1 Optimised assembly of DNA-lipid nanostructures

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16 Abstract

17

18 Liposomes, aqueous vesicles enclosed by lipid bilayers, are widely used in research as simple, synthetic analogues of cell membranes. Membrane-binding DNA nanostructures have been 19 developd whichh can modify the shape, porosity and reactivity of liposomes. Lipid-DNA 20 21 binding is moderated using strands with hydrophobic or amphipathic chemical groups such as cholesterol. However, the factors that affect the binding interactions of cholesterol-modified 22 23 DNA and membrane bilayers have not been systematically investigated. Here we characterise 24 the effect of buffer and lipid composition and DNA structure near the cholesterol motif on the 25 strength of DNA-lipid binding. We observed that DNA-membrane binding is inhibited at increasing ionic concentrations and that binding is severely reduced in strongly acidic 26 27 conditions. Background membrane cholesterol content demonstrated a more varied effect, dependent on lipid composition. The composition of the DNA, whether simplex or duplex, 28 29 showed little effect on binding, as did the presence or absence of a single-stranded 'overhang' to protect the cholesterol and prevent DNA strand aggregation. Our results inform the design 30 31 and modelling of the membrane binding of cholesterolated DNA nanostructures.

32 Introduction

33

34 Liposomes are aqueous vesicles bound by one or more bilayers of lipids, a diverse group of amphipathic and hydrophobic small molecules. Due to their similarities to membrane bilayers, 35 36 which are ubiquitous in nature and form the basis of biological compartmentalisation, liposomes have proven a powerful research tool for modelling cellular membranes in simplified 37 38 synthetic systems. Liposomes can also be used for encapsulating therapeutic payloads in order 39 to increase a drug's circulation time and alter its distribution profile (Storm and Crommelin, 40 1998). Various strategies for engineering liposomes for therapeutic applications have been 41 developed in order to increase circulation time, allow targeted payload release or deliver a payload to a cell's cytosol (Dou et al., 2017; Veronese and Harris, 2002). A number of 42

43 liposome-encapsulated 'nanodrugs' are FDA-approved (Bulbake et al., 2017).

DNA nanotechnology is a 'bottom-up' approach to designing and building nanometre-scale
structures based on DNA based on Watson-Crick base pairing (Seeman and Sleiman, 2018).
Since the development of DNA nanotechnology (Seeman, 1982), a variety of two and three-

47 dimensional structures have been created and described (Wang et al., 2017) as well as

48 environment-sensing mechanisms which allow DNA nanostructures to change state in response

49 to an external trigger (Singh et al., 2018).

50 DNA and lipid nanotechnologies can be combined by modifying DNA with hydrophobic chemical groups such as cholesterol to enable membrane binding (Bell and Keyser, 2014). 51 52 Using this approach, a variety of membrane-binding and membrane-spanning DNA 53 nanostructures have been developed (Darley et al., 2019). Membrane-binding DNA nanostructures have been used to functionalise liposome surfaces (Akbari et al., 2017), control 54 55 the shape of liposomes by inducing membrane curvature and tubulation (Franquelim et al., 56 2018; Grome et al., 2018) and form membrane-spanning nanopores (Burns et al., 2013; 57 Langecker et al., 2012). Such nanpores can have dimensions which exceed those of natural 58 protein pores (Diederichs et al., 2019) and feature gating mechanisms that can be triggered 59 externally (Burns et al., 2016; Mendoza et al., 2017).

60

61 Despite the widespread use of cholesterol for DNA-lipid mediation, little is known about the kinetics and energetics of DNA nanostructure insertion in bilayers (Darley et al., 2019). 62 63 Increasing the efficiency of membrane attachment is thus of great interest to the design and 64 application of membrane-bound DNA nanostructures. In particular, currently large numbers of 65 hydrophobic groups are necessary for spontaneous and stable membrane insertion to occur (Krishnan et al., 2016), and to overcome the substantial energy penalties associated with the 66 67 insertion of membrane-spanning DNA nanopores (Göpfrich et al., 2016). It has been observed 68 that both the quantity and position of TEG-cholesterol anchors on DNA nanostructures affects 69 their affinity for lipid bilayers (Khmelinskaia et al., 2016; Langecker et al., 2012). Monovalent 70 and divalent cations are necessary buffer components in order to assemble and maintain the 71 stability of DNA duplexes and nanostructures (Kielar et al., 2018; Nakano et al., 1999), yet 72 also are also known to affect the physical characteristics of membrane bilayers (Böckmann et 73 al., 2003; Velikonja et al., 2013) and may affect the binding activity of cholesterol-modified

- 74 DNA. Thus far, the optimal environmental conditions to promote binding interactions between
- cholesterol-modified DNA and have not been systematically investigated.

