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Development and evaluation of a novel intranasal spray for the delivery of amantadine
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The aim of this study was to develop and characterise an intranasal delivery system for amantadine (AMT). Optimal formulations (F) consisted of a thermosensitive polymer Pluronic® 127 (P127) and either carboxy methyl cellulose (CMC) or chitosan (CS) which demonstrated gel-transition at nasal cavity temperatures (34 °C ± 1 °C). Rheologically, the loss tangent (Tan δ) confirmed a three-stage gelation phenomena at 34 °C ± 1 °C and non-Newtonian behaviour. Storage of FCMC and FCS at 4 °C for 8 weeks resulted in repeatable release profiles at 34 °C when sampled, with a Fickian mechanism earlier on but moving towards anomalous transport by week 8. Polymers (P127, CMC and CS) demonstrated no significant cellular toxicity to human nasal epithelial cells up to 4 mg/mL and up to 1 mM for AMT (IC50: 4.5 mM ± 0.05 mM). FCMC and FCS demonstrated slower release across an in-vitro human nasal airway model (43-44 % vs 79 % ± 4.58 % for AMT). Using a human nasal cast model, deposition into the olfactory regions (potential nose-to-brain) was demonstrated upon nozzle insertion (5 mm) whereas tilting of the head forward (15°) resulted in greater deposition in the bulk of the nasal cavity.

KEYWORDS
Parkinson's disease; rheology; nasal; RPMI 2650; nose to brain; thermoresponsive;
INTRODUCTION

Parkinson’s disease (PD) is an ageing associated progressive neurological disorder and affects 1.5 % of the population over 65-years of age 1, with age-adjusted prevalence rates thought to be around 150 per 100,000 with a suggested mean onset in the 70’s 1. Ageing is the largest single risk factor for the development of PD and the pathogenesis of PD is composed of a range of events leading towards neuronal apoptosis, primarily of degradation of dopamine neurones in the sustantia nigra. The mainstay of clinical drug therapy focuses on increases the levels of dopamine (DA) in the brain, with oral dosing of levodopa being the most commonly used to pharmacological intervention treatment to reverse the symptoms of PD. It is also common to use adjunct therapies in conjunction with levodopa, such as the weak NMDA-type receptor antagonist amantadine, which can aid in blocking dopamine reuptake.

One of the major complications of PD is the emergence of dysphagia affecting the ability to ingest/swallow and is a thought to affect between 45-95 % of PD patients 1,2 and has significant implications for orally dosed therapeutics such as amantadine and other anti-Parkinsonian therapeutics agents. Furthermore the occurrence of bradykinesia and morning akinesia is often associated with delayed gastric emptying following the oral administration of anti-Parkinsonian therapeutics. This is again confounded by the use of dopaminergic and other concomitant medication interventions. At least 24% of PD patients are thought to suffer from gastroparesis, the delay in emptying of the stomach, which will inevitably impact upon clinical outcome of any orally administered medication. A more recent approach for therapeutic delivery is through the intranasal route to obtain systemic deposition as a practical alternative to oral and parenteral routes 3,4.

The primary function of the nasal cavity is respiration and olfaction. The nasal cavity (in humans) is divided in to two symmetrical halves by a septum with each cavity sub-dived into four areas namely vestibules, atrium, respiratory region and olfactory regions. With a surface area of 150 cm², total volume of 15-20 mL the nasal cavity provides an optimal absorption area for drugs into the systemic circulations which would inevitably enhance the bioavailability of the therapeutic agent, particularly those which demonstrate poor oral absorption (intestinal permeation) or significant first-pass metabolism (extra-hepatic and hepatic) 5,6. Although originally exploited for the delivery of locally acting agents (allergic of infections rhinitis, nasal pluposos and sinusitis) the highly vascularised nature of the nasal cavity has provided a portal for systemic delivery of small molecules and biomolecules 7. The nasal cavity can provide larger surface area for rapid systemic absorption due to its highly vascularised surface 8. This non-invasive route not only provides rapid onset of action, but often increased bioavailability in low doses as it bypasses the first pass metabolism 9.

Although intranasal drug delivery offers many advantages, one of the main limitations is the short residence time of the formulation in the nasal cavity, as a result of mucociliary clearance (MCC) 10. This movement involves the beating of mucocilia at a frequency of 1,000 strokes per minute and hence movement of mucus of 5 mm/minute 11. Thus the clearance of any nasally administered formulation further back towards to the nasopharynx is often a limiting factor in clinical administration and results in a short nasal residency half-life of 15-20 minutes 12.

An ideal formulation would be one that was capable of prolonging residence time within the nasal cavity and therefore enhance the bioavailability of the drug. To achieve this goal, the use of intelligent polymer based systems have now become more commonplace in this field and particularly the use of responsive systems which may respond to the nasal cavity temperature 13-22 or pH 23-27 have increased amongst researchers. Furthermore, the use of mucoadhesive polymers 24,27,33 to enhance the adhesion of the formulations onto the nasal mucosa has further demonstrated the ability to prolong contact time with the mucosa and hence increase bioavailability 10,15,18,23,34-36.

