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Formulation, characterisation and stability assessment of a food derived 1 tripeptide, Leucine-Lysine-Proline loaded chitosan nanoparticles

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1 **Formulation, characterisation and stability assessment of a food derived**
2 **tripeptide, Leucine-Lysine-Proline loaded chitosan nanoparticles**

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24 **Abstract**

25 The chicken or fish derived tripeptide, Leucine-Lysine-Proline (LKP), inhibits the
26 Angiotensin Converting Enzyme and may be used as an alternative treatment for pre-
27 hypertension. However, it has low permeation across the small intestine. The formulation of
28 LKP into a nanoparticle (NP) has the potential to address this issue. LKP-loaded NPs were
29 produced using an ionotropic gelation technique, using chitosan (CL113). Following
30 optimisation of unloaded NPs, a mixture amount design was constructed using variable
31 concentration of CL113 and tripolyphosphate at a fixed LKP concentration. Resultant
32 particle sizes ranged from 120-271 nm, zeta potential values from 29-37 mV and
33 polydispersity values from 0.3-0.6. A ratio of 6:1 (CL113: TPP) produced the best
34 encapsulation of approximately 65%. Accelerated studies of the loaded nanoparticles
35 indicated stability under normal storage conditions (room temperature). Cytotoxicity
36 assessment showed no significant loss of cell viability and *in vitro* release studies indicated
37 an initial burst followed by a slower and sustained release.

38

39 **Keywords:** chitosan nanoparticles; food derived peptide; mixture amount design; accelerated
40 thermal stability analysis; ACE inhibition

41 **1. Introduction**

42 A number of synthetic antihypertensive drugs (ACE inhibitors) are currently available on the
43 market (e.g. captopril, lasinopril and enalapril) but all have been reported to have associated
44 adverse side effects, such as coughing, dizziness, loss of taste and skin rashes and poor
45 pharmacokinetics with a short half life, resulting in the requirement of frequent dosage
46 (Bougatef et al., 2008). Therefore, natural ACE inhibitors, isolated from food sources, have
47 attracted increasing attention in recent years, in the search to find a safer and more
48 economical approach for the consumers (Bougatef et al., 2008). Peptides derived from food
49 sources have been reported to have health benefits such as hypotensive activity, due to their
50 Angiotensin Converting Enzyme (ACE) inhibitory activity (Berilyn et al., 2016; Li et al.,
51 2016). Using DNA microarray experiments it was found that other mechanisms can also
52 contribute to the decrease of blood pressure for bioactive tripeptides (Yamaguchi,
53 Kawaguchi, & Yamamoto, 2009) reducing the side effects of ACE inhibition. However,
54 exploitation of the potential nutraceutical benefits of these peptides in general is known to
55 face a number of challenges. Insufficient gastric residence time, low permeation and/or
56 solubility within the gut, chemical degradation within the gastrointestinal tract (GIT) due to
57 low pH, enzymatic degradation, and the presence of other nutrients (in food), all limit the
58 bioavailability of bioactive peptides by the oral delivery route (Braithwaite et al., 2014; Ma,
59 2014; Segura-Campos, Chel-Guerrero, Betancur-Ancona, & Hernandez-Escalante, 2011). A
60 number of researchers have attempted to formulate peptides into oral delivery systems for
61 example by the addition of absorption enhancers (Choonara et al., 2014), enzyme inhibitors
62 (Bruno, Miller, & Lim, 2013), hydrogels (Sharpe, Daily, Horava, & Peppas, 2014), liposomes
63 (Takahashi, Uechi, Takara, Asikin, & Wada, 2009) and nanoparticles (Yao, McClements, &
64 Xiao, 2015).

65 Leucine-Lysine-Proline is a tripeptide, derived from chicken muscle, which has shown *in*
66 *vitro* ACE inhibitory activity, having a mean inhibitory concentration (IC₅₀) of 0.32μM
67 (Zhou, Du, Ji, & Feng, 2012). In addition, LKP has been shown to elicit a significant
68 reduction of blood pressure in spontaneously hypertensive rats (SHR) when delivered
69 intravenously, 10mg/kg⁻¹ producing a reduction in systolic blood pressure of 75mmHg,
70 compared to an oral dose of 60mg/kg⁻¹, which resulted in a reduction of 18mmHg (Fujita,
71 Yokoyama, Yoshikawa, Iroyukifujita, & Eiichiyokoyama, 2000). Captopril, a synthetic oral
72 ACE inhibitor drug used for the treatment of hypertension, has been reported by Quiñones *et*
73 *al.* (2015) to show a maximum *in vivo* change in SHR of 60.5 ± 2.7 mmHg, 4 hours post-
74 administration, when 50mg/kg was given orally . Recent studies have shown that LKP is
75 stable in the GIT but with lower permeation than the market drugs, across the intestine at its
76 target site, the small intestine (Gleeson, Heade, Ryan, & Brayden, 2015). Formulation into
77 nanoparticles (NPs) for oral delivery can enhance the bioavailability of an encapsulated
78 peptide drug and consequently improve its pharmacokinetics and stability (Patel, Patel, Yang,
79 & Mitra, 2014; Ryan et al., 2013). Stable NPs with particle sizes ranging between 100-500nm
80 (des Rieux, Fievez, Garinot, Schneider, & Pr at, 2006), zeta potential values (ZP) ≥ 30mV
81 (Lakshmi & Kumar, 2010), polydispersity (PDI) < 0.400 (Abdel-Hafez, Hathout, &
82 Sammour, 2014a) and maximum encapsulation efficiency are ideal characteristics for oral
83 supplementation.

84 Chitosan is a linear polysaccharide, prepared by N-deacetylation of chitin (Rinaudo, 2006).
85 Chitosan NPs have shown promising results for oral delivery/supplementation due their
86 GRAS properties and their intrinsic properties, including, non-immunogenic, mucoadhesion
87 and the ability to transiently open the tight junctions of the intestinal barrier, which can help
88 facilitate transport of macromolecules and has the potential to act as an enhancer (Chuah,
89 Kuroiwa, Ichikawa, Kobayashi, & Nakajima, 2009; de Moura et al., 2009; Madureira,

90 Pereira, & Pintado, 2016). LKP NPs were produced using an ionotropic gelation technique.
91 This technique allows the preparation of chitosan NPs in aqueous solution and avoids the use
92 of organic solvents, high dispersion energy and harsh conditions, making the technique
93 suitable for the inclusion of nutraceuticals (García, Forbe, & Gonzalez, 2010). In this process,
94 a chitosan with a high degree of deacetylation is used, which increases the viscosity and
95 results in an extended conformation with a more flexible chain because of the charge
96 repulsion in the molecule (Franca, Freitas, & Lins, 2011). Chitosan can be ionically cross-
97 linked by counterions, such as sodium tripolyphosphate (TPP), to form a hydrogel of
98 microparticles, and when the relative concentrations of chitosan and these anions are
99 appropriate, NPs may be generated (Sureshkumar, Das, Mallia, & Gupta, 2010).

100 The formulation of NPs from different constituent components can be troublesome, due to the
101 different variable parameters used (concentration, temperature and pH). Empirical
102 optimisation using Response Surface Modelling (RSM) can help to rationalise the process
103 and has found applications in different fields, such as engineering, pharmaceutical,
104 biomedical, environmental and epidemiological research (Singh, Singh, Saraf, & Saraf,
105 2011). RSM has been shown to be useful for optimisation of experimental parameters in
106 nanoparticle formulation, and has been adopted by a number of research groups (Abdel-
107 Hafez, Hathout, & Sasmour, 2014b; Bezerra, Santelli, Oliveira, Villar, & Escalera, 2008).