76 Here we have quantified the binding of cholesterol-modified DNA strands to synthetic

77 liposomes using fluorescence microscopy. We examined the effects of pH, ion concentration

- 78 and membrane cholesterol content on the binding of cholesterol-modified DNA strands to
- 81 Ohmann et al. to reduce aggregation during nanostructure assembly (Ohmann et al., 2019).

82 Materials and Methods

- 83 Preparation of Buffers and solutions
- 84 In order to investigate the effect of salts on DNA/Lipid interaction, Liposomes and DNA stocks

85 were diluted in Liposome Buffer (210 mM D-Sorbitol [S1876, Sigma], 5 mM Tris-HCl

86 [**T3253**, **Sigma**], pH 7.5) containing NaCl [AJA465, Ajax-Finechem] (12.5 mM to 400 mM)

- and $MgCl_2$ [AJA296, Ajax-Finechem] (0 mM to 80 mM) as required.
- 88 To investigate the effect of pH on DNA/Lipid interaction we used a modified Liposome Buffer
- 89 (210 mM D-Sorbitol, 100 mM NaCl) with pH adjusted to approximate pH values of 2, 4, 6, 7,
- 90 8 and 10. The pH was adjusted to ± 0.2 of the target pH value by using 200 mM NaOH or
- 91 HCl 3 hours prior to imaging.
- 92

Buffer	Composition
Slide buffer	100 mM NaCl, 10 mM Tris-HCl, pH 7.5
DNA duplex buffer	100 mM NaCl, 5 mM Tris-HCl, pH 7.5
Electroformation buffer	210 mM sorbitol, pH 7.5
Extrusion buffer [Standard]	210 mM sorbitol, 100 mM NaCl, 5 mM Tris-
	HCl, pH 7.5
Extrusion buffer [NaCl]	210 mM sorbitol, X mM NaCl, 5 mM Tris-HCl,
	pH 7.5
Extrusion buffer [MgCl ₂]	210 mM sorbitol, 100 mM NaCl, X mM MgCl ₂ ,
	5 mM Tris-HCl, pH 7.5
Extrusion buffer [pH]	210 mM sorbitol, 100 mM NaCl, pH X

93

94 Design and assembly of oligonucleotides and DNA duplexes

DNA strands used for colocalisation experiments were 23 nt-long, used alone (ssDNA), or hybridised to a complimentary oligo (dsDNA) or a complimentary sequence with a 5' 6 nt single stranded 'overhang' (dsDNA-6 nt) (Supplementary Table 1). The oligonucleotide sequences were generated using NUPACK design software (Zadeh et al., 2011) and selected

99 to prevent the formation of unwanted secondary structures. The 6 nt 'overhang' sequence 100 introduced at the 5' end of oligos was chosen from Ohmann, et al., 2019. Oligos were modified

101 at the 3' end with a tetraethylene glycol cholesterol moiety (TEG-cholesterol) and with a 5'

102 Alexa 647 fluorescent group respectively. All oligos were purchased from IDT (Integrated

103 DNA Technologies, Inc., USA).

104 DNA stocks (100 µM, 1000x) were prepared using MilliO water [Milli-O, Millipore] and

105 stored at 4° C. Alexa 647-labelled DNA was stored in foil at -20° C. DNA duplexes were

- annealed at 10 μ M final concentration in duplex buffer. All oligos were heated to 90°C for five
- 107 minutes then cooled in a termocycler at 5° C /minute for 15 minutes to a final temperature of
- 108 15°C, then stored at 4°C. For duplex assembly, unmodified complementary strands were added
- in a 3-fold excess. DNA was diluted in extrusion buffer to a final concentration of 100 nM after
- 110 the melting and annealing steps.