The primary aim of this study was to develop an intranasal thermoresponsive hydrogel based delivery system for the delivery of anti-Parkinson’s drugs, using amantadine as a model candidate drug. The study objectives were to utilise thermoresponsive polymers, for example Pluronic F127 (P127) 13, mucoadhesive polymers (carboxymethyl cellulose and chitosan) and surfactant (PEG4000 and PEG 12000) to aid and enhance formulation delivery and
retention within the nasal cavity using a multi-dose nasal spray and human nasal cast. In order to develop and assess the formulations a range of studies were conducted and included rheological, stability, biological compatibility and nasal targeting of the formulations.
METHODS

Materials
Eagle’s Minimum Essential Medium (MEM), Dulbecco’s phosphate buffered saline (PBS), L-glutamine 200 mM, non-essential amino acids (NEAA), penicillin/streptomycin and trypsin-EDTA solution were obtained from PAA Laboratories (Austria); fetal bovine serum (FBS) (Labtech Intl Ltd. East Essex, UK); polyethylene glycol 4000 (PEG4000), polyethylene glycol 12000 (PEG12000), chitosan-medium molecular weight (CS), sodium chloride, potassium chloride, magnesium sulphate, calcium chloride, acetonitrile, orthophosphoric acid, acetic acid, ethanol, sodium hydroxide, potassium chloride and sodium chloride were obtained from Fisher Scientific (Loughborough, UK); gentamycin, (3-(4, 5-dimethylthiazol-2-yi)-2, 5-diphenyl tetrazolium bromide) MTT, Trypan blue dye, Pluronic® F127 (P127), sodium carboxymethyl cellulose (CMC), amantadine hydrochloride (AMT), sodium metabisulphate, mucin from porcine stomach, boric acid, dimethyl sulfoxide (DMSO) and benzalkonium chloride were obtained from Sigma-Aldrich (St. Louis, MO, USA).

Formulation development
Formulations were initially prepared by a modified ‘Cold method’ approach as originally described by Schmolka (1973). Briefly fixed quantities of excipients were utilised in all formulations and consisted of d-sorbitol (humectant) (0.5 % w/v), sodium metabisulphate (antioxidant) (0.1 % w/v) and benzalkonium chloride (surfactant) (0.1 % w/v). Additional polymers were incorporated and included mucoadhesives (CMC and CS) and permeation enhancers/ modulators of gelation temperature (PEG4000 and PEG 12000) at concentrations between 0.5-1.5 % w/v. These components were combined with ultrapure water under constant stirring. This was followed by AMT (0.5-1.5 % w/v) and the resulting solution was kept in an ice bath for 2 hours prior to the dropwise of P127 (15-20 %) and storage at 4 °C for 12 hours. Prior to use, the pH of all formulations were adjusted to 5.5. Formulations were developed according to variation in the concentration of P127, mucoadhesive polymer, and drug content.

Sol-gel transition \( (T_{\text{sol-gel}}) \)
A modified method described by Zaki et al. was used to determine the sol-gel transition temperature \( (T_{\text{sol-gel}}) \). A sample of each formulation (1 g) was transferred to a glass vial and heated in a dry block (Techne Dri-Block® DB-2D), initiated at 20 °C and increased by 1 °C after every 5 minutes of equilibration time. The \( T_{\text{sol-gel}} \) point was defined as the temperature whereby the upper meniscus of the gel did not move upon tilting the vial by 90 ° and this was used to demark optimal formulations. The optimal subset of formulations were then further characterised.

Rheological characterisation
To assess the rheological properties of the formulations, rheological measurements were carried out on AR-G2 Rheometer (TA Instruments, USA), using parallel plate geometry with 40 mm steel plates having a gap 1.0 mm. The approximate sample volume used was 1.26 mL and the instrument was used in the oscillatory mode in the linear viscoelastic range of the sample. Steady state behavior was assessed at ambient (18 °C) and nasal (34 °C) temperatures over a shear rate range from 0.1 to 100 s\(^{-1}\).

To quantify the rheological properties, the Ostwald-de Waele relationship (often called the power law) was applied to relate the shear stress \( (\sigma) \) is related to a consistency coefficient \( (k) \), the shear rate \( (\dot{\gamma}) \) and an index value \( (n) \):

\[
\sigma = k\dot{\gamma}^n
\]

which is applicable to the formulations in light of the constant shear viscosity curves. In the context of rheology, the index value can be used to determine the type behavioural flow, with Newtonian behaviour when \( n = 1 \). When the magnitude of \( n \) is < 1, the underlying rheological process is shear-thinning (pseudoplastic flow) and then when \( n > 1 \) the fluid is shear-thickening.
In-vitro mucoadhesion

The method of Nakamura et al. (1996) was adapted to measure the mucoadhesion of the formulation systems. Briefly, 100 g of a hot 1% agar and 2% mucin mixture in PBS (pH 6.0) was cast onto a glass plate and left to gel at 4 °C for 12 hours. The gel plate was then equilibrated at 34 °C for 1 hour followed by the application of a 250 µL sample of each formulation when the plate was kept at a 45° angle at 34 °C. This process was repeated using a 1% agar mixture (without mucin) and the mucoadhesion of the formulations were calculated by a ‘displacement’ value (measured in cm) reflecting the difference in gel movement in the presence and absence of 2% mucin.