108 The aim of this work is to formulate and investigate the feasibility of LKP encapsulated
109 chitosan NPs, determining the physico-chemical characteristics, stability to storage,
110 bioactivity and low cytotoxicity properties. The formulation of the LKP NP was optimised
111 using a RSM approach. The physico-chemical characteristics of the NPs were assessed using
112 dynamic laser scattering, scanning electron microscopy, and Fourier transform infrared
113 spectroscopy, with the aim of producing the optimal NPs as an oral delivery system.
114 Accelerated thermal conditions are employed to explore the stability for future storage

115 conditions; particle size, polydispersity, zeta potential and bioactivity were assessed. In
116 addition, the cytotoxicity and release profiles in simulated gastric and intestinal fluids were
117 assessed.

118 **2. Materials and Methods**

119 LKP (Mw 356.47, purity = 96% according to the manufacturer's specifications) was
120 synthesised by ChinaPeptides Co. Ltd, (Shanghai, China). CL113 (Mw = 110 kDa,
121 deacetylation degree (DD) = 86% according to manufacturer's specifications) was obtained
122 from Pronova Biopolymer (Norway). TPP, Angiotensin-I converting enzyme (from rabbit
123 lung), captopril, N- α -hippuryl-L-histidyl-L-leucine hydrate salt (HHL) and all other materials
124 were obtained from Sigma-Aldrich (Ireland). CellTitre 96® AQu_{eous} One Solution Cell
125 Proliferation Assay was supplied by Promega (Madison, USA). Caco-2 cells (passage 24-26)
126 were obtained from European Collection of Cell Cultures (Salisbury, UK). HepG2 cells
127 (passage 32-34) were obtained from American Type Culture Collection. Ultrapure water was
128 used for all experiments and was obtained from a Milli-Q water purification system
129 (Millipore Corporation, USA).

130 2.1 Unloaded nanoparticle formulation design

131 Unloaded NPs formulation was optimised using varying concentrations of CL113 and TPP,
132 following a 3 block Central Composite Design (CCD) with 2 variable parameters and 3
133 responses (particle size, ZP and PDI).

134 2.2 LKP nanoparticle formulation design

135 A Mixture Amount Design (MAD) was employed using the concentration ranges of chitosan
136 (CL113) and TPP (see table 1) around the optimal point (1.5mg/mL CL113 and 0.3mg/mL
137 TPP) suggested from preliminary unloaded particle experiments (Section 3.1). The

138 experimental design and data analysis was performed using Minitab 17 software (Minitab
139 Inc, USA).

140 **Table 1** MAD for LKP nanoparticles at optimised CL113, TPP concentration

Sample	CL113 (mg/ml)	TPP (mg/ml)	Ratio (CL113/TPP)
1	1.64	0.21	8.0
2	1.52	0.33	4.5
3	1.58	0.27	5.9
4	1.45	0.40	3.6
5	1.39	0.46	3.0

141

142 2.3 Preparation of LKP NPs

143 Preparation of LKP NPs was based on a modified ionotropic method (Calvo, Remu, Pez,
144 Vila-Jato, & Alonso, 1997; Vimal et al., 2013). Stock solutions of 10mg/mL CL113 and TPP
145 were prepared. CL113 was dispersed in acetate buffer (pH3) and TPP in 0.01M sodium
146 hydroxide solution. The stock solutions of CL113 and TPP were diluted to different
147 concentration ratios at a fixed volume mixture of 2.5:1 CL113: TPP containing solution. A
148 fixed concentration of 0.1mg/mL LKP was added to the diluted TPP solutions. The
149 TPP/peptide solution was added dropwise to the CL113 solution while stirring (800rpm for
150 30mins). NPs were separated using ultrafiltration-centrifugation (Centriplus YM-30, MWCO
151 of 30kDa, Millipore, USA). 10mL of sample were placed in the sample reservoir of the
152 centrifugal filter device and centrifuged for 30mins at 3000rpm. After separation, the volume
153 of the solution in the filtrate vial was measured and the filtrate was assayed for the amount of
154 LKP by Reverse Phase High Performance Liquid Chromatography (RP-HPLC). The wet

155 pellet was re-suspended in purified water and immediately characterised using a range of
156 physico-chemical techniques.

157 2.4 Physico-chemical characterisation of LKP NPs

158 2.4.1 Size, zeta potential and polydispersity index

159 The nanoparticle size (number distribution) and electrophoretic mobility measurements were
160 performed using folded capillary cells in a Nanosizer ZS fitted with a 633 nm laser (Malvern
161 Instruments Ltd.). Each analysis was carried out at 25°C with the equilibration time set to 2
162 min using size by intensity distribution.

163 2.4.2 Fourier transform infrared spectroscopy

164 The chemical properties of the NPs were monitored using Fourier transform infrared
165 spectroscopy (FT-IR), performed using a Perkin Elmer Spotlight 400 Series Spectrometer
166 (with Universal Attenuated total reflectance (ATR) accessory). FT-IR spectra of LKP,
167 unloaded NPs, and LKP NPs were obtained in the spectral range 650 to 4000 cm^{-1} in
168 triplicate. NP samples were stored at -80°C in glass vials and then lyophilised prior to
169 analysis using a Labconco FreeZone 6 Liter Benchtop Freeze Dry System.

170 2.4.3 Scanning electron microscopy

171 The morphology of the freeze-dried NPs was studied using scanning electron microscopy
172 (SEM) (Hitachi SU6600 FESEM), at an accelerating voltage of 20kV using the secondary
173 electron detector. The freeze dried NPs (0.5mg) were dispersed in deionised water (10mL)
174 and sonicated for 4min. One drop of the dispersion containing LKP NPs was placed on a
175 silicon wafer and dried at room temperature. This was sputter coated with 4nm Au/Pd prior to
176 imaging.

177 2.4.4 Determination of association efficiency and loading capacity of LKP nanoparticles

178 The association efficiency (AE) and loading capacity (LC) of NPs was calculated by the
179 indirect method of Al-Qadi *et al.* (2012). The supernatant was assayed for the content of LKP
180 by RP-HPLC. This quantity of LKP is referred to as the non-associated peptide. The RP-
181 HPLC analysis was performed on a Waters 1525 pump (Waters, Milford, Massachusetts)
182 with a Photo Diode Array detector 2487 (Waters) using a Luna C18 column (5 μ m, 250mm x
183 4.6mm, Phenomenex). Analytes were detected at the wavelength of $\lambda_{\text{max}}= 220\text{nm}$. The
184 column was eluted at a flow rate of $1\text{mL}\cdot\text{min}^{-1}$ with an isocratic system (15% Acetonitrile,
185 0.05% TFA in water). The AE% and loading capacity (LC %) was calculated using the
186 following equations.

$$187 \text{ AE \%} = \frac{(\text{Total amount Peptide} - \text{free amount Peptide in supernatant})}{\text{Total amount of Peptide}} \times 100 \quad (4.1)$$

$$188 \text{ LC\%} = \frac{(\text{Total amount Peptide} - \text{free amount Peptide in supernatant})}{\text{Nanoparticle weight}} \times 100 \quad (4.2)$$

189 2.4.5 ACE inhibition assay

190 The ACE inhibition of LKP was determined as previously described with minor
191 modifications (Henda *et al.*, 2013; Lahogue, Réhel, Taupin, Haras, & Allaupe, 2010). All
192 solutions were pre-filtered with 0.22 μ m nylon syringes prior to analysis. HHL (5mM) was
193 dissolved in pH 8.3 buffer (0.1M borate buffer in 0.3M NaCL). In a 96-well plate, 100 μ l
194 substrate solution and 25 μ l inhibitor were incubated for 10min at 37°C. 10 μ l ACE solution
195 (100mU/mL) was then added and incubated for another 30min at 37°C. The assay was
196 terminated using 100 μ l of 1M HCL. HPLC was performed using a C8 column (2.7 μ m, 3.0 x
197 100mm, Agilent Technologies UK & Ireland Ltd) and wavelength detection at $\lambda_{\text{max}}= 228\text{nm}$.
198 An isocratic method was used at a flow rate of $0.4\text{mL}\cdot\text{min}^{-1}$, 25% Acetonitrile, 0.1% TFA in
199 water for 5min. Controls were prepared by replacing the inhibitor with assay buffer (negative

200 control) and captopril (positive control). 100% ACE inhibition (negative control) was used to
201 calculate the % of ACE activity.

$$202 \text{ ACE inhibition (\%)} = \left(1 - \frac{A_{\text{inhibitor}}}{A_{\text{blank}}}\right) * 100 \quad (4.3)$$

203 where $A_{\text{inhibitor}}$ and A_{blank} are the peak areas of HA (product of HHL) and negative control,
204 respectively. The IC_{50} of the inhibitor was determined using the Hill-Step equation (Prism 5,
205 GrapPad Software Inc., USA).