111 Preparation of Liposomes

112 Liposomes were produced from two main lipid mixtures, DOPE/DOPC liposomes [49.9% 1-

113 palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (DOPE 18:1, 850725 P, Avanti),

114 49.9% 1-palmitoyl-2-oleoyl-glycero-3-phosphocholine (DOPC 18:1, 850375 P, Avanti)] or

115 DPhPC liposomes [99.8% 1,2-diphytanoyl-sn-glycero-3-phosphocholine (DPhPC 850356P

116 Avanti Polar Lipids)] (Supplementary Table 2). Both lipid mixtures were doped with 0.1%

117 PE-rhodamine [1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-lissamine rhodamine B

sulfonyl, 810150P Avanti Polar Lipids] for fluorescence imaging and 0.1% PE-biotin [1,2-

119 dioleoyl-sn-glycero-3-phosphoethanolamine-N-biotinyl, 870282P Avanti Polar Lipids] for

120 surface tethering. All percentages indicate weight to weight ratios.

121 Liposomes with cholesterol were prepared by replacing either DPhPC or equal parts of DOPE

and DOPC with cholesterol [700000P Avanti Polar Lipids]. All lipids stocks were dissolved in
chloroform at 10 mg/mL and stored at -20 °C.

124 Large unilamellar vesicles (LUVs) were produced by extrusion using a Mini-Extruder kit

125 (Avanti Polar Lipids Inc., USA). Briefly, lipid stocks were added to a round-bottom glass tube

126 and dried into under gentle nitrogen flow into a thin film and resuspended in extrusion buffer

127 to a final concentration of 1 mg/mL by vortex mixing and sonication. The resulting suspension

128 $\,$ was transferred to a 500 μL glass syringe (Hamilton Company, UK) and passed back and forth

129 through a 100 nm polycarbonate filter (Whatman plc, USA) 41 times to produce a clear

130 suspension of homogenous unilamellar liposomes. Liposomes were then diluted 100-fold in

131 Liposome Buffer solution prior to loading onto tunnel slides for imaging.

132 Giant unilamellar vesicles (GUVs) were produced by electroformation using the Vesicle Prep

133 Pro machine (Nanion Technologies GmbH, Germany). 30 μ l of 3.5 mg/mL lipid dissolved in

134 chloroform was added to a conductive indium tin oxide-coated glass slide and spread over a 135 spot approximately 12 mm in diameter and allowed to air-dry for two minutes into a circular

film. A 1.5 mm thick rubber gasket of 15 mm diameter was placed around the film, forming a

137 well into which 250 µl of electroformation solution was added. A second indium tin oxide-

138 coated glass slide was placed face-down on top of the gasket and clamped in place, creating a

139 sealed chamber of liquid between the two slides. The machine was run using the default

140 protocol of 3 V AC for 120 minutes to form GUVs. Giant unilamellar liposomes in

141 electroformation solution were then diluted at a 1:1 ratio in buffer consisting of 210 mM

142 sorbitol, 80 mM NaCl, 10 mM Tris-HCl, giving a final external solution of 210 mM sorbitol,

- 40 mM NaCl, 5 mM Tris-HCl. Liposome dissolution was tested by titration of increasing
 concentration of the detergent Polysorbate-20 (Supplementary Figure 4).
- 145 Construction of tunnel slides for microscopy

146 To form a tunnel slide for imaging DNA-liposome interactions, a 50 mm cover slip (#1 thickness) (Menzel Glaser GmbH) was fixed to a glass slide (Suzhou Upline Medical Products 147 Co., China (PRC)) using two parallel strips of double-sided tape (Nichiban Co., Japan) 148 149 approximately 150 µm thick placed 2 mm apart, forming a channel of approximately 15 µL 150 volume. A thin layer of CoverGrip Coverslip Sealant (Biotium Inc., USA) was applied over 151 remaining exposed tape to prevent the contamination of solutions by adhesive residue and left 152 to cure for 24 hours. Solutions were added to one end of the channel with a pipette while simultaneously drawing solution from the opposite end with an absorbent paper wipe 153 154 (Kimberly-Clark Professional, USA).