AMT derivatisation and HPLC detection

To quantify the release of AMT from the formulations, a pre-column derivatisation method for memantine was modified in order to enable detection of AMT. Briefly, to 50 µL of sample or standard solution of AMT, 40 µL of 0.015 M FMOC and 50 µL of 0.5 M borate buffer were added and kept at room temperature for 20 minutes prior to the addition of 360 µL of a 0.05M borate buffer:acetonitrile (50:50 v/v) diluent. A Shimadzu HPLC system was used with a Phenomenex Luna C18 (150 x 4.6 mm) 5 µm column. The mobile phase consisted of phosphate buffer:acetonitrile (20:80 v/v). The column was maintained at 30 °C and 10 µL injected. The flow rate was maintained at 2 mL/minute with run time of 7 minutes. The UV detection of derivatised AMT was measured at 265 nm. The intra/inter-day precision was determined and the limit of detection (LOD) and limit of quantitation (LOQ) were calibrated according to ICH (QR (R1)) recommendations.

Stability assessment

Optimised formulations were stored in stability cabinets maintained at 4 ± 1 °C (Sanyo Medicool, UK) and 25 ± 2 °C (Firlabo, France) at a humidity of 60 % ± 5 %. The stability of the formulation was assessed based on the gelation of a sample of the formulation at 34 °C within 5 minutes, followed by assessment of the extent of drug release.

The release of AMT from the formulations were assessed by a ‘membrane-less’ diffusion system. A mass of formulation (0.25 g) appropriate for the nasal cavity volume was transferred to a glass vial and allowed to equilibrate in a dry block at 34 °C. Artificial nasal electrolyte (ANE) (0.8 g/L NaCl, 3 g/L KCl, and 0.45 g/L CaCl2, pH 6.8) was used as release media and 100 µL transferred onto the gelled formulation. This volume was completely withdrawn and replaced with fresh ANE at each time point. Samples were analysed by a pre-column derivatisation method followed by HPLC analysis.

In-vitro drug release kinetics

Several kinetic drug release mathematical models were used to assess drug release from the formulations. The best-fit to the mathematical models described below confirmed the appropriate release kinetics:

Higuchi model: \[ \frac{M_t}{M_\infty} = k_H \cdot t^{\frac{1}{2}} \]
where \( M_t/M_\infty \) is the drug fraction released at time \( t \) and \( k_H \) is the Higuchi constant.

Zero order model: \[ \frac{M_t}{M_\infty} = k_0 \cdot t \]
where \( M_t/M_\infty \) is the drug fraction released at time \( t \) and \( k_0 \) is the zero-order release constant.

First order model: \[ \frac{M_t}{M_\infty} = 1 - e^{-k_1 \cdot t} \]
where \( M_t/M_\infty \) is the drug fraction released at time \( t \) and \( k_1 \) is the first-order release constant.

Power law: \[ \frac{M_t}{M_\infty} = kKP \cdot t^n \]
where \( M_t/M_\infty \) is the drug fraction released at time \( t \), \( kKP \) is a kinetic constant which describes the structural and geometrical elements of the formulations and \( n \) is the release exponents which is used to indicate the mechanism of drug release. The power law, otherwise known as the Korsmeyer-Peppas power law, has been used in many pharmaceutical formulations. The value of \( n \) is important in understanding the mechanism of release when it is unknown and often polymeric formulations can be categorised accordingly to this scale. When \( n \leq 0.45 \) drug release is diffusion controlled and sometimes referred to as Fickian diffusion and when \( n > 0.89 \) the diffusion is indicative of erosion controlled drug release or class-II kinetics. For situations where \( 0.45 < n \leq 0.89 \)
the diffusion is a complex mixture of both processes and often termed anomalous transport. In all cases this is based on the assumption of release from a cylinder and applied to cumulative release rates < 60 % \(^{45}\).

**Human nasal epithelial cell culture model**

The immortalised human nasal epithelial cell line RPMI 2650, was used to assess the compatibility of the formulations with human nasal epithelia and to develop an in-vitro nasal epithelial cell culture model \(^{50}\) to assess drug release/transport. Cells were grown in MEM supplemented with 10 % FBS, 1 % L-glutamine, 1 % NEAA, 1% penicillin-G/streptomycin in a humidified 37 °C incubator with 5 % (v/v) CO\(_2\) \(^{51}\). The media was changed every 2 days.

**(3-(4,5-Dimethylthiazol-2-yl)-2,5-Diphenyltetrazolium Bromide) (MTT) assay**

To assess cellular toxicity, RPMI 2650 cells were seeded at a density of 4x10\(^4\) cells per well of a 96-well plate and allowed to attach for 24 hours. The media was removed and replaced with 200 µL of media containing either AMT (0.01-50,000 µM), mucoadhesive polymers or P127 (10-4000 µg/mL) and incubated for 24 hours at 37 °C in a 5 % (v/v) CO\(_2\) humidified air environment. Subsequently 20 µL of 5 mg/mL MTT dissolved in PBS was added to each well and incubated at 37 °C in a humidified air environment for 4 hours. After incubation the medium was removed and 100 µL of DMSO was added and the plates left to incubate for 15 minutes in the dark. The UV-absorbance of the formazan product was determined at 595 nm. Each concentration was assayed in eight wells and run in three independent experiments and results expressed as percentage cytotoxicity relative to a control (0.5% DMSO).