206 2.4.6 Accelerated stability analysis

207 The optimal formulation was further analysed under accelerated stability conditions. 10mL of
208 a NP formulation equivalent to 19.5mg of NPs were resuspended in aqueous solution (pH7)
209 and stored at accelerated conditions; 60°C for 720min, 70°C for 300min and 80°C for
210 120min. The particle size and colloidal stability over different time intervals were measured
211 using the Nanosizer ZS (Malvern Instruments Ltd), and the order of degradation in aqueous
212 LKP suspension was determined. The end point of each sample was further assessed for ACE
213 inhibition activity (Method 2.4.5). Stability analysis was analysed using R software (R Core
214 Team, 2015).

215 The kinetic model used to describe the stability was of zero order. The temperature
216 dependence of the kinetic parameters of LKP NPs stability was measured by calculating the
217 observed rate constants. This was plotted according to the Arrhenius equation and apparent
218 activation energy, E_a and reaction rate, k_{ref} were calculated according to Equation 4 (Brauner
219 & Shacham, 1997).

$$220 C = C_0 + e^{\ln(k) - \frac{E_a}{R} \left(\frac{1}{T} - \frac{1}{T_{\text{ref}}} \right)} t \quad (4.4)$$

221 where C is the property (particle size or PDI) at time t, C_0 is the initial property conditions, k
222 is the apparent zero order reaction constant, E_a is the energy of activation, R is the universal

223 gas constant, T is the temperature of the experiment (K) and T_{ref} is the reference temperature
224 (70°C).

225 2.4.7 MTS assay

226 Caco-2, heterogeneous human epithelial colorectal adenocarcinoma cells and HepG2, a
227 human liver cancer cell line, were seeded at a cell density of 2×10^4 cells/well and cultured
228 on 96 well plates in DMEM and EMEM respectively, supplemented with 10% fetal bovine
229 serum, 1% L-glutamine, 1% penicillin-streptomycin and 1% non-essential amino acids, and
230 incubated for 24h at 37°C in a humidified incubator with 5% CO₂ and 95% O₂. Specified
231 exposure times were used for Caco-2 and HepG2, in order to mimic *in vivo* conditions. The
232 maximum time NPs will be exposed to the intestines are 4h, hence a 4h exposure time was
233 used in Caco-2 cell lines (Neves, Martins, Segundo, & Reis, 2016). In addition to this, 72h
234 exposure time was used for HepG2 cell line to mimic the liver (Brayden, Gleeson, & Walsh,
235 2014). LKP (native), unloaded NP and LKP NPs at 1, 5 and 10mM concentration were
236 assessed. Triton X-100™ (0.05%) was used as a positive control. After exposure, treatments
237 were removed and replaced with MTS (3-(4,5-dimethylthiazol-2-yl)-5-(3-carbo
238 xymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium. Optical density (OD) was measured at
239 490 nm. Each value presented was normalised against untreated control and calculated from
240 three separate experiments, each of which included six replicates.

241 2.4.8 *In vitro* controlled release studies

242 LKP release from loaded formulation was carried out using a dialysis bag diffusion technique
243 (Hosseinzadeh, Atyabi, Dinarvand, & Ostad, 2012) over 24h (Calderon *et al.*, 2013; Yoon *et*
244 *al.*, 2014). To ensure sink conditions, NPs were solubilised and sonicated 3 times for 20
245 seconds (Branson Ultrasonics; Ultrasonic processor VCX-750W, Wilmington, North
246 Carolina, USA). 5mL of LKP formulation was placed in the dialysis bag (cellulose ester

247 membrane, molecular weight cut-off 100kDa, Float-A-Lyzer[®]G2, Sigma-Aldrich, Ireland)
248 and immersed in a vessel containing 50mL of release fluid using simulated gastric fluid
249 (SGF) or simulated intestinal fluid (SIF) specified according to the British Pharmacopoeia,
250 respectively. SGF was composed of 0.1 M HCL and SIF, as the buffering stage, was
251 composed of 1 volume of 0.2 M trisodium phosphate dodecahydrate and 3 volumes of 0.1 M
252 HCL (adjusted to pH 6.8), without enzymes (British Pharmacopoeia Commission, 2016).
253 Each experiment was agitated at 100rpm, 37°C using a thermostatic shaker. At predetermined
254 time points over 24h, 1mL of release fluid was analysed and replaced with simulated fluid.
255 The LKP release was measured using RP-HPLC. The following equation was employed to
256 determine the % cumulative drug release:

$$257 \quad \% \text{ cumulative release} = \frac{\text{LKP release}}{\text{LKP initial}} * 100 \quad (4.5)$$

258 where LKP release and initial represents the concentration of LKP release and the amount of
259 LKP initially loaded into the NPs, respectively.

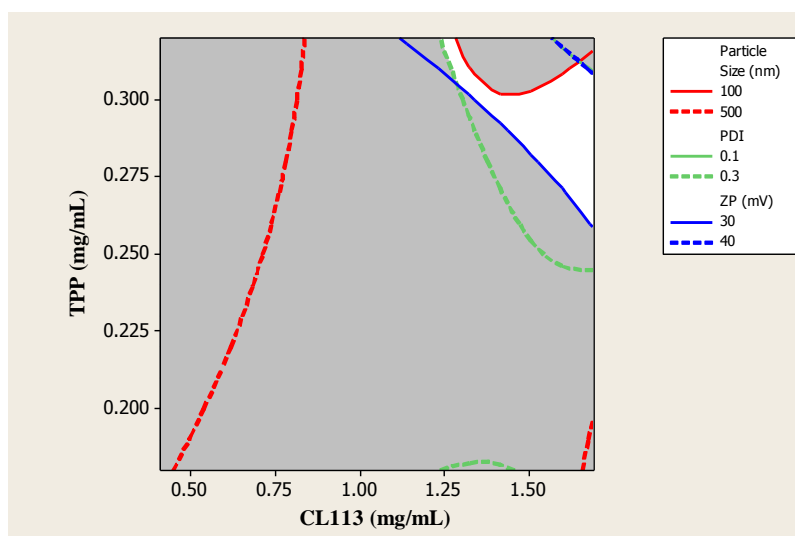
260 **3. Results and discussion**

261 LKP has an isoelectric point (pI) of 9.17, calculated using the Henderson-Hasselbach
262 equation (Henriksson, Englund, Johansson, & Lundahl, 1995). This is the net charge of a
263 molecule indicating that, at a pH of 9.17, LKP will have minimal solubility. At pH values
264 below the pI, peptides carry a net positive charge and above the pI, a negative charge. LKP
265 has a similar charge to TPP, and hence LKP was dissolved in the TPP solution. TPP-LKP
266 was then added dropwise to the CL113 solution, resulting in the formation of opalescent NPs.

267 **3.1 Unloaded nanoparticle production feasibility zone identification**

268 The formulation of CL113 NPs was optimized using a CCD factorial design to analyse the
269 effect of the pH, CL113: TPP ratio and acetic acid concentration on the size and ZP of the
270 particles. Preliminary studies were conducted to select the most feasible region for the

271 formulation of peptide NPs (figure 1). This showed that > 1.25 mg/mL CLL113 and 0.25-0.3
272 mg/mL TPP resulted in unloaded NPs of optimal sizes of 300 nm and ZP > 30 mV.