155 Tethering of liposomes for TIRF imaging

156 The imaging system was based on a protocol developed by Jungmann, et al. for DNA-PAINT

- 157 super-resolution microscopy of cells (Jungmann et al., 2014), with modifications made to 158 buffer compositions, volumes and solution concentrations.
- 159 First, 15 µL of a 9:1 mixture of bovine serum albumin (BSA) and biotinylated bovine serum
- 160 albumin (BSA-biotin) in slide buffer at a combined concentration of 1 mg/mL was added to
- 161 the channel and incubated for 10 minutes to block and coat the surface of the cover slip. Excess
- 162 BSA and BSA-biotin in solution was then removed by flushing 60 μ L of slide buffer through
- 163 the channel.
- 164 Next, 15 μ L of streptavidin at 0.1 mg/mL in buffer A was added to slide and incubated for 10

165 minutes. Unbound streptavidin remaining in solution was removed from the slide by flushing

166 60 μ L of extrusion buffer through the slide. Afterwards, 15 μ L of biotinylated liposome

- 167 solution was introduced into the slide and incubated for 30 minutes to allow streptavidin-biotin
- 168 conjugation. Finally, 15 μ L of Alexa 647-labelled DNA solution (100 nM) in extrusion buffer
- 169 was added to the slide and incubated for 30 minutes prior to imaging.
- 170 Fluorescence microscopy of extruded liposomes (Large Unilamellar Vesicles)
- 171 Surface-tethered liposomes were imaged using on a Zeiss Elyra PALM/SIM Microscope in
- 172 Total Internal Reflection Fluorescence (TIRF) mode with a 63x/1.4 Oil Iris M27 oil immersion
- 173 objective (Carl Zeiss AG, Germany) and Andor iXon 897 EMCCD camera (Oxford
- 174 Instruments, United Kingdom).
- 175 Two-channel images were collected to visualize the fluorescence from liposomes ('liposome
- 176 channel', 561 nm laser) and fluoreophore-tagged DNA ('DNA channel', 642 nm). Signal in the
- 177 'liposome channel' was imaged using an emission dichroic filter (570-650 nm band pass plus
- 178 750 nm long pass) with a camera integration time of 100 ms and line averaging of two. The
- 179 'DNA channel' was imaged using an emission dichroic filter (655 nm long pass) with a camera
- 180 integration time of 33 ms and line averaging of two.

181 Fluorescence microscopy of electroformed liposomes (Giant Unilamellar Vesicles)

182 Binding interactions between DNA fluorophores and the surface of micron-scale GUVs were

imaged using a Leica TCS SP8 DLS confocal microscope with a HC PL APO CS2 63 x oil

184 immersion objective lens and Acousto-Optical Beam Splitter, a programmable crystal-based

185 beam splitter (Leica Microsystems GmbH, Germany). Two-channel images were acquired

186 showing Liposomes and DNA. In channel one, Rhodamine B-labelled liposomes were excited

- 187 with a 561 nm laser and imaged between 569-611 nm. In channel two, DNA labelled with was
- 188 excited with a 640 nm laser and imaged between 690-734 nm.

189 *Quantifying DNA-liposome colocalisation*

190 A custom macro script was developed using FIJI in ImageJ (Schindelin et al., 2012) to quantify 191 the colocalisation of DNA and liposomes using a method inspired by Manders Overlap 192 Coefficient (Dunn et al., 2011). Briefly, the pixel intensity data for the 'liposome channel' of 193 all images within a dataset comprised of two-channel images (representative of a single 194 experimental condition) was aggregated and analysed to determine the pixel intensity threshold 195 used to define the boundary of liposomes against the slide background (Supplementary Methods, Supplementary Fig. 1). A unique binary mask of the liposome channel was then 196 197 generated based on this threshold for each image in the dataset to separate liposome-covered 198 section from the background. The method used showed no bias or correlation with liposome 199 area (percentage coverage), in comparison with Pearson's correlation, whichh did 200 (Supplementary Fig. 2/3).

201 The mean pixel intensities of the DNA channel for each of these two sections was then

202 compared to produce a ratiometric colocalisation score, *C*, indicating the relative fluorescent

203 intesity of DNA attached to liposomes (within the binary mask) compared to the fluorescence

204 of DNA in the background of the image (outside the binary mask). For example, a

205 colocalisation score of C = I is indicative of an image where DNA fluorescence is evenly

- 206 distributed across both sections and therefore displays an equal mean pixel intensity in both 207 channels.
- 207 channels.