**Human nasal epithelial airway cell culture model: AMT transport**

To assess the release and transport of AMT from formulations, an in-vitro nasal epithelial airway cell culture model was developed with RPMI 2650 cells seeded onto collagen-coated 6-well permeable inserts (ThinCerts\textsuperscript{TM}) at a seeding density of 4x10\(^5\) cells/cm\(^2\) \(^{52}\)\(^{50}\) and grown for 14 days. On days 10-14 an air-liquid interface (ALI) was initiated with the removal of apical media. The integrity of the monolayer was determined by measuring the trans-epithelial electrical resistance (TEER) using an EVOM epithelial voltohmmeter (World Precision Instruments Inc.). Inserts were deemed acceptable where the TEER value was above 180 Ω.cm\(^2\) \(^{52}\)\(^{50}\).

Immediately prior to the start of the transport study, maintenance media was replaced with transport media (HBSS with 25 mM HEPES) and the cells left to acclimatise for 30 minutes in a humidified 37 °C incubator with 5% (v/v) CO\(_2\). A 1 mL sample of formulation was placed into the apical chamber, after removal of transport media from apical and basolateral compartment, and returned to the incubator for 5 minutes. Once gelled, fresh transport media was placed into the basolateral chamber and 250 µL samples withdrawn at 5, 15, 30, 60, 120, 150 and 180 minutes and replaced with fresh pre-warmed media. Withdrawn samples were derivatised and AMT release quantified by HPLC analysis.

**Nasal spray systems: droplet size distribution**

To assess the potential in-vivo deposition characteristics of the proposed nasal formulation, a multi-dose pump spray delivery system was employed to characterise the formulation droplet size distribution using a laser diffraction technique. The HELOS (Sympatec, UK) system was used with an R3 lens (measuring range 0.5-175 µm). The nasal pump was vertically mounted 3 cm away from the laser path and a vacuum source was mounted anterior to the pump system. The pump systems were pre-actuated prior to mounting, and actuated three times to detect the particle size distribution. Data was reported as volume diameters at 10 %, 50 % and 90 % of the cumulative undersized volumes. Span was calculated as: \([Dv90-Dv10]/Dv50\).

**Nasal spray system: human nasal cast deposition**

To assess the potential in-vivo deposition characteristics of the proposed nasal formulation, a multi-dose pump spray delivery system was employed to assess the deposition of formulations into an anatomically correct human nasal cast model (Koken Ltd, Japan). The inner surface of each nostril was evenly coated in a water-indicated dye (Kolor Kut, USA). Formulations were loaded into the spray pump, pre-actuated, before one actuation was
delivered into the nasal cast where the cast was fixed in an upright head position. The impact of spray angle and nostril insertion depth on nasal deposition was analysed through photographing colour changes. The captured images were subsequently processed through pixel quantification software to quantify the deposition patterns and calculate the deposition surface area. The depositions of within the nasal cast was classified according to the predominate regions of deposition within the cast, i.e. lower nasal regions (nasal vestibule), the middle/upper nasal regions (turbinates) and the olfactory regions were assessed.

**Statistical analysis**

Unless otherwise stated, three independent experiments were carried out for each study. Statistical significance was evaluated by one-way ANOVA or paired two-tail Students t-test using GraphPad Prism version 6.00 for Windows (GraphPad Software, La Jolla California USA, www.graphpad.com). Calculations of IC50 were determined using a four-parameter logistic sigmoidal fitting function within GraphPad Prism. Unless otherwise states, data is reported as mean ± standard deviation (SD). A significance level (p-value) of < 0.05 was considered as statistically significant.
RESULTS AND DISCUSSION
The treatment of age-associated degenerative CNS disorders such as Parkinson’s diseases, poses particular difficulties in the design of appropriate drug delivery systems capable of overcoming the hindrances associated with dyskinesia and dystonia that are inherent in current pharmacological interventions for PD. Although reformulation of existing orally dosed formulations may provide some benefit there is an inherent need to provide an alternative delivery routes of a range of CNS disorders which can both provide ease of clinical use but also improve the bioavailability of the therapeutic.

The nasal mucosa fulfils this purpose in light of the large and vascularised route of access for drugs to the systemic circulation and can provide a relatively non-invasive approach to delivery drugs.

A key concern of any nasal administered formulation is the residency of the formulation within the nasal cavity. Existing formulations are often associated with a nasal drip or ‘run-off’ effect which potentially diminish the bioavailability of nasal formation. A novel development in this area has been the formulation of intelligent response based hydrogel nasal formulations which act to enhance residency within the nasal cavity.

Sol-gel transition ($T_{\text{sol-gel}}$)
The initial screening and optimisation resulting in 32 formulations, of which 3 demonstrated optimal gelation at the nasal cavity temperature of 34 °C ± 1°C. These formulations were identified as FCMC (1 % w/v CMC), FPEG (1 % w/v PEG4000) and FCS (0.1 % w/v chitosan) and which all contained 17 % w/v P172 and 1% w/v amantadine (in addition to excipients).

Rheological behaviour
When assessing the properties of thermoresponsive gel-like systems, the storage/elasticity modulus (G’) and the loss modulus (G") are key metrics to describing the rheology of the formulation. The elasticity modulus is an important metric in this regard and measures the energy stored and which is subsequently recovered for each cycle of deformation. It presents with lower values with reduced temperatures but significantly increases with an increase in temperature, and the point at which G’ overtakes G” indicates the initiation of the gelation phenomena. To assess this phase transition the ratio of G’” and G’ (the loss tangent, Tan δ) was used, and where a phase-transition occurs an abrupt change in Tan δ is observed with values of < 1 indicating a greater G’ compared to G”", indicative of gel formation. This phase transition is often associated with three distinct phases: (i) an initial stable liquid phase plateau at low temperatures; (ii) an abrupt transition phase during the gelation process; (iii) a late stage stable gel plateau.