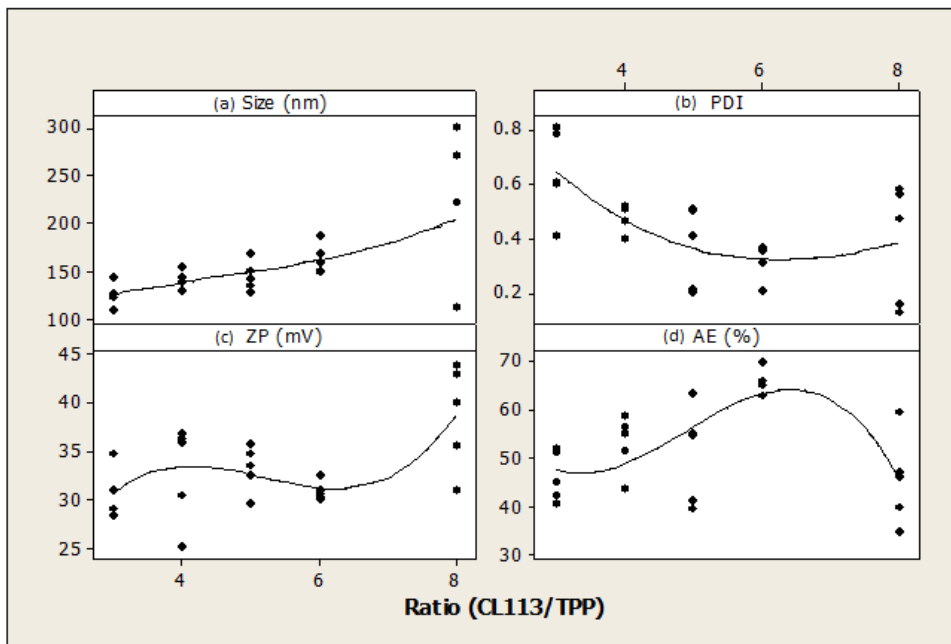


273
274 **Figure 1** Overlaid contour plots of Size and ZP: the targeted area is highlighted (white).

275 This result is in agreement with Calvo et al., (1997) who reported that concentrations which
276 exceed 4mg/mL and 0.75mg/mL respectively for chitosan and TPP resulted in the formation
277 of large aggregates. Nanoparticles of the desired characteristics were found to be produced at
278 concentrations of 0.28 mg/mL (TPP) and 1.25 mg/mL (Chitosan). In addition, studies done
279 by de Pinho Neves et al. (2014) showed similar results within the 5 to 6:1 ratio. Unloaded
280 CL113 NPs were used as control for peptide-loaded experiments.

281 3.2 Nanoparticle size, zeta potential analysis and AE % of LKP nanoparticles

282 From the preliminary optimisation analysis of unloaded CL113 NPs; a CL113 concentration
283 of 1.5mg/mL was chosen as the centre point for the MAD. LKP NP size values ranged from
284 120 to 271nm, with ZP values above 30mV. With regards to the AE % (figure 2), at ratios
285 above 5.9, there is less variability (error bars), but at a ratio of 8, stable agglomerates (>
286 30mV ZP) of variable sizes are observed.



287

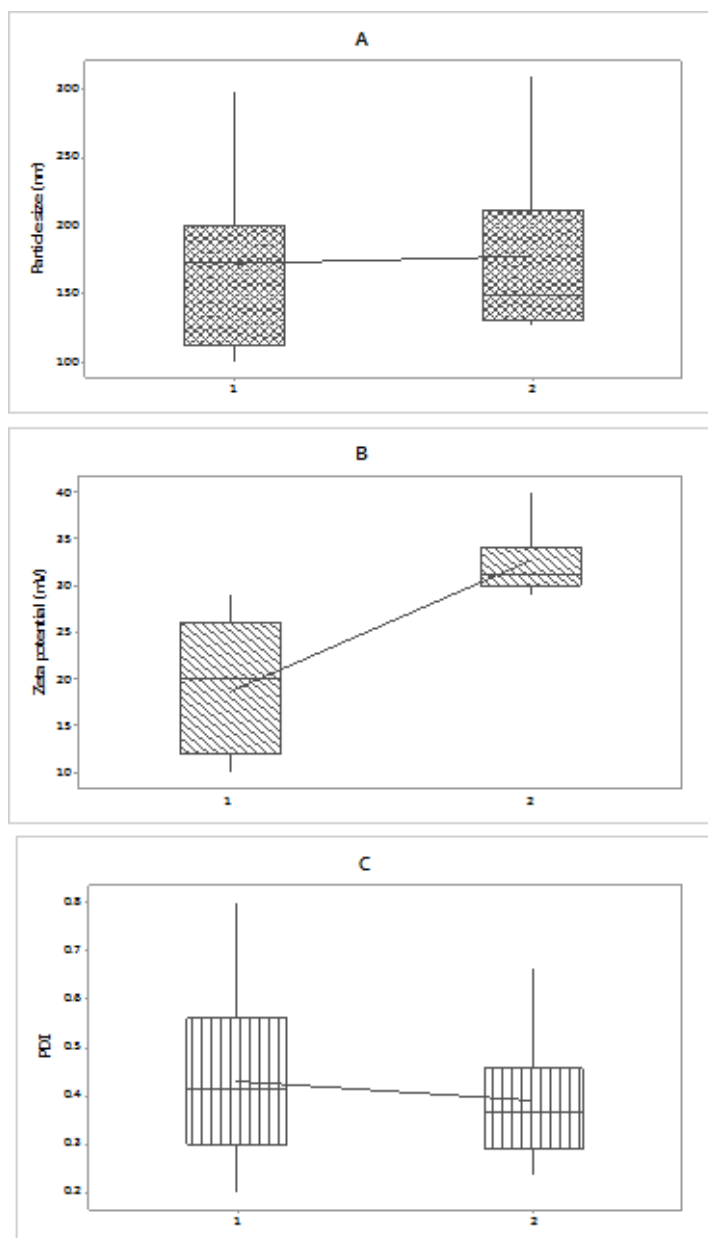
288 **Figure 2** Scatterplot of (a) Size (nm), (b) PDI, (c) ZP (mV) and (d) association efficiency
 289 (AE %) of LKP NPs against different ratios of (CL113/ TPP). Error bars represent the
 290 individual 95% Confidence interval for the average

291 An initial observation of the results at different ratios seems to indicate that the ratio of 5.9 is
 292 the best performing in terms of higher AE% and lower PDI, while still maintaining a particle
 293 size (150nm) and a ZP (>30mv) that indicates stability (figure 2). LC% for all experiments
 294 showed no significant differences, values ranged from 2-3% (see supplementary material, S2)
 295 were attained. Studies from other groups showed that an increase of counterion (TPP)
 296 concentration results in a decrease of LC due to the high level of crosslinking, causing the
 297 encapsulated material to come out of the particle (Woranuch & Yoksan, 2013).

298 Similar profiles are seen for the 3 responses, size (nm), ZP (mV) and PDI, suggesting
 299 agglomeration and colloidal instability above a ratio (CL113/TPP) of 7. The degree of
 300 crosslinking can be assessed by the chitosan concentration and the available NH₃⁺ able to be
 301 crosslinked with TPP functional groups. At intermediate operation conditions, i.e. ratios
 302 between 7 and 5.5, most responses present a lower variation between replicates with respect

303 to all responses and a minimum PDI. Analysis for all experimental results shows particle
304 sizes within the optimal size range (100-500nm) and ZP above 30mV values. However, for
305 some experimental conditions, the AE% and PDI exhibited values far from the optimal
306 physico-chemical characteristics and had high variability between replicates (N=3). At
307 (CLL113/TPP) ratios above 7 and below 4.5, the NPs exhibited higher PDI and, below a ratio
308 of 6, significant variability of AE% amongst replicates is evident.