208 **Results**

209 Characterisation of DNA-liposome interactions through colocalisation analysis

210 We immobilised a 100-fold dilution of extruded liposomes on the surface of a coverslip using

- 211 biotin-avidin conjugation (Figure 1). Fluorescent DNA colocalised with fluorescent liposomes
- only when cholesterol tags were present (Figure 1e-h). DNA without a cholesterol tag did not
- colocalise with liposomes and was distributed evenly throughout the image independently of
- 214 the position of liposomes (Figure 1a-d). The role of cholesterol tags in causing DNA-liposome
- 215 colocalisation was further verified by confocal images of cholesterol-tagged DNA colocalising
- 216 with giant unilamellar vesicles, while DNA with no cholesterol tag did not colocalise with
- 217 liposomes (Figure 1d/1h)

We quantified the colocalisation of DNA to extruded liposomes by determining a colocalisation score. The liposome channel was first converted into a binary image according to a standardised pixel intensity threshold (SI). This binary image of the liposome channel was

then used as a mask to divide the DNA channel into two sections: liposome and background,which were used to calculate a colocalisation score via:

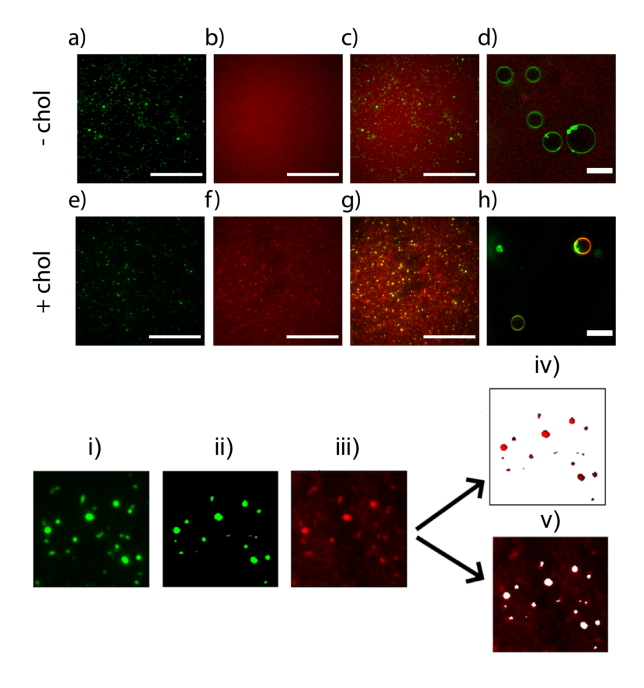
223

224
$$C = \frac{D}{B}$$

225

Where C is the reported colocalisation score, D is the mean pixel intensity of the fluorescent DNA in the liposome region of the DNA channel, and B is the mean pixel intensity of the

fluorescent DNA in the background region of the DNA channel (Figure 1).



- 230
- 231

Figure 1: Image acquisition and analysis of DNA-liposome interactions with fluorescence microscopy. Top row – without cholesterol: DNA (a) rhodamine-doped liposomes, (b)

234 *Alexa647-DNA (c) merge. (d) GUVs + Alexa647-DNA visualised using confocal fluorescence.*

235 Second row – with cholesterol (e) rhodamine-doped liposomes, (f) Alexa647-DNA, (g) merge,

236 (h) GUVs + Alexa647 DNA. Bottom row: Image analysis process for quantifying DNA-

liposome colocalization: the image of rhodamine-doped liposomes (i) is converted into a
binary mask (ii). This mask is then used to partition the Alexa647-DNA image (iii) into two

- sections: liposomes (iv) and background (v). The mean pixel intensity of DNA in the liposome
- section (iv) is divided by the mean pixel intensity of DNA in the background section (v) to give
- 241 *a ratiometric colocalisation score*. Scale bars: 20 μm.
- 242

243 The effect of solution composition on DNA-liposome colocalisation

244 We investigated the effect of common DNA origami buffer components on the interactions of

cholesterol-tagged DNA and lipid bilayers by quantifying the colocalisation of DNA and

unilamellar liposomes in various conditions. For each condition, the co-localisation of ssDNA,

dsDNA and dsDNA-6 nt was measured in order to compare the overall binding yield for each
 DNA configuration. Tests were repeated for 1:1 DOPE/DOPC liposomes and DPhPC

249 liposomes to detect for lipid-dependant responses to variables.