All formulation demonstrated a gelation phenomenon, which initiated at approximately 26-28 °C, and was preceded by the stable plateau region (Figure 1). FCS and FCMC (Figure 1A) demonstrated a complete profile with three distinct phases culminating in a stable gel formulation, whereas FPEG (Figure 1A) only demonstrated two phases, plateau followed by a partial gelation phase, but did not form a stable a gel at the termination of the study (34 °C).

The rate of gelation is also a key determinant of residency within the nasal cavity and this was assessed through a time-sweep analysis at a fixed temperature (34 °C). FCMC demonstrated the quickest gelation of 28 ± 2 seconds which was followed by FCS (37 ± 3 seconds) and FPEG (68.5 seconds) (Figure 1B).

At nasal cavity temperatures, a significant shear thinning behaviour was observed in all formulations, indicative of temperature-induced gel formation when compared to ambient temperatures (Figure 2), with viscosities for 34 °C being statistically significantly different to those at 18 °C ($P < 0.0001$). Furthermore, at 34 °C no significant difference in viscosity was reported over the shear rate studied ($\rho = 0.227$), however FPEG demonstrated a statistically significant lower viscosity compared to FCS and FCMC ($P < 0.01$).
As the shear rate increased from 0.091 to 100 s$^{-1}$, the viscosity of the formulations dropped by 990-fold (FCMC), 409-fold (FCS) and 207-fold (FPEG) (Figure 2). However, all formulations exhibited non-Newtonian behaviour at nasal temperatures compared to Newtonian behaviour at ambient temperatures (Table 1).

From an interpretation of the Ostwald model, the gel structure can be changed as a result of any deformation induced changes in the gel particles, through changes in the alignment of the polymer chains and also any changes in the interactions between polymer chain segments and any side chains. Therefore, upon heating the values of $n$ will be lower in for stronger gels due to an increase in the noncovalent forces of between neighbouring particles $^{56}$.

It was also apparent that the presence of different mucoadhesives can significantly alter the behaviour of the formulations. For examples, the viscosity of FCMC was always higher than all other formulations, and particularly at 34 °C where there is an approximate 8-fold difference between FCMC and FPEG. Furthermore, the FCMC demonstrated a much lower $n$-value compared to FCS and FPEG (Table 1), suggesting a stronger gel formation at 34 °C. This phase represents the gradual conversion into a highly viscous/solid state. During this phase the intricate hydrogen-binding network between the unimers of F127 and water molecules are disrupted and drives the formation of micelles, which further aggregate to form the physical gel-like structure. $^{57}$ On the basis of $^{13}$C NMR studies Pisal $^{58}$ suggested that at high temperature, conformational changes in the methyl group of the polyoxypropylene within the hydrophilic micellar region and in the motion of the hydrophilic end chains takes place. This results in dehydration and end chain friction, which causes the gelation.

**In-vitro mucoadhesion**

Mucoadhesion is defined as a state where at least one biological material is held with another material for a prolonged period of time $^{59}$. In the context of drug delivery, this can be defined as the adhesion of an artificial material (such as a polymer system) to a biological substrate material $^{60}$. We assessed the mucoadhesion of optimized formulations using an inclined-plate method in the presence and absence of mucin $^{24}$, with mucoadhesion assumed to have occurred if the displacement of the formulation in the absence of mucin is greater than in the presence of mucin.

The displacement transfer test demonstrated that all formulations showed a statistically significant ($P \leq 0.001$) displacement in the absence of mucin (Figure 3). The difference between the absence and presence of mucin is indicative of the displacement effect and FCMC demonstrated the least extent of displacement (2.26 cm) with FPEG demonstrating the greatest extent of displacement (3.8 cm).

The superiority of FCMC and FCS, compared to FPEG, is presumed to be a function of the enhanced rheological properties FCMC and FCS possess (Figure 1 and 2). The fact that displacement is observed in the absence of mucin would indicated that the use of is able to recover some of the mucoadhesive properties of the mucoadhesive polymers included within the formations, and although known to show less biophysical properties as compared to native mucin, this purified mucin has less batch-to-batch variation and hence provides better reproducibility in measuring mucoadhesive force $^{61}$.

**AMT derivatisation and HPLC detection**

The HPLC was successfully able to detect the derivatisation of AMT with a retention time of 4.4 minutes. Percentage RSD for system precision (0.120 %) and method precision (0.121 %) were within the acceptable limits of 1 % and 2 % respectively.

The linearity response of amantadine was determined using concentrations ranging from 0.0078 to 1.0 mg/mL and the response was linear over the specified range with a correlation coefficient of 0.9998, a slope of 1385904 ± 33080 and an intercept of -13515 ± 1045. The validity was verified by ANOVA with no deviation from linearity.
Stability assessment

To assess the stability of the optimised formulations, batches were prepared and stored in stability cabinets at 25 °C/60 % RH and 4 °C for 8 weeks, with sampling at weekly intervals followed by assessment of AMT release conducted at nasal temperatures (34 °C). A prerequisite for conducting release studies was the successful gelation of the stability samples prior to initiation of release studies.