309 The results obtained were consistent with those observed for unloaded CL113 NPs in terms
310 of particle size, although a significant increase in colloidal stability for the loaded NPs was
311 observed (see figure 3).



312

313 **Figure 3** Boxplot for LKP NPs where each Group represents 1: control (unloaded NPs) and

314 2: LKP NPs. The changes of each group are examined for A: particle size, B: ZP and C: PDI.

315 A significant increase in ZP is seen for loaded NPs with a decrease of variability.

316 For the MAD design, a third order polynomial regression model was employed to describe

317 the variation of size (nm), ZP (mV), PDI and AE % of the LKP NPs against the ratio

318 (CL113/TPP). The polynomial equation parameters for each response against ratio were

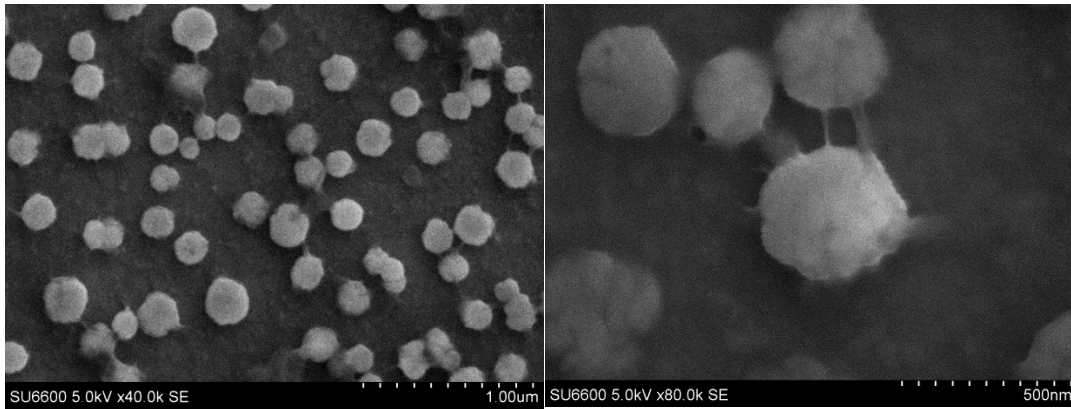
319 fitted. The models were built with the aim of identifying conditions within the experimental

320 design where NPs would be present as monodispersed, stable NPs with maximum peptide
321 encapsulation (see supplementary information S1).

322 From the results obtained, a ratio of 5.9 provided the most promising results, for optimal oral
323 delivery fulfilling the formulation constraints; 100-500 nm, PDI <0.4 and |ZP| >30 mV. In
324 comparison, NPs produced at a ratio of 7.8, presented the highest variability for size ($206 \pm$
325 73 nm), PDI (0.4 ± 0.2), ZP (38 ± 6 mV) and AE ($41 \pm 11\%$). Notably, the NPs produced at a
326 ratio of 5.9 (CL113/TPP) yielded a substantially higher AE % of around 65%. It should be
327 noted that the LKP of isoelectric pH (9.17) was added to a higher pH TPP (pH 12) solution.
328 This provided more negatively charged molecules to interact with chitosan, consequently
329 increasing the AE % (Acton, 2012). This finding is in agreement with Silva et al.,(2013), who
330 observed higher AE % of daptomycin at higher pH values relative to the isoelectric pH. In
331 addition, LKP has a low Mw and studies of lower Mw actives showed higher AE %. This
332 trend was observed by Jarudilokkul et al., (2011), who showed that α -Lactalbumin (17.4 kDa)
333 has higher AE % than Fibrinogen (340 kDa).

334 3.3 Morphological characterisation of LKP chitosan nanoparticles

335 Further characterisation of the NPs produced with the optimal (CL113/TPP) ratio of 5.9 was
336 performed. Figure 4 represents an SEM image of LKP formulation, confirming the formation
337 of the NPs. Spheroidal NPs were obtained, of sizes ranging from 150-250nm, consistent with
338 the DLS measurement.



339

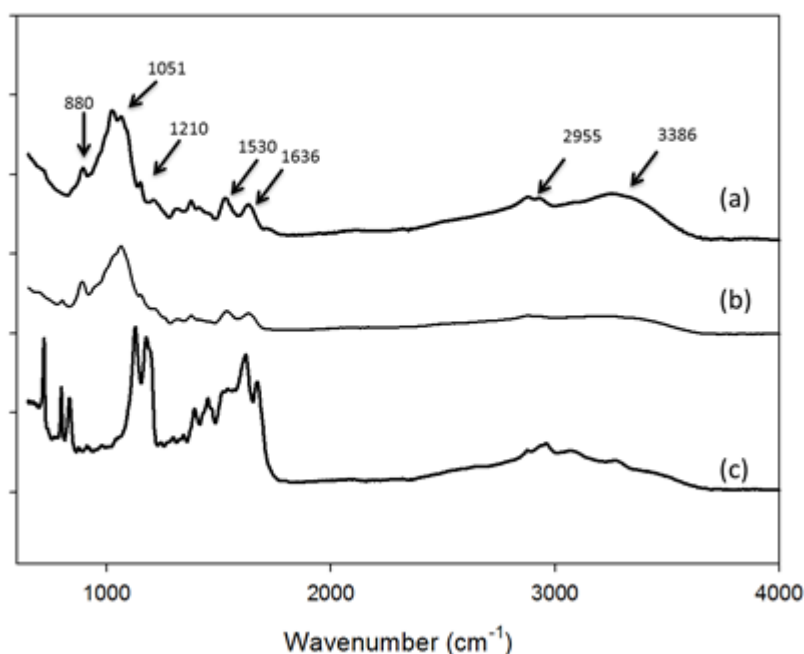
340 **Figure 4** SEM image of optimal formulation (Ratio 5.9) of LKP NPs

341 3.4 Chemical Characterisation of LKP nanoparticles

342 FT-IR was used to identify whether there were variations in chemical functional groups
343 presented in the LKP loaded NPs with respect to their raw materials. An FT-IR analysis was
344 conducted for pure LKP powder, unloaded NPs and LKP NPs (Figure 5). Chitosan NPs have
345 been previously characterised using FT-IR (Mohammadpour Dounighi et al., 2012;
346 Sureshkumar et al., 2010; Vimal et al., 2013).

347 Comparison of the FT-IR spectra of loaded and unloaded NPs indicated that the spectrum of
348 the unloaded NPs is largely unchanged by the presence of LKP, as may be expected due to
349 the relatively low LKP content. Characteristic peaks of unloaded NPs are seen in both, at
350 1530 , 1636 and 880cm^{-1} , representing the amide I and amide II bands of CL113 and pyranose
351 (P-O) of TPP. For the optimal formulation, 18.5mg of NPs is needed to encapsulate 1mg of
352 LKP. Hence, no distinctive peaks of LKP can be seen in the LKP spectrum. The FT-IR
353 spectrum of LKP NPs does, however, show some changes from that of the unloaded NPs,
354 potentially indicative of localised conformational changes of the CL113 as a result of
355 interaction with the LKP (figure 5). An increased absorption at 3386cm^{-1} compared to that of
356 the unloaded NPs is observed. Absorption in this region of the spectrum represents O-H
357 bonding; a possible explanation may be due to the interaction of the hydrogen acceptors (O-

358 in LKP) and hydrogen donors (NH_3^+ in chitosan). In addition, increased absorption is also
359 seen for LKP NPs at 2955cm^{-1} , representing an increase in (C-H) hydrogen bond stretching
360 with presence of the peptide. A shift of 1605cm^{-1} to 1530cm^{-1} was also observed which
361 represent the amide carbonyl stretch (Mohammadpour Dounighi et al., 2012). The peaks at
362 1210cm^{-1} and 880cm^{-1} represent phosphate group (P-O) and pyranose ring (Woranuch &
363 Yoksan, 2013). The peak at 1051cm^{-1} shows a split for the loaded NPs, suggesting a
364 conformational change due to the interaction with the LKP.