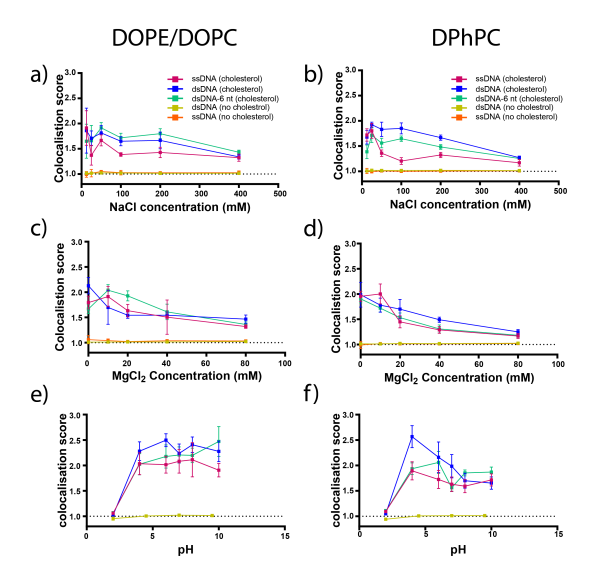
- 250 Imaging DNA-liposome interactions in extrusion buffer [NaCl] containing between 12.5 and
- 400 mM NaCl and extrusion buffer [MgCl₂] containing between 0 and 80 mM MgCl₂ revealed
- that both salts had a significant impact on DNA-liposome colocalisation. The colocalisation of
- 253 cholesterol-tagged DNA and liposomes was inhibited at increasing concentrations of NaCl
- 254 (Figure 2 A & B) and MgCl₂ (Figure 2 C & D). For all three configurations of cholesterol-
- tagged DNA on both DOPE/DOPC liposomes and DPhPC liposomes, a significant decrease
- 256 was observed in colocalisation scores between 12.5 mM and 400 mM NaCl (P < 0.05) and
- between 0 mM and 80 mM MgCl₂ (P < 0.05). Linear regression analysis for all three configurations on both 1:1 DOPE/DOPC liposomes and DPhPC liposomes showed a significant (P < 0.05) trend of decreasing co-localisation scores with increasing NaCl and
- 260 MgCl₂ concentration.
- 261 Imaging DNA-liposome interactions in extrusion buffer adjusted to approximate pH values of
- 2, 4, 6, 7, 8 and 10 showed that the colocalisation of cholesterol-tagged DNA and liposome 262 was inhibited in highly acidic conditions. At pH 2, the colocalisation of all three configurations 263 264 of cholesterol-tagged DNA with both DOPE/DOPC liposomes (Figure 2e) and DPhPC liposomes (Figure 2f) was strongly inhibited and significantly less than at all other pH values 265 266 (P < 0.05). For DOPE/DOPC liposomes, all configurations of cholesterol-tagged DNA produced similar colocalisation scores at pH values between 4 and 10. For DPhPC liposomes, 267 cholesterol-tagged dsDNA showed increased binding at moderately acidic conditions which 268 269 decreased with increasing pH, while ssDNA and dsDNA-6 nt showed similar binding at all pH 270 values between 4 and 10.
- 271

DNA that was evenly distributed throughout a slide independently of liposome location would be expected to produce a colocalisation score of approximately 1.0, while membrane-binding

2/3 be expected to produce a colocalisation score of approximately 1.0, while membrane-binding

- DNA should produce a colocalisation score greater than 1.0. In the absence of cholesterol tags,
- 275 we did not detect any significant colocalisation of DNA and liposomes at any NaCl
- 276 concentration or pH value. For MgCl₂, no significant colocalisation of DNA with no cholesterol

tag was observed in 19 out of 20 samples. The remaining sample, ssDNA with no cholesterol tag on DOPE/DOPC liposomes at 40 mM, produced a colocalisation score of C = 1.01, significantly greater than one (P < 0.05) but far below those produced by cholesterol-tagged DNA (mean: 1.63 range: 1.18-2.12).



282

283 *Figure 2: The effect of NaCl, MgCl₂ and pH on DNA-liposome colocalisation. Colocalisation* scores and standard deviations are shown for Alexa647-labelled cholesterol-tagged single 284 stranded DNA (ssDNA, pink), cholesterol-tagged double stranded DNA (dsDNA, blue) and 285 286 cholesterol-tagged double stranded DNA with a 6 nt overhang (dsDNA-6nt, green) as well as dsDNA with no cholesterol tag (vellow) and ssDNA with no cholesterol tag (orange) and 287 288 rhodamine-labelled DOPE/DOPC liposomes (left column, a/c/e) and DPhPC liposomes (right column, b/d/f). Conditions tested included extrusion buffer [NaCl] containing 12.5, 25, 50, 100, 289 290 200 and 400 mM NaCl (a/b), extrusion buffer [MgCl₂] containing 0, 10, 20, 40 and 80 mM 291 $MgCl_2$ (c/d) and extrusion buffer [pH] adjusted to pH values of 2, 4, 6, 7, 8 and 10 (e/f).