All formulations stored at 4 °C demonstrated successful sol-gel transition prior to the initiation of the release studies, confirming the thermoresponsive nature of the gel systems was maintained during the stability study duration. However only FCS demonstrated successful sol-gel transition when stored at 25 °C/60 % RH for 2 weeks, confirming that storage of formulations would be more appropriate at ‘refrigerated’ conditions to ensure stability of the products.

The cumulative percentage release at preparation and after 1 week storage at 4 °C were not significantly different for all formulations, reaching 15-17 % release (Figure 4A). Following 8 weeks storage at 4 °C, the quantity of AMT released from FCMS (16.15 % ± 0.82 %) was not significantly different to that at week 0 (16.86 % ± 1.13 %) (Figure 4B), however at earlier time-points during the release study (30-120 minutes) significant differences (p < 0.05) in the AMT quantity released were apparent (Figure 4B). For FCS, the amount of AMT released following 8 weeks storage at 4 °C (18 % ± 0.99 %) was significantly (p < 0.05) greater than that at week 0 (15.32 % ± 0.30 %) (Figure 4B). Significant differences in the release profile were also apparent during the initial phases of the release study (30-60 minutes) (Figure 4B).

As the formulations presented are inherently based on ‘swellable’ type polymer systems, the diffusion was also modelled by the Korsmeyer-Peppas power law and generally demonstrated Fickian-type diffusion early on (except for FCS). By week 8 this had moved towards anomalous transport (Table 2), whereby drug diffusion and the polymer relaxation were contributing to the overall release kinetics.

The movement towards anomalous transport can be explained by considering the relationships between the two driving forces influences the diffusional process within the polymer system, namely the rate of solvent diffusion (R0) and the rate at which polymer chains relax (Ra).

In Fickian-type diffusion (case-I), R0 is much slower compared Ra. For non-Fickian diffusion (case II, anomolous and super case II) the main differences lies with the R0 whereby for case II Rk >> R0, for anomalous-type kinetics R0 = Ra and for super case II R0 >> Ra. Initially, formulations demonstrated Fickian-type diffusion (Table 2) but the Korsmeyer-Peppas exponent increased over the stability period to > 0.5 by week 8 moving towards anomalous transport (Table 2). This would suggest that polymer relaxation (swelling/erosion) plays a critical role in the release of AMT at earlier stability time points. The mathematical description and modelling of the casual factors for anomalous transport is challenging, and is beyond the scope of this study however a detailed review has been published discussing this issue.

The mass transport of AMT into the polymer network can be assumed to be comprised of 3 processes, whereby the solvent (water) is absorbed onto the surface of the polymer. The solvent molecules may then enter the lattice network of the polymer, which causes a ‘swelling’ of the network and elongation. This elongation is then counter balanced by an elastic-restrictive force to oppose the swelling. Eventually a state of equilibrium is reached when both forces are balanced, these process can significantly alter the kinetics of drug diffusion through the polymer network and are highly influenced by time-scale associated structural changes in the polymer. The move towards anomalous transport over 8 weeks may suggest a polymer network time-associated affect, which becomes more apparent over the longer stability study. The equilibrium between swelling and elastic-restrictive force within the polymer may be associated with a time-dependent equilibrium, where equilibrium is not established immediately.
Although the formulations are in liquid-phase during the stability storage period, the interactions between both mucoadhesive polymers and thermoresponsive polymers may influence eventual formulations of the polymer networks, altering the equilibrium processes and hence contributing towards the anomalous transport.

**Human nasal epithelial cell culture model: MTT assay**

To investigate the toxicity of amantadine and polymers used within the formulations, a cellular viability study was conducted using the human nasal epithelial cell line RPMI 2650. Cell viability was generally maintained for AMT up to 1 mM, with an IC50 of 4.6 mM ± 0.05 mM (Figure 5A) when exposed for 24 hours. For all polymers tested, no significant decrease in cellular viability was observed up to 4 mg/mL (Figure 5B-D) and confirmed the formulations developed would be tolerable, from a cellular toxicity perspective, to human nasal epithelia. Furthermore, no previous reports have examined the in vitro toxicity profile of AMT towards human nasal epithelia and our reports demonstrated that 1 mM would be suitable for prolonged (up to 24 hours) exposure.

**Human nasal epithelial airway cell culture model: AMT transport**

The in vitro release of AMT alone and from formulations was further assessed in an in vitro nasal epithelial airway cell culture model over 3 hours (Figure 6). In all formulations a statistically significant (p ≤ 0.01) release of 43-44 % was observed over the duration of the transport study in comparison to that of AMT (79 % ± 3.58 %) (Figure 6). The mechanism of release, as described by the Korsmeyer-Peppas power law, was modelled according to a super case-II effect (n > 0.89) where the sorption of solvent results in breaking of the polymer network (termed solvent crazing) as a result of the swelling of the gel within the vitreous nucleus and can be a result of crosslinking density, drug loading and copolymer composition. The super-case II diffusion of AMT from all gel systems was surprising, and therefore the release mechanism of AMT from the formulations is not clear as it is beyond the limits of the power law. It suggests that a rapid release of AMT occurred later in the experiment resulting in rapid relaxation governing/controlled transport of AMT from the polymer network. The ionic composition of the media, comprising of a pH of 7.4, may have directly altered the state of ionisation within the polymer gel network, compared to the pH of the ANE (pH = 5.5) leading to an altered release profile, with the rapid release being a time-dependant reflection of the shift in equilibrium as the solvent (media) penetrates the polymer network.