365
366 **Figure 5** FT-IR spectra of (a) LKP NPs (b) unloaded NPs and (c) Pure LKP powder.
367 Absorbance spectra are normalised and offset for clarity.

368 3.5 IC_{50} determination of inhibitor

369 The inhibitory activities of captopril (reference molecule) and LKP were determined using a
370 synthetic substrate, HHL, and varying concentrations over the range $0.0001 - 10\mu\text{M}$
371 (captopril) and $0.001 - 10\mu\text{M}$ (LKP). The IC_{50} obtained was $0.006 \pm 0.002 \mu\text{M}$ for captopril
372 and $0.30 \pm 0.08 \mu\text{M}$ for LKP. These values are consistent with previously reported values,

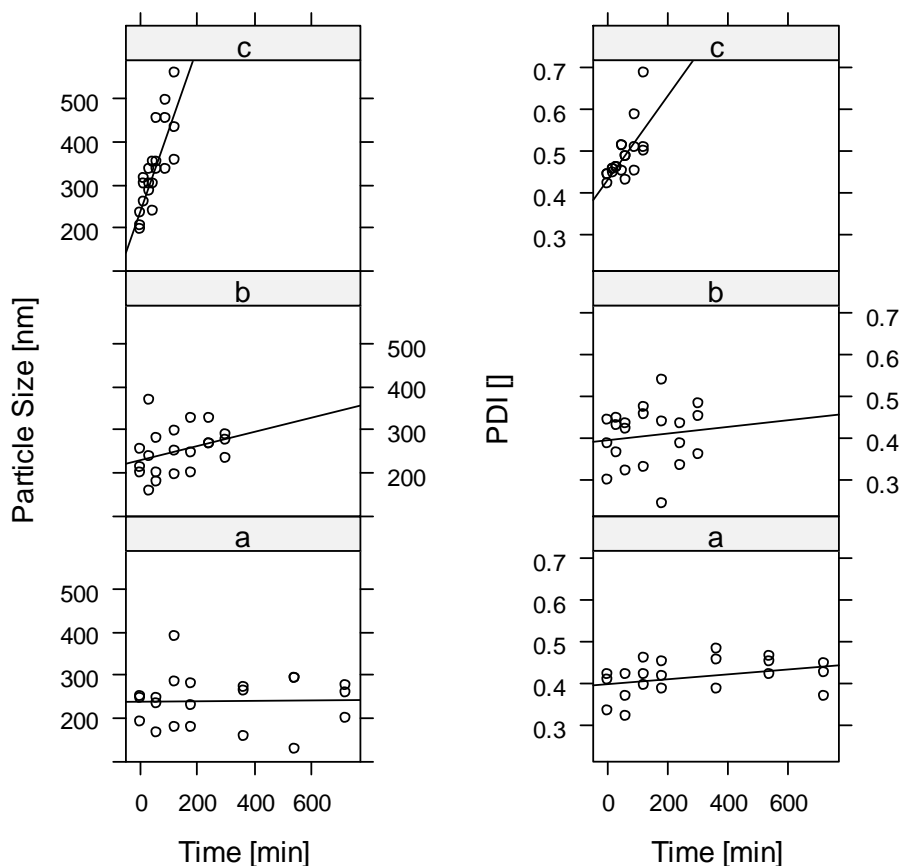
373 which range from 0.001-0.039 μ M for Captopril and 0.2-0.32 μ M for LKP (Fujita &
374 Yoshikawa 1999; Fujita *et al.* 2000; Henda *et al.* 2013).

375 3.6 Accelerated stability analysis of LKP NPs

376 In accelerated stability testing, a product is stressed at high temperatures and
377 degradation/stability of the product at normal storage conditions is then predicted (Bajaj,
378 Singla, & Sakhuja, 2012; Rauk, Guo, Hu, Cahya, & Weiss, 2014; Waterman & Adami,
379 2005). A number of factors can affect the solution stability of NPs, for example, the pH of the
380 aqueous solvents, light, oxygen, co-solutes, buffer salts, surfactants and antioxidants.
381 Common degradation routes include hydrolysis/solvolysis, photolysis/oxidation and
382 racemisation (Weber, Coester, Kreuter, & Langer, 2000). A number of groups have found
383 that CL113 NPs synthesised by ionic gelation lose their integrity in aqueous media, even in
384 the absence of enzymes (López-León, Carvalho, Seijo, Ortega-Vinuesa, & Bastos-González,
385 2005). Jonassen *et al.* (2012) looked at the effect of different ionic strength over the course of
386 a month; the main findings were that the most stable NPs with respect to the size and
387 compactness of the particles were produced in saline conditions (Jonassen, Kjøniksen, &
388 Hiorth, 2012). A similar study was also conducted, preparing NPs in different ionic strength
389 and buffers, and results showed that the least stable NPs were produced in non-buffered
390 solutions or low ionic solutions (López-León *et al.*, 2005).

391 The stability of formulations can be tested using a number of testing protocols, which include
392 real time stability testing, accelerated stability testing, retained sample stability testing and
393 cyclic temperature stress testing (Bajaj *et al.*, 2012). In the current study, accelerated stability
394 testing was used, by which NPs are subjected to stress and then assayed simultaneously to
395 predict the likelihood of instability based upon the Arrhenius equation. Suspensions of NPs in
396 buffered solutions (PBS pH 7), formulated at a ratio of 5.9 CL113:TPP (optimal LKP loaded

397 NPs), were exposed to three different storage temperatures, 60°C, 70°C and 80°C, over a
398 time course of 120, 300 and 720min at each temperature. Figure 6 shows the kinetic
399 behaviour of the particle sizes at different temperatures. The stability of the NPs decreased
400 with increasing temperature. At 60°C, no change in particle size was observed over the
401 720min. Figure 6 indicates a more pronounced increase in both particle size and PDI at 70°C,
402 while at 80°C, both particle size and PDI increase significantly over the time course. At this
403 temperature, the particle size was seen to increase monotonically from 200 to 600nm, while
404 the PDI increases from 0.4 to 0.7. While some curvature is apparent in the trends at the
405 highest temperature, within the present experimental error, an apparent zero order mechanism
406 fitted better to all the data, compared to an apparent first or second order model.



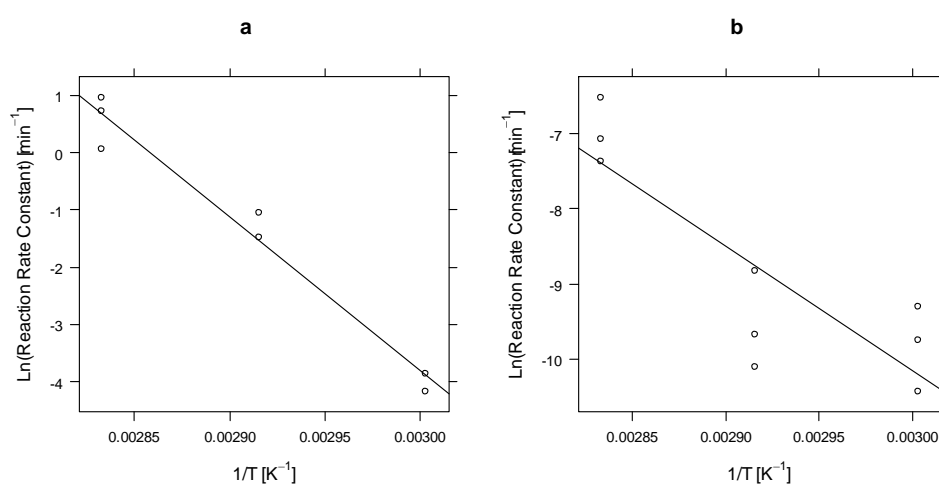
407

408 **Figure 6** Particle size and PDI analysis of LKP loaded NPs exposed at (a) 60°C (b) 70°C and
 409 (c) 80°C over time periods of 120, 300 and 720min, respectively. N= 3

410 An Arrhenius plot of the apparent zero order reaction rate constants, derived from the
 411 analysis of the individual experiments at the different temperatures, indicates that the kinetics
 412 of the particle size and PDI followed this temperature relationship, consistent with an energy
 413 activated process, with similar energies of activation for particle size and PDI (see Figure 7).

414 The one-step nonlinear regression analysis of the kinetic experiments shows that the particle
 415 size fits to a zero order kinetic behaviour and an Arrhenius dependence with $\ln(k_{\text{ref}@70\text{C}}) = -$
 416 $3 \pm 1 \text{ min}^{-1}$ and an E_a of $360 \pm 103 \text{ kJ/mol}$. For the PDI, a one-step nonlinear regression was
 417 fitted to zero order kinetics with an Arrhenius dependence of $\ln(k_{\text{ref}@70\text{C}}) = -8.9 \pm 0.3 \text{ mins}^{-1}$
 418 1 and $E_a = 196 \pm 33 \text{ kJ/mol}$. A linear correlation is evident between $1/T$ and $\ln k$ in figure 7.

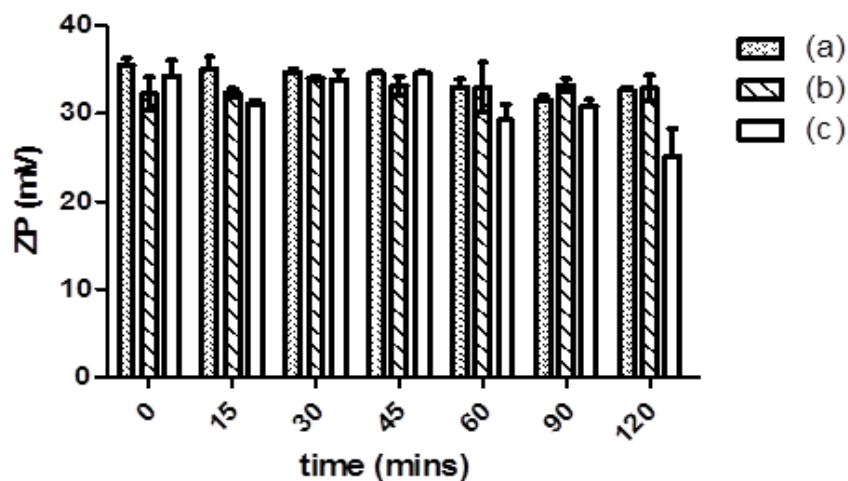
419 From this analysis, it suggests the nanoparticle formulation would be stable (in terms of
420 particle physico-chemical properties) in a neutral pH solution at ambient storage temperature,
421 with negligible increases in particle size or PDI of the NPs, confirming the higher stability of
422 NPs prepared in buffered solutions. Only after 90min destabilisation of NPs was observed at
423 the highest temperature conditions. These results are in agreement with previous literature,
424 reporting that when NPs are produced in salt or buffered environment, stability is improved
425 (López-León et al., 2005).



426

427 **Figure 7** Arrhenius Plots for the (a) Particle Size and (b) PDI accelerated studies of LKP
428 loaded NPs. N=3

429 The results suggest either (i) swelling of the chitosan NPs in the aqueous environment (Bajpai
430 & Maan, 2012) or (ii) agglomeration of chitosan NPs due to electrostatic interactions with an
431 increase of temperature. In order to confirm the type of degradation, the zeta potential was
432 assessed. Destabilisation of NP suspensions which would give rise to aggregation should be
433 reflected in changes in ZP values when NPs aggregate. However, no significant changes were
434 seen for the colloidal stability (i.e. ZP of LKP NPs). This suggests that swelling could be the
435 primary mechanism of the CL113 nanoparticle changes with time, followed by destabilisation
436 at higher temperatures and longer time (see figure 8).



437

438 **Figure 8** Zeta potential analyses of LKP-loaded NPs at (a) 60°C, (b) 70°C and (c) 80°C. No
 439 significant changes were observed with One-Way ANOVA with Dunnetts's post-test. Each
 440 value represents the mean \pm SD (n=3)

441 In addition, DD of CL113 used for this experiment is greater than 85%, it has been
 442 previously reported that with an increase of DD the aggregation stability decreases due to the
 443 CL113 more prone to TPP bridging which causes it to become more lyophobic near
 444 physiological pH which may contribute to the stability of the NPs (Haung, Cai, & Lapitsky,
 445 2015). Overall, the accelerated experiments on LKP NPs indicate that the formulations will
 446 be stable under normal storage conditions. This could be due to the conditions (presence of
 447 salt) used to prepare the NPs, in addition to the strong bonding between the bioactive and
 448 complex.

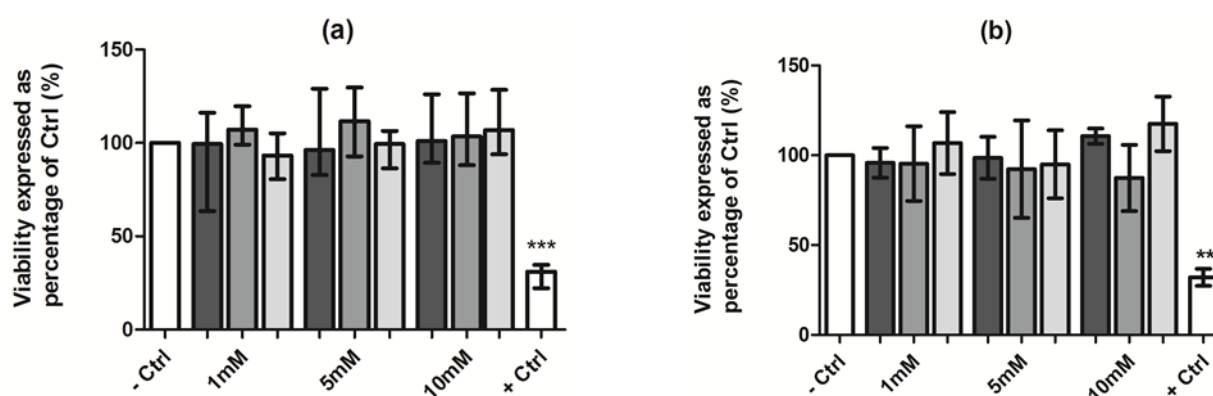
449 3.7 Cytotoxicity assessment of LKP nanoparticles

450 The MTS assay was used to assess the cytotoxicity of LKP and LKP NPs. The therapeutic
 451 dose of LKP is 10mg/kg (Fujita et al., 2000; Fujita & Yoshikawa, 1999b). Hence, LKP
 452 loaded or unloaded NPs at the different concentrations (1, 5 and 10mM) when exposed to
 453 Caco2 (4h) and HepG2 (72h) cell lines. No cytotoxicity was observed for LKP, indicating

454 negligible overall cytotoxicity of the formulation (figure 9). No significant changes were
 455 observed. A number of deviations were observed above 100% viability for NPs with and
 456 without LKP, proliferation may occur due to increase in fibroblasts production caused by the
 457 presence of polymeric chitosan (Rajam, Pulavendran, Rose, & Mandal, 2011) or interference
 458 of NPs with the cell assay (Casey et al., 2007).

459

460



461

462 **Figure 9** Cytotoxicity assessment of LKP, unloaded NPs and LKP NPs

463 exposed for (a) 4h in Caco2 cell lines and (b) 72h in HepG2 cell line at 1mM, 5mM and
 464 10mM concentration. Percentage (%) of MTS converted was compared to untreated control.

465 1-Way ANOVA with Dunnetts's post-test *** P< 0.001, ** P< 0.01, N=3

466 3.8 *In vitro* release studies

467 The release of a bioactive can take place by several different mechanisms, for example
 468 surface erosion, disintegration, diffusion and desorption (Hosseini, Zandi, Rezaei, &

469 Farahmandghavi, 2013). Such a phenomenon can be influenced by a number of factors such

470 as the type of polymer used, the polymeric swelling capability, the solute diffusion and

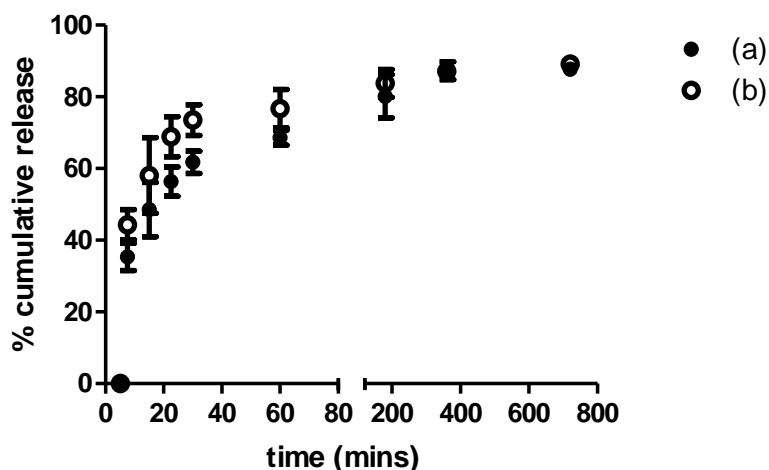
471 material degradation (Fu & Kao, 2009; Siepmann & Göpferich, 2001). The *in vitro* LKP

472 release profiles of the NP formulation in SGF and SIF were measured over 24h, using RP-

473 HPLC at 220nm at different time points. The site of target for LKP is in the jejunum, small

474 intestine, therefore it is important to bypass the acidic stomach environment. LKP NPs results
475 showed an initial burst followed by a slow release. Similar results were reported by other
476 groups, Hosseini *et al.* 2013 and Luo *et al.* 2010. Hosseini *et al.* 2013, they observed a
477 “biphasic release” mechanism (initial burst followed by slower release) with oregano
478 essential oil when encapsulated into CL113 NPs. Luo *et al.* 2010, encapsulated selenite in
479 chitosan NPs, demonstrating the effect of CL113 concentration on the release profile. They
480 found that, at high concentrations of chitosan, more dense particles were found which
481 ultimately lowered the epithelial membrane permeability. Conflicting results have been
482 reported of the release mechanism of chitosan nanoparticles. Some groups observed the
483 inability of chitosan based nanoparticles to sustain the release of an active following an initial
484 burst release at higher ratios of CL113 and TPP (Stoica & Ion, 2013). Others reported a
485 controlled release; for example, Nallamuthu, Devi, & Khanum, (2015) observed the 69%
486 release of chlorogenic acid over 100 h. Release profiles at 1.5mg/ml CL113 also showed a
487 burst within the first 30min in the SGF followed by a slower release from 30min to 2h then a
488 more slow sustained release up to 24h, as shown in figure 10. For LKP NPs, the initial burst
489 may possibly represent loosely bound LKP around the CL113 NP. Within the first hour, 62%
490 ± 3 and 74% ± 4 LKP was released in (a) SGF and (b) SIF, significance was observed for SGF
491 vs SIF using a t-test, where $P = 0.0074$, respectively (figure 10). 12% more release was
492 observed in the SIF (site of target) at pH 6.8. Chitosan has an isoelectric point of 6.5, at
493 which pH chitosan holds a charge of zero (no charge), causing it to become unstable and
494 precipitate out; consequently releasing more of the loaded peptide. It has been suggested by
495 Gan & Wang, (2007) that a burst release of protein molecules may correspond to the fast
496 swelling and degradation of the nanoparticles. This swelling phenomenon can be observed in
497 (figure 10) After 1h, a slower release is observed, this may be attributed to the more strongly
498 bound LKP loaded within the CL113 nanoparticle. A burst release has previously been

499 reported from other groups working with chitosan NPs, Sarmiento *et al.* (2007) showed a
500 similar undesirable burst of 50% insulin chitosan NPs when complexed with alginate. Ryan *et*
501 *al.* (2013) also reported an initial burst of 40% of salmon calcitonin when complexed into a
502 NPs system using chitosan and hyaluronic acid. For oral delivery systems, NPs remain in the
503 system for up to 6h after intake. Our studies show after 6h up to 85% release was obtained.
504 However, a prolonged release is desired. Additional of an outer surface coating is a technique
505 which has been widely used by a number of researchers in order to improve the integrity of
506 the NPs and to better control the release profile (Elgadir *et al.*, 2015). A popular approach to
507 yield coated chitosan NPs is by polyelectrolyte complexation, which exploits the interaction
508 between positively charged chitosan and negatively charged polyelectrolytes such as alginate
509 (Garrait, Beyssac and Subirade, 2014), dextran (Sarmiento *et al.*, 2006), hyaluronic acid
510 (Mero *et al.*, 2014) or zein (Luo, Teng and Wang, 2012). Further work on LKP encapsulation
511 will involve the polyelectrolyte complexation of LKP to achieve the desired release.
512 .



513

514 **Figure 10** Cumulative release profile of LKP from NPs in (a) SGF and (b) SIF for 24h.

515 **4. Conclusions**

516 LKP loaded NPs were formulated successfully by applying the ionotropic gelation technique.
517 The optimal NPs were found at a ratio 5.9:1 (CL113: TPP), using the design of experiment
518 approach, which resulted in reproducibility of the desirable physico-chemical characteristics.
519 Optimally LKP NPs were spheroidal particles of size ~200nm, as shown by SEM, and had
520 enhanced colloidal stability compared to the unloaded particles. A 5.9:1 ratio provided high
521 encapsulation efficiency of $65\pm 3\%$ with loading capacity of approximately $5\pm 0.8\%$. Stability
522 analysis showed long term physico-chemical stability. ACE inhibitory studies presented no
523 change in bioactivity of LKP over the different temperature conditions and after formulation
524 indicating it is quite stable. In addition, no cytotoxicity was observed from both the LKP
525 loaded and unloaded NPs. *In vitro* release studies indicated an initial burst within 1h,
526 suggesting the presence of more loosely bound tripeptide in the nanoparticle complex,
527 followed by peptide bounded within the nanoparticle which is released at a slower rate.
528 Chitosan based delivery systems is a feasible for the formulation of bioactive peptide,
529 physicochemical analysis and stability was efficient. The present results indicate that the
530 addition of an enteric coating to is recommended in order to bypass the stomach acidic pH
531 conditions, providing an efficient delivery system.

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773

Supplementary Material

S1: Fitted results of 3rd order polynomial regression against the PDI, ZP, AE % and particle size for LKP-loaded NPs

Polynomial regression equation: $a + b\text{Ratio} + c\text{Ratio}^2 + d\text{Ratio}^3$	Coefficient of determination (R^2)
Size (nm) a= - 284.90 c= - 64.68 b= 293.40 d= 4.51	$R^2 = 85.6\%$
PDI a = 2.31 c = 0.12 b= - 0.88 d = 0.01	$R^2 = 83.2\%$
ZP (mV) a = - 16.80 c = - 6.36 b = 31.05 d = 0.42	$R^2 = 76.1\%$
AE (%) a = 335.60 c = 43.79 b = - 202.40 d = - 2.90	$R^2 = 50.2\%$

S2: Loading capacities of LKP-loaded NPs, n=3

Sample	CL113 (mg/ml)	TPP (mg/ml)	Ratio (CL113/TPP)	LC %
1	1.64	0.21	8.0	2.2±0.01
2	1.52	0.33	4.5	3.3±0.00
3	1.58	0.27	5.9	2.3±0.08
4	1.45	0.40	3.6	2.7±0.03

5	1.39	0.46	3.0	2.8±0.02
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