292

293 The effect of DNA configuration on DNA-liposome colocalisation

- 294 We compared the binding of cholesterol-tagged DNA in different configurations (ssDNA,
- dsDNA and dsDNA-6 nt) across all experiments to assess if one particular DNA configuration
- 296 yielded higher colocalisation than other configurations. For each condition and configuration,
- 297 12 images and were recorded, giving a total of n = 204 colocalisation scores recorded for each
- 298 DNA configuration on each liposome composition.
- We tested for a difference in means of colocalisation scores for each pair of DNA configurations. This *t*-test was repeated for both DPhPC liposomes and DOPE/DOPC liposomes. Our results (Table 1) show there is are significant differences in mean DNAliposome colocalisation dependant on DNA configuration. For DOPE/DOPC liposomes, we found DNA to colocalise in the order $C_{(dsDNA)} \approx C_{(dsDNA-6 nt)} > C_{(ssDNA)}$. For DPhPC liposomes, we found DNA to bind in the order $C_{(dsDNA)} > C_{(dsDNA-6 nt)} > C_{(ssDNA)}$.
- 305
 - Liposome composition
 ΔC(dsDNA) C(ssDNA)
 ΔC(dsDNA)
 C(dsDNA-6
 ΔC(dsDNA-6 nt)
 C(ssDNA)

 nt)
 nt)
 0.0121 (not sig.)
 0.144 (P<0.05)</td>
 0.144 (P<0.05)</td>

 liposomes
 0.203 (P<0.05)</td>
 0.145 (P<0.05)</td>
 0.0577 (P<0.05)</td>

306

Table 1: Mean differences in colocalisation scores for different configurations of DNA binding to DOPE/DOPC liposomes and DPhPC liposomes. Results are pooled from data shown in section 4.2. Paired t-tests of colocalisation scores were conducted comparing each DNA configuration pair-wise on both types of liposomes (n = 204) against the hypothesis 'mean difference = 0'. For example, ' $\Delta C_{(dsDNA)}$ C_(ssDNA)' represents the mean difference in colocalisation scores of dsDNA and ssDNA.

313

314 The effect of membrane cholesterol content on DNA-liposome colocalistion

Colocalisation scores of DNA and liposomes in extrusion buffer showed different trends for the two lipid mixtures as cholesterol content was increased between 0% and 40% mass. For DOPE/DOPC liposomes, colocalisation scores of all three configurations of cholesterol-tagged DNA showed a significant increase between 0% and 40% cholesterol (P < 0.05). Linear regression analysis shows a gradient significantly greater than zero across the observed range

- 320 of membrane cholesterol content (P < 0.05) (Figure 3A).
- 321

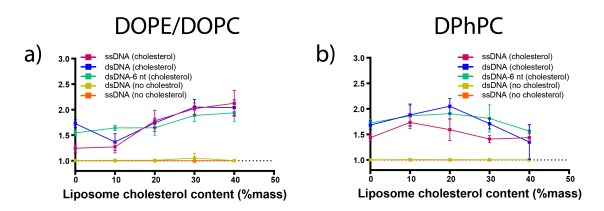
322 For DPhPC liposomes, colocalisation scores of cholesterol-tagged DNA increased to a

maximum at 10%-20% membrane cholesterol, then decreased as membrane cholesterol content

324 was increased above this point. All three configurations of cholesterol-tagged DNA (ssDNA,

- dsDNA and dsDNA-6 nt) showed both a significant increase in colocalisation between 0% and
- 326 20% (P < 0.05) and a significant decrease in colocalisation between 20% and 40% (P < 0.05).

- Linear regression analysis for all three configurations showed a slight overall decreasing trend across the observed range (P < 0.05) (Figure 3 B).
- 329
- Using the test C > 1.0, both ssDNA and dsDNA with no cholesterol tag did not show significant
- 331 colocalisation with liposomes of any cholesterol content.
- 332



333

334 Figure 3: The effect of cholesterol content on DNA-liposome concentration. Colocalisation

335 scores and standard deviations are shown for Alexa647-labelled cholesterol-tagged ssDNA

336 (pink), cholesterol-tagged dsDNA (blue) and cholesterol-tagