface that reduces the contact points between particles leads to an improved aerosolization characteristic of powders (22,47,51). Similar reports have also demonstrated the enhanced aerosol performance with L-leu containing formulations (22,47,51,52). Moreover, the MMAD values show an efficient delivery of NCMPs containing BSA to the deep lungs mainly to the bronchial-alveolar region (3). A study by Todoroff *et al.* have shown that more intense specific immune responses could be achieved by targeting the antigen to the deep lungs than to the upper airways (53). Also, Menzel *et al.* have shown that upon inhalation of Pneumovax®, a pneumococcal polysaccharide vaccine, by healthy volunteers the vaccine deposited in the alveolar region displayed increased serum antibody levels compared to that deposited in the larger airways (54). Thus, this deposition to the deep lungs may generate stronger immune responses.

The NGI data suggests a deposition mainly in the bronchial-alveolar region of the lungs (3), thus the cell viability studies were performed on A549 cell line (adenocarcinomic human alveolar basal epithelial cells) and 16HBE14o- cell line (human bronchial epithelial cells). The results show that both the cell lines were tolerant to the NCMPs up to 1.25 mg/ml concentration encouraging further investigation in animals.

CONCLUSIONS

PGA-co-PDL NPs of appropriate size to target DCs were successfully produced using Taguchi L18 orthogonal array DoE. BSA adsorption onto NPs in the ratio of 100: 20 (NP: BSA) for 1 h at room temperature produced the maximum adsorption of BSA (10.23 ± 1.87 µg of protein per mg of NPs). The BSA adsorbed NPs were successfully spray-dried using L-leu into NCMPs producing a yield of $\pm 3.17\%$ and the NCMPs had irregular and corrugated morphology. The BSA released from the NCMPs was shown to be maintaining its structure under SDS- PAGE and CD analysis with 77% of relative residual esterolytic activity. Moreover, FPF of $76.49 \pm 6.26\%$ and MMAD of 1.21 ± 0.67 µm values indicate deep lung deposition with NCMPs showing a low toxicity profile. This study suggests that PGA-co-PDL NCMPs could be used as a novel carrier for pulmonary vaccine delivery.

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