

# Investigation Into The Effect of Varying L-leucine Concentration on the product characteristics of Spray-dried Liposome Powders

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- 1 <u>Title:</u> Investigation Into The Effect of Varying L-leucine Concentration on the product
- 2 characteristics of Spray-dried Liposome Powders
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- 11

#### 12 Abstract

Spray-dried formulations offer an attractive delivery system for administration of drug 13 14 encapsulated into liposomes to the lung, but can suffer from low encapsulation efficiency and poor aerodynamic properties. In this paper the effect of the concentration of the 15 anti-adherent I-leucine was investigated in tandem with the protectants sucrose and 16 17 trehalose. Two manufacturing methods were compared in terms of their ability to offer small liposomal size, low polydispersity and high encapsulation of the drug indometacin. 18 Unexpectedly sucrose offered the best protection to the liposomes during the spray drying 19 process, although formulations containing trehalose formed products with the best powder 20 21 characteristics for pulmonary delivery; high glass transition (Tg) values, fine powder fraction (FPF) and yield. It was also found that L-leucine contributed positively to the characteristics 22 23 of the powders, but that it should be used with care as above the optimum concentration of 24 0.5% (w/w) the size and polydispersity index increased significantly for both disaccharide formulations. Relating to the method of manufacture it was found that while both the 25 sucrose and trehalose conferred protection on the liposomes produced using either method 26 the ethanol-based proliposome method offered improved drug incorporation and did not 27 suffer from loss of drug caused by dilution effects. 28

29 (200 words)

30

31 Keywords. liposomes; spray drying; disaccharide; leucine

2 R

#### 1. Introduction 32

Liposomes are suited to encapsulation of a variety of drugs from small molecular 33 weight compounds to macromolecules and including both hydrophilic and lipophilic entities. 34 This is reflected in the range of therapeutics that have been tested in liposomal 35 formulations including cytotoxic agents [3]; bronchodilators and anti-asthmatics [4; 5]; 36 antibiotics [6] as well as photosensitizing agents [7] and genetic material [8-10]. The use of 37 spray drying to produce stable powder formulations for pulmonary administration is 38 attractive since it offers several advantages over the parenteral route [1]. Aerodynamic 39 diameter is a crucial factor in determining deposition of particles in the different sites of the 40 respiratory tract. Particles in the range 1-6 µm are best suited to deposition in the lower 41 airway following inhalation [2]; those with diameters > 6  $\mu$ m are deposited in the oropharynx, 42 whereas smaller particles (< 1  $\mu$ m) are exhaled during normal tidal breathing. In addition, 43 fine particle fraction (FPF, the fraction of powder emitted from the inhaler with a particle size 44  $\leq 5 \,\mu$ m) is a critical parameter to predict the proportion of the emitted dose that can deliver 45 deeply into the lower respiratory system. The Use of liposomes as carriers offers benefits 46 including protection of drug from enzymatic degradation; prolonging retention time and 47 reducing side effects. 48

Many methods are available for the manufacture of liposomes, including thin-film 49 hydration [11], organic solvent injection [12], reverse-phase evaporation [13] and 50 51 dehydration-rehydration [12; 14]. Both the ethanol injection and proliposome methods of 3

52 liposome preparation offer good potential for scale-up and have been used to encapsulate a wide range of substances [15-25] [26-31]. The ethanol injection method involves the 53 rapid injection of a lipid-ethanol solution into an excess of aqueous medium to. 54 spontaneously form large unilamellar vesicles (LUVs) [12]. Advantages of the technique 55 include simplicity and low risk of lipid degradation or oxidation. The ethanol-based 56 proliposome method is based on the preparation of hydrated stacked bilayer sheets in a 57 water-ethanol solution termed proliposomes. Spontaneous formation of liposomal 58 suspensions (multilamellar vesicles, MLVs) is achieved by addition of excess aqueous 59 solution to a lipid mixture [26]. The MLVs produced can be further processed for the 60 preparation of oligolamellar and unilamellar liposomes. 61

62

The effect of disaccharide protectants on the stability of spray dried liposomes has 63 been examined previously [32-34], while amino acids have been shown to play an 64 important role in improving the aerosol behaviour of spray-dried powders by reducing 65 moisture sorption and surface tension of dried particles [37; 38]; they can also protect 66 proteins against thermal stresses and denaturation [39]. Leucine, in particular, tends to 67 improve powder aerosol properties, this anti-adherent effect has been attributed to its 68 hydrophobic character and surfactant-like properties that allow it to migrate rapidly to the 69 70 surfaces of the particles during drying [40; 41] and it has been shown to interact well with 71 lipid membranes [42]. Two studies have used leucine in combination with sugars at very 72 high fixed concentrations [43; 44]. To the best of our knowledge, the effect of varying leucine concentration on the spray drying of liposomal systems has not been investigated 73 previously. In this paper we demonstrate the effect of varying concentrations of the 74 anti-adherent L-leucine in combination with optimised concentrations of disaccharides and 75 show for the first time that formulation effects can lead to significant differences in the 76 product characteristics for spray dried liposomes, especially in terms of for sucrose 77 formulations. Furthermore, at higher concentrations leucine increases liposome size. 78 Additionally, we investigated whether switching from the ethanol injection to the 79 proliposome method of liposome preparation affected liposome properties or drug release. 80 Indometacin was chosen as it has been used previously as a model drug in liposomal 81 82 studies [45; 25].

83

#### 84 2. Materials and methods

#### 85 2.1 Materials

Cholesterol (PhEur grade), sucrose (>99%), L-Leucine (PhEur grade), NaCl (≥99%), Trifluoroacetic acid (99%), Iron (III) chloride hexahydrate (≥99%), Ammonium thiocyanate (≥99%), Sephadex G-50 (20-80 µm), Indometacin (≥99%) phosphotungstic acid hydrate and ethanol (≥99.5%) were purchased from Sigma Chemicals.  $\alpha$ , $\alpha$ -Trehalose Dihydrate (high purity, low endotoxin) was purchased from Ferro Pfanstiehl. Soy phosphatidylcholine (LIPOID S 75, 80%) and soy phosphatidylcholine (Lipoid S PC, >98%) were purchased from Lipoid. Methanol (HPLC grade, >99.8%) and chloroform (99.2%) were purchased from
VWR. Phosphate buffered saline tablets was purchased from Invitrogen Corporation. All
aqueous solutions were prepared with de-ionized water.

- 95
- 96 **2.2 Methods**

#### 97 2.2.1 Preparation and drying of liposomal dispersions

# 98 **2.2.1.1 Preparation of liposomes by ethanol injection**

99 Small unilamellar vesicles (SUV's) were prepared by ethanol injection method followed by sonication in order to reduce liposomal size. Each batch was prepared on a 10g scale. 100 101 0.02 g Indometacin was dissolved in 0.7 g ethanol (at 57 °C) together with the lipids composed of 1g SPC and 0.115 g cholesterol. Hydration media were prepared by 102 dissolving varying amounts (2.5, 5, 7.5, 10 and 15% w/w) of disaccharides (sucrose or 103 trehalose) in 0.9% NaCl solution. Once the optimum concentration of these protectants 104 had been determined, liposomal dispersions were prepared using various concentrations 105 (0.25, 0.5 and 1% w/w) of L-Leucine added into the hydration medium also containing the 106 optimum concentration of each protectant. The lipid solution was rapidly injected into 107 8.165g of hydration medium at room temperature with stirring. After 2 hr hydration, the 108 prepared liposomal suspension was then submitted to a probe sonication process with a 109 sequence of 40 s of sonication and 20 s of rest in an ice bath to the desired size. In all 110 111 cases, the initial turbid liposomal suspension was translucent after sonication. Then, the 6

112	sonicated liposomes were annealed at 4 $^{\circ}$ C overnight before centrifugation (12,000 rpm, 30
113	min) and diluted 4-fold to give a final lipid concentration of 25 mg/mL of lipid.
114	
115	2.2.1.2 Preparation of liposomes by proliposome method
116	1.5g of the appropriate hydration medium was added to a lipid dispersion containing
117	soya lecithin (750 mg), cholesterol (86.4 mg) and indometacin (30 mg) in ethanol (600 mg)
118	at 60 °C. The dispersion was stirred for 10 min at 60 °C, cooled to room temperature and
119	then converted to a 25 mg/mL liposome suspension by drop-wise addition of the rest of
120	hydration medium with continuous stirring for 2 hr following sonication and centrifugation.
121	
122	2.2.1.3 Spray drying <u>of liposomes</u>
123	Spray drying was performed with a Mini Spray-dryer (Büchi 190). Applied spraying
124	parameters were: inlet temperature 100 °C, outlet temperature 70 °C, air-flow 600-650 NI/h,
125	aspirator setting 20 (100%), pump setting 2.5- 3 ml/min. A 0.5-mm nozzle was used.
126	
127	2.2.2 Liposome size analysis
128	The average liposome size was determined with a ZetaSizer 3000HS (Malvern
129	Instruments Ltd, Malvern, United Kingdom) at a temperature of 25 $\pm$ 0.1 $^{\circ}$ C. Samples of the
130	dispersion were diluted with hydration medium and the Z-average vesicle size and

polydispersity were determined at 25  $^{\rm 0}{\rm C}$  by dynamic light scattering. The values of the  $^{\rm 7}$ 131

132 viscosity and refractive index used in the calculation of the liposome size of the light scattering data were modified in terms of protectant concentration. The intensity of the laser 133 light scattered by the samples was detected at an angle of 90° with a photomultiplier. For 134 each specimen 10 autocorrelation functions were analyzed using a Contin analysis. From 135 this analysis, the z-average diameter  $(D_z)$  was obtained, which is an approximation of the 136 diameter of the liposomes. The particle size distribution was characterized using the 137 polydispersity index (PI). The spray-dried liposomes were reconstituted with de-ionized 138 water to attain the original lipid content according to the method of Bligh and Dyer [46] and 139 then performing the Stewart assay [47]. Samples were further diluted with hydration 140 141 medium for liposome size analysis.

142

# 143 **2.2.3 HPLC analysis of indometacin**

HPLC was carried out using a Waters system (Waters 1525 Binary HPLC Pump, 144 Waters IN-Line Degasser AF, Waters 2487 Dual  $\lambda$  Absorbance Detector, Waters 717 plus 145 Autosampler). Luna C-18 column (100A<sup>0</sup>, 150 X 4.6 mm 5µm, Phenomenex) was used. 146 The detector wavelength was set at 260 nm. The mobile phase consisted of a gradient of 147 methanol and 0.1% v/v trifluoroacetic acid. The gradient schedule was: (a) 0-4.5 min, 85% 148 149 methanol, flow rate of 1ml/min; (b) 4.5-5.5 min,  $85 \rightarrow 100\%$  methanol, flow rate of 1ml/min; (c) 5.5-7 min, 100% methanol, flow rate of 1.2 ml/min; (d) 7-9 min, 100% methanol, flow 150 151 rate of 1.5 ml/min; (e) 9-15 min, 100% methanol, flow rate of 1.5 ml/min; (f) 15-16 min,

100% methanol, flow rate of 1 ml/min (g) 16-17 min,  $100 \rightarrow 85\%$  methanol, flow rate of 1.0 152 ml/min; (h) 17-23 min, 85% methanol, flow rate of 1.0 ml/min. In order to determine 153 encapsulation efficiency (EE %) 50 µL of separate liposome solutions and reconstituted 154 dispersions were applied to a Sephadex G-50 column and eluted with hydration medium. 155 The fractions were diluted with methanol and the concentrations of indometacin were 156 determined by HPLC. Loading efficiency was determined by dividing the encapsulated drug 157 content (µg) by the lipid content (mg). The encapsulation efficiency (EE (%)) of indometacin 158 was determined from the ratio of encapsulated to total drug concentration. 159

160

#### 161 **2.2.4 Water content and thermal analysis of the powders.**

Thermogravimetric studies were carried out to measure the water content of the 162 spray-dried liposomes using a TGA (Q500, TA instruments). Samples were heated from 20 163 to 25 °C at a scan rate of 10 °C /min. Modulated differential scanning calorimetry (MTDSC) 164 measurements of the dried products were performed on TA Q100 Differential Scanning 165 Calorimeter (Q100, TA Instruments, which had been calibrated for temperature, enthalpy 166 and heat capacity). The product was sealed into a hermetic aluminium pan and after 167 equilibration at 0 °C, was heated at 2 °C /min to 200 °C with a modulation of  $\pm$  0.4 °C/40 168 sec. Tg values are recorded as onset values from the reversing heat flow signal and 169 reported as mean values (n=4-6) with standard deviation. 170

171 **2.2.5 Scanning electron microscopy (SEM)** 

The spray-dried powders were coated with gold in a sputter coater and their surface morphology was observed using a scanning electron microscope (JEOL 6500F field emission scanning electron microscope).

## 175 **2.2.6 Powder particle size analysis**

The volume mean diameter of spray-dried liposomes was measured by Laser Light Diffraction Analyzer (HELOS/BR, Sympatec, Clausthal-Zellerfeld, Germany). Approximately 5 mg of powder were suspended in chloroform in a 50 ml glass cuvette and stirred with a magnetic bar at 1000 rpm. A short period of sonication (60 s) at a power of 60W (CUVETTE, Sympatec; 8.5 mm diameter ultrasound tip) was applied before sizing [48;

181 49]. A R4 lens was used allowing measurements in the range of 0.45-875  $\mu$ m.

#### 182 **2.2.7** *in vitro* release of indometacin from liposomes

Free drug was removed from the reconstituted liposome suspensions by use of centrifugal filter tubes (Amicon Ultra 15 MW Cut-off 10 KDa, Millipore). 1 ml portions were sealed into dialysis tube (MW cut-off 7 KDa, Thermo Scientific), and added to 50 mL of pH 7.4 PBS release medium in a shaking incubator (37° ±0.5 °C, 60 rpm, 25 mm throw; Unitron, Infors HT, Switzerland). 2-ml samples were drawn periodically and the amount of drug release determined using the HPLC method. The release volume was kept constant throughout.

# 190 2.2.8 Aerodynamic study

191 A twin-stage liquid impinger was used to determine the emitted dose and fine particle 10 192 fraction (FPF) of the spray-dried powders. The dried liposome powders were filled into number 3 gelatin capsules. A dry powder inhaler (Cyclohaler, Teva, UK) was attached to 193 the mouthpiece of emitted dose apparatus. Ten capsules were pierced and the dried 194 liposome powders were emptied at 60 L/min for 10 seconds. The powders deposited at 195 each stage were washed out and recovered. The powders deposited in the inhaler and 196 capsules were also collected. After being diluted with methanol to a suitable concentration, 197 each indometacin solution was assayed by HPLC. The FPF values were defined as the 198 powder mass recovered at stage 2. The results are expressed as the percentage of drug 199 dose emitted to the capsule content (loaded dose). 200

#### 201 2.2.9 Statistical analysis:

<u>Kruskal-Wallis analysis with Dunn's all pairwise multiple comparisons or Mann-Whitney</u>
 <u>U test were used</u> to calculate the p values using SigmaPlot 8 software. Differences were
 deemed significant if p values were <0.05.</li>

205

## 206 3. Results and discussion

#### **3.1Initial Determination of Optimum Disaccharide and L-leucine concentrations**

# 208 3.1.1 Effect of disaccharide concentration on liposomal size and PI

- 209 The size change of any liposomes prior to and after spray drying is a critical parameter
- in the assessment of liposomal stability so this was used as an initial screening parameter
- in choosing which formulations to take forward to the next step of formulation optimisation.

Initially the optimum concentration of either sucrose or trehalose as protectants during thespray drying process was determined.

214 <Figure 1>

As shown in Figure 1a, the addition of 2.5% (w/w) sucrose to the hydration medium did 215 not prevent the aggregation and fusion of reconstituted liposomes after spray drying, but 216 increasing the concentration to 5% (w/w) inhibited liposomal size increase prior to spray 217 drying and after reconstitution more effectively. A further increase to 7.5% (w/w) showed no 218 significant change in the PI values. 10% (w/w) Sucrose was more protective still; this was 219 the only formulation containing disaccharide alone that showed no statistical difference in 220 221 liposomal size after drying. A further increase in concentration to 15% (w/w) resulted in an increase in liposome size. This effect can also be seen by comparing the  $S_F/S_I$  ratios, 222 where is  $S_F$  is the final liposomal size after rehydration and  $S_I$  is the initial liposomal size 223 [50]. Previous research into freeze-drying of liposomes demonstrated that solute 224 incorporated by liposome in the presence of lyoprotectant in a mass ratio of sugar: lipid of 225 2:1 could be effectively retained after freeze-drying [51], in the 5% (w/w) sucrose 226 dispersions the mass ratio of sucrose to lipid is close to this ratio. A similar effect was 227 observed when trehalose was used as the protectant (Figure 1b). However, the protective 228 effect of trehalose was not so good; the addition of 15% (w/w) trehalose dihydrate to the 229 hydration medium did not prevent the liposome size increasing after reconstitution (p < p230 231 0.05).

#### 232 **3.1.2 Effect of disaccharide concentration on recovery rate and water content of**

#### 233 spray-dried liposome powders

The recovery rate of the spray-dried powders significantly improved when increasing the 234 concentration of protectants from 2.5% (w/w) to 5% (w/w) for both sugars (Figure 2), 235 although further addition of either disaccharide showed little additional effect. Further, the 236 237 spray-dried liposomes with trehalose had a much higher recovery rate than those with sucrose, which is ascribed to its higher Tg (101 °C compared with 64 °C for sucrose [36]. 238 The sticky point (*Ts*) of an amorphous powder is generally considered to lie 10-20 °C above 239 Tg; if the outlet temperature of the dryer surpasses *Ts* then particle cohesion increases 240 sharply and there may also be increased adhesion to the dryer walls [35]. Because the 241 powder temperature would have been greater than the Tg for the sucrose formulation, the 242 particles would have been more prone to adherence to the walls of the spray-dryer. 243 Increasing the concentration of either protectant reduced the water content of the 244 spray-dried powders (Figure 2). Since Tg data are inversely related to water content this 245 would be expected. Based on the results in 3.1.1 and 3.1.2 the formulations containing 10% 246 sucrose and 15% trehalose were selected for further optimisation by addition of the 247 anti-adherent L-leucine. 248

249 <Figure 2>

250 <Figure 3>

251 **<u>3.2 Effect of varying L-leucine concentration</u>** 

#### 252 **<u>3.2.1 Effect of L-leucine concentration on liposomal size</u>**

Figure 3a indicates that the inclusion of L-Leucine at a concentration of 0.5% (w/w) 253 was the most effective in preventing size changes for both the 10% (w/w) sucrose and the 254 15% (w/w) trehalose formulations, with no statistical difference seen in the sizes prior to 255 spray drying compared with the reconstituted dispersions. Increasing the concentration to 256 257 1% (w/w) L-Leucine had the effect of markedly increasing the liposomal size in the reconstituted dispersions. This might be caused by partitioning of the hydrophobic amino 258 acid into the lipid membrane during drying, causing vesicle fusion. While this phenomenon 259 has not been investigated for spray dried liposomes it has been observed previously during 260 261 freezing and freeze-drying of liposomes; Anchordoguy et al [52] found that the amino acids 262 with hydrocarbon side chains increased membrane damage during freeze/thaw trials. Popova et al [53] observed that the amphiphilic aromatic amino acids tryptophan and 263 phenylalanine induced solute leakage and membrane fusion during freezing studies of 264 liposomes, while Mohammed [54] et al demonstrated that basic, polar amino acids 265 stabilized liposomes during lyophilisation but that at higher concentrations these amino 266 acids promoted vesicle fusion. 267

268 <Figure 4>

#### 269 <u>3.2.2 Effect</u> of L-leucine concentration on recovery rate and water content

Addition of 0.5% (w/w) L-Leucine markedly improved the recovery rate of the 10% (w/w)
 sucrose formulations (Figure 4) but had little effect on the liposomes formulated with 15%

272 (w/w) trehalose, which exhibited good yields already. L-Leucine did not have a significant effect on the water contents of the powders (Figure 4) but a marked improvement in the 273 appearance of the powders was apparent to the naked eye. Formulations without L-leucine 274 were clumped into aggregates in the mm size range while increasing L-leucine reduced the 275 size. This effect was especially apparent for the sucrose formulations. 276

Taking all data from 3.1 and 3.2 into account the two formulations selected for further 277 testing were those containing 0.5% (w/w) leucine and either 10% (w/w) sucrose or 15% 278 (w/w) trehalose. 279

280

#### 3.3 Effect of Liposome manufacturing method on liposomal size and drug content 281

282 Once the optimal levels of disaccharide and anti-adherent had been determined, the effects of manufacturing process variables were investigated. Table 1 shows that liposomal 283 size was clearly larger for the formulations prepared by the proliposome method. It has 284 been suggested that the proliposome-prepared liposomes produce multilamellar vesicles 285 [31], while those prepared by the ethanol-injection method are primarily unilamellar, which 286 are more amenable to size reduction under the same sonication conditions. Based on 287 encapsulation efficiency and loading efficiency, the two methods have the same capability 288 to incorporate indometacin. However, lipid loss during the alcohol injection method could 289 not be avoided owing to this process involving injection of lipid solution into hydration 290 291 medium. It has previously been reported that ethanol concentration is a decisive factor in 15

292 liposome size reduction using a high-pressure homogenizer, whereby the liposomal size293 and range decreased with increasing concentration of ethanol [55].

294 Statistically significant changes were seen between freshly prepared and reconstituted liposomes prepared by the ethanol injection in terms of PI value, encapsulated content and 295 loading efficiency. The small increases in encapsulated content and loading efficiency were 296 attributed to unentrapped drug being reincorporated into the liposomes during 297 reconstitution, as has previously been demonstrated for liposomes prepared using the DRV 298 (dried-rehydrated vesicle) method [56]. Even though the formulation was optimised using 299 the ethanol injection method, excellent encapsulation efficiency in liposomes prepared by 300 301 the proliposome method was obtained. Dispersions produced using the proliposome method showed small statistically significant reductions in PI values for both formulations, 302 while for the trehalose formulation there was also a small statistically significant reduction in 303 liposomal size after drying. However, the liposomal indometacin content and its loading 304 efficiency did not significantly change during spray drying. In short, while sucrose enabled a 305 better loading efficiency when comparing samples produced by the ethanol injection 306 method; the combination of either 10% (w/w) sucrose and 0.5% (w/w) leucine or 15% (w/w) 307 trehalose and 0.5% (w/w) leucine in the formulations could effectively protect liposomes 308 prepared by either method against spray drying stress. In terms of drug loading and 309 efficiency liposomes prepared by the proliposome method incorporated drug more 310 311 effectively than those produced using the ethanol-injection method (Table 1). In addition, a 16

dilution effect was observed when comparing loading efficiency of liposomal stock solutions and the final dispersions prepared by the ethanol-injection method, which was attributed to encapsulated drug leaking out of the liposome stock solutions upon dilution as observed by. Foldvari et al [57]. TEM images (data not shown) indicate that the vesicular structure of liposome in the presence of the optimised disaccharide and anti-adherent formulations could be preserved very well through spray drying .

318

319 <Figure 5>

#### 320 **<u>3.4 Imaging</u>**, size analysis and aerodynamic properties of the spray-dried powders

SEM analysis showed that particles of the optimally formulated sucrose powder 321 exhibited a smooth surface (Figure 5a and b), while the powders containing trehalose were 322 wrinkled (Figure 5d, e and f), which potentially prevents particles from adhering tightly to 323 324 each other, thus preventing aggregation and lowering the energy required to disperse them [58]. This may explain why the sucrose formulation appeared to contain more aggregates. 325 Furthermore, the fine particle fraction was higher for the trehalose/L-leucine formulation 326 than for that containing sucrose/L-Leucine (Figure 6). 327 <Figure 6> 328

Bosquillon et al [59] had reported that the type of sugar incorporated did not affect morphology of the spray-dried powders. The wrinkled surface perhaps can be attributed to the inclusion of additives that alter the surface tension that controls droplet shape during 17 332 drying as Adler et al [60] demonstrated that surfactant could change the interface viscosity and that consequently, the dried particle morphology also changed. By increasing the ratio 333 of surfactant to other additives, more spherical, smooth particles were obtained. It is 334 proposed that L-leucine at a concentration of 0.5% (w/w) functions as a surfactant. Hence, 335 it is suggested that the ratio of 0.5% (w/w) leucine to 15% (w/w) trehalose is below the 336 critical point of powder morphology conversion between roughness and smoothness and 337 so it tends to exhibit a wrinkled surface. The formulation containing 10%(w/w) sucrose 338 and 0.5% (w/w) exhibited a smoother morphology. To investigate whether this was a 339 concentration effect rather than being specific for trehalose, spray dried liposomes were 340 prepared with a sucrose concentration of 15% (w/w), but these still had smooth surfaces 341 342 (Figure 5c and d), indicating that the wrinkling is not caused by surface tension effects alone. The SEM images also show that the diameters of all the spray dried powders were 343 less than 10µm (Figure 5). Accurate size distribution data are given in Table 2, which show 344 that all formulations the volume mean diameters were measured to be 3~4 µm with the 345 exception of the formulation containing 10% (w/w) sucrose and 0.5% (w/w) L-leucine, 346 prepared by ethanol injection, which gave a larger diameter of 5.40 µm. This size increase 347 cannot be attributed to hygroscopicity of the sucrose formulations [34] as the water content 348 values for all formulations were similar. Although the sucrose formulations exhibited lower 349 350 Tg values than the trehalose formulations, as might be expected, there was no significant 351 difference between those obtained for the liposomes prepared by the ethanol injection 18

rather than the proliposome method. The emitted dose and fine particle fraction of spray-dried liposome formulated with trehalose and L-leucine were higher than those formulated with 10% (w/w) sucrose and 0.5% (w/w) L-leucine. (Figure 6). Therefore, spray dried liposomes with 15% (w/w) trehalose and 0.5% (w/w) leucine exhibited better aerosol powder performance than those in the presence of 10% (w/w) sucrose and 0.5% (w/w) leucine in terms of emitted dose, aerodynamic diameter and fine particle fraction.

358 <Figure 7>

#### 359 **<u>3.5 Release</u> of indometacin from reconstituted liposomes**

Release data for Indometacin from reconstituted liposomes for both optimised 360 formulations using both methods of manufacture are shown in Figure 7. Diffusion of free 361 362 indometacin through the dialysis membrane was measured as a control. The release profile of free drug shows over 80% diffusion in 4 hr. Over the same interval, the reconstituted 363 liposomes prepared by the ethanol-injection method released 15.1 % and 16.9% drug 364 (optimised formulations containing sucrose & trehalose respectively) and released ~45% of 365 the encapsulated drug over 24 hr. Those prepared by the proliposome method in the 366 presence of disaccharide (sucrose or trehalose) and leucine exhibited drug release of 367 21.8% and 24.3% in 4 hr and in total released 54.5% and 61.5% over 24 hours respectively 368 (Figure 7). These differences are not significant suggesting that the mode of release is the 369 same for all formulations. The mechanism responsible for the release of drug from the 370 371 liposomes may be due to diffusion phenomena, degradation effects, or a combination of 19

both processes. To examine the drug release kinetics and mechanism, the release data were fitted to models representing zero-order, first-order, and Higuchi's square root of time and the Korsmeyer–Peppas models (Table 3). All systems showed best correlation with the Higuchi model and anomalous (non-Fickian) diffusion (n > 0.5). It is reasonable to propose that this is because the indometacin is located within the phospholipid membrane of the liposomes and must diffuse through in order to be released.

378

#### 379 **4. Conclusions**

The inclusion of either 10% (w/w) sucrose or 15% (w/w) trehalose dihydrate and 0.5% (w/w) L-leucine protected liposomes prepared by ethanol injection or proliposome method against spray drying stress in terms of size change, polydispersity index, encapsulated drug content and loading efficiency of the reconstituted liposomes. Formulation design was of more importance than the method of liposome manufacture. The method for preparing the liposomes had no effect on the stability or encapsulation efficiency of spray-dried liposomes with optimal protectant and anti-adherent.

This paper has demonstrated for the first time that L-leucine should be used with care as an additive for spray dried liposomes in combination with disaccharide. While there was a clear advantage in using this material at an optimum level; at higher concentrations it caused an increase in liposomal size upon rehydration of the spray dried powders that might be attributed to it partitioning into the lipid membrane during drying, causing vesicle  $_{20}$  392 fusion. This is an area for future research.

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**Table 1.** The characterization of liposomes prepared by ethanol injection and by proliposome. The hydration buffers were in either a combination of 10% (w/w) sucrose and 0.5% (w/w) L-Leucine or 15% (w/w) trehalose dihydrate and 0.5% (w/w) L-Leucine. \* denotes P<0.05 (Mann-Whitney U test) in comparison with the corresponding samples prior to spray-drying. + P<0.05 (Mann-Whitney U test) compared to formulations with the inclusion of trehalose/L-Leucine. Each value represents the mean  $\pm$  SD (three different batches) and n denotes replicate measurements of each batch.

Preparative methods	Ethanol injection		Proliposome				
	Before spray-drying	After reconstitution	Before spray-drying	After reconstitution			
Liposome with 10% (w/w) sucrose and 0.5% (w/w) L-Leucine							
Liposome size (nm) & (Pl)	$107.4 \pm 13.8$ (0.20 ± 0.02)	$115.1 \pm 11.5^{*}$ (0.25 ± 0.04*)	$137.9 \pm 4.9$ (0.48 ± 0.02)	$134.0 \pm 8.2$ (0.39 ± 0.03*)			
Total drug content (μg/mL)	407.5 ± 7.9	384.8±7.3*	846.4 ± 23.5	797.2 ± 13.1*			
Encapsulation efficiency (%)	33.3±6.1	53.2 ± 15.1*	45.4 ± 1.0	45.9 ± 2.3			
Encapsulated drug content (µg/mL)	$135.6 \pm 24.9$	204.8±58.1*	397.6±8.3	366.3 ± 23.7			
Lipid content (mg/mL)	23.57 ± 0.37	21.60 ± 1.17*	26.75 ± 0.47	25.87 ± 0.26			
Loading efficiency (μg drug/ mg lipid)	5.7 ± 1.0	9.5 ± 2.9* <sup>+</sup>	14.4±0.6	14.3±1.8			

#### Liposome with 15% (w/w) trehalose dihydrate and 0.5% (w/w) L-Leucine

Liposome size (nm) &	$130.7 \pm 2.7$ (0.31 ± 0.03)	132.7 ± 4.2 (0.36 ± 0.02*)	$138.5 \pm 4.8$ (0.52 ± 0.04)	$127.3 \pm 3.6^{*}$ (0.40 ± 0.03*)
Total drug content (μg/mL)	442.1 ± 7.4	428.2 ± 10.9*	$899.9 \pm 8.1$	868.6±16.7*
Encapsulation efficiency (%)	37.4 ± 6.4	42.5 ± 3.8	49.7 ± 7.1	52.0 ± 7.0
Encapsulated drug content (µg/mL)	165.3 ± 27.4	181.8 ± 16.5	447.5 ± 67.7	449.1 ± 66.0
Lipid content (mg/mL)	$26.21 \pm 0.70$	25.51 ± 1.80	$29.60 \pm 0.10$	27.85 ± 0.39
Loading efficiency (μg drug/ mg lipid)	6.3±0.9	7.2 ± 0.70	15.1 ± 2.2	15.7 ± 2.0