**Nasal spray systems: droplet size distribution**

Prevention of pulmonary deposition of nasally administered spray formulations is paramount to ensure residency within the nasal cavity. Droplet size distribution is key in determining the potential for pulmonary deposition (< 2 µm), however the droplet size is often dictated by the design of the orifice of the actuator device used to deliver the spray plume. For ensuring retention of drug within the nasal passages, droplet sizes of in-excess of 5 µm are recommended and represented our target cut-off. Our formulations all demonstrated a diameter in excess of the traditional cut-off of 5 µm. (Table 3). FCS demonstrated a 10% fraction of 7.27 µm ± 0.28 µm (Table 3) closest to the cut-off for pulmonary deposition, whereas both FCMC and FPEG were above this. FCS also demonstrated the smallest diameter (VMD = 92.41 µm ± 1.72 µm). However, when considering the span of particle sizes, FCS was the broadest (0.96 ± 0.30) with both FCMC and FPEG showing a smaller distribution spread (Table 3). A one-way ANOVA confirmed droplet VMD were significantly different across all formulations (p < 0.0001), suggesting that the incorporation of different mucohesive polymers contributed to the differing polymer sizes with FCS demonstrating the small resulting droplets (92.41 µm ± 1.72 µm) and FCMC the largest (120.87 µm ± 0.59 µm). When considering the viscosity of the polymers used in each formulation (Figure 1), CMC resulted in the most viscous formulations and this may have contributed to the larger particle sizes, with similar trends observed for CS and PEG.

**Nasal spray system: human nasal cast deposition**

To assess the potential in vivo application of formulations, FCS and FCMC were taken forward to assess spray deposition within a human nasal cast model. The angle of spray administration was altered from 60-80° and the
impact on deposition patterns assessed. For both formulations the administration angle had a significant effect on deposition, with an increased angle (80°) having a significantly pronounced localisation in the nasal vestibules and lower regions of the nasal cavity. For FCMC (Figure 7) this deposition area was significantly smaller greater than that for FCS (Figure 8).

Interestingly, spray angles of 60° and 70° lead to deposition of the formulation in the olfactory regions of the nasal cavity, suggesting the possibility for exploitation in nose-to-brain delivery of amantadine. FCS sprayed at an angle of 60° with insertion lead to olfactory deposition of 0.6-0.7 cm². For FCMC a greater olfactory deposition of 0.91 cm² was observed at 60° with insertion. Although the higher deposition for FCS may be a result of the smaller particle sizes for FCS (92.41 ± 1.72 μm) compared to 120.87 ± 0.59 μm, the impact of nozzle insertion for FCMC may enhance delivery onto the olfactory regions (Figure 7 and 8). When considering the surface area of the human olfactory mucosa (2-10 cm²) FCMC may be a viable candidate for further development targeting nose-to-brain drug delivery.

In both formulations, the impact of head position (15° forward) was important in governing nasal depositions, leading to significantly higher deposition within the middle-upper regions of the cast irrespective of the insertion angle.

The higher viscosity of FCMC compared to FCS resulted in significantly larger particle sizes ($p < 0.001$) for the optimal formulations and may have resulted in a more focussed (less disperse) spray plume resulting in more localised deposition patterns. In contrast, the lower viscosity of FCS, leading to small particles would lead to a wider plume and greater deposition in the nasal cavity.

As a viable vehicle for nasal or olfactory delivery and residency of formulations within the nasal cavity, the optimised formulations demonstrated rapid gelation (FCS: 36.6 s; FCMC: 28 s) (Figure 1) with minimal displacement within the nasal cavity (FCS 0.035 cm/s; FCMC: 0.038 cm/s) (Figure 3). When considering their potential residency within the nasal cavity, the sol-gel conversion and displacement would result in minimal nasal cavity displacement (FCS: 1.28 cm; FCMC: 1.06 cm). In-vitro release studies were conducted for 3-hours, with the gel structure intact following the release studies, hence the residence in-vivo would be expected to be at least 3-hours. Furthermore, the identification of potential nose-to-brain delivery, through olfactory deposition, would suggest that these formulations may be adapted to better exploit this ‘direct’ transfer route to the brain.
CONCLUSION

This study has investigated the potential to administer the anti-Parkinsonian drug amantadine through the nasal route using a mucoadhesive and thermoresponsive hydrogel formulation systems. We have demonstrated that both carboxymethyl cellulose and chitosan can provide the requirements for mucoadhesion in these formulations, whilst also leading to a relatively stable formulation which can be targeted to delivery drug in specific regions of the nasal cavity dependant on administration angle and nozzle insertion depth. Furthermore, our results have indicated the potential for nose-to-brain delivery of amantadine, yielding a potentially novel avenue therapeutics delivery route to avoid the blood-brain barrier.
REFERENCES


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Figure 4: Amantadine cumulative % release for optimised formulations following storage at 4°C and 25°C for up to 8 weeks. (A) Cumulative % AMT release for samples successfully gelling at nasal cavity temperature (34 °C); (B) Comparison of AMT release from FCMC and FCS at preparation (Week 0) and at the end of the stability period (Week 8). * p < 0.05, ** p < 0.01.

Figure 5: Cellular toxicity of amantadine, CMC, PEG4000 and F127 on RPMI 2650 cells. Cells were grown on a 96-well plate and exposed to various concentrations of amantadine (0.1 nM – 10 mM) or polymer (10-4,000 µg/mL) for 24 hours. Data is reported as mean ± SD with up to 8 replicates per compound in at least 3 independent experiments.

Figure 6: Transport of AMT across an in vitro cell culture model of the nasal epithelial, from optimised formulations in HBSS-HEPES transport media. ** p < 0.01.

Figure 7: Representative nasal deposition patterns of FCMC. Brown colours represent the applied Kolor-Kut and green colours resent the depositions patterns of the spray. Angles refer to nasal spray position relative to the horizontal place (60, 70 or 80°), insertion of the spray orifice into the nostril (5 mm) or with the nasal cast angled 15° forward.

Figure 8: Representative nasal deposition patterns of FCS. Brown colours represent the applied Kolor-Kut and green colours resent the depositions patterns of the spray. Angles refer to nasal spray position relative to the horizontal place (60, 70 or 80°), insertion of the spray orifice into the nostril (5 mm) or with the nasal cast angled 15° forward.
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![Graph showing drip transfer](image-url)
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![Graph showing cumulative transport of AMT over time](image)
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<table>
<thead>
<tr>
<th>Formulation</th>
<th>Temperature (°C)</th>
<th>k</th>
<th>n</th>
<th>Behaviour</th>
</tr>
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<tbody>
<tr>
<td>FCMC</td>
<td>18</td>
<td>0.157</td>
<td>0.972</td>
<td>Newtonian</td>
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<tr>
<td>FPEG</td>
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<td>0.0545</td>
<td>0.971</td>
<td>Newtonian</td>
</tr>
<tr>
<td>FCS</td>
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<td>1</td>
<td>Newtonian</td>
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<td>FCMC</td>
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<tr>
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<td>13.59</td>
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<tr>
<td>FCS</td>
<td>34</td>
<td>56.56</td>
<td>0.1762</td>
<td>Non-Newtonian</td>
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</tbody>
</table>
**Table 2:** Drug release kinetics analysis of formulations stored at ambient (18 °C) and nasal cavity (34 °C) temperature for up to 8 weeks.

<table>
<thead>
<tr>
<th>Week</th>
<th>Formulation</th>
<th>Korsmeyer-Peppas</th>
<th>4 °C</th>
<th>25 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>FCMC</td>
<td>$kKP = 2.859 \pm 0.899$; $n = 0.391 \pm 0.051$</td>
<td>FPEG</td>
<td>$kKP = 2.931 \pm 1.160$; $n = 0.352 \pm 0.076$</td>
</tr>
<tr>
<td>0</td>
<td>FPEG</td>
<td>$kKP = 2.004 \pm 0.027$; $n = 0.438 \pm 0.24$</td>
<td>FPEG</td>
<td>$kKP = 2.12 \pm 0.032$; $n = 0.423 \pm 0.013$</td>
</tr>
<tr>
<td>8</td>
<td>FCMC</td>
<td>$kKP = 1.034 \pm 0.170$; $n = 0.538 \pm 0.024$</td>
<td>FCMC</td>
<td>$kKP = 1.614 \pm 0.351$; $n = 0.474 \pm 0.042$</td>
</tr>
<tr>
<td>1</td>
<td>FCS</td>
<td>$kKP = 1.997 \pm 0.659$; $n = 0.433 \pm 0.059$</td>
<td>FCS</td>
<td>$kKP = 1.614 \pm 0.351$; $n = 0.474 \pm 0.042$</td>
</tr>
</tbody>
</table>

$kKP$: Korsmeyer-Peppas constant; $n$: release exponent. Mean ± SD.
Table 3. Laser diffraction particle size analysis

<table>
<thead>
<tr>
<th></th>
<th>Diameter (µm)</th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Dv10</td>
<td>Dv50</td>
<td>Dv90</td>
<td>VMD</td>
<td>Span</td>
</tr>
<tr>
<td>FCS</td>
<td>7.27 ± 0.28</td>
<td>109.07 ± 3.42</td>
<td>157.77 ± 0.99</td>
<td>92.41 ± 1.72</td>
<td>1.38 ± 0.21</td>
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<tr>
<td>FCMC</td>
<td>39.81 ± 1.02</td>
<td>129.86 ± 0.29</td>
<td>167.39 ± 0.09</td>
<td>120.87 ± 0.59</td>
<td>0.98 ± 0.30</td>
</tr>
<tr>
<td>FPEG</td>
<td>43.93 ± 0.88</td>
<td>104.67 ± 1.88</td>
<td>150.95 ± 1.04</td>
<td>100.35 ± 1.53</td>
<td>1.02 ± 0.06</td>
</tr>
</tbody>
</table>

Mean ±SD, 6 replicate spray actuations per formulation.

Dv10: 10% of the cumulative undersized (volume) fraction; Dv50: 50% of the cumulative undersized (volume) fraction; Dv90: 90% of the cumulative undersized (volume) fraction; VMD: volume mean diameter; Span: relative span (Dv90-Dv10)/Dv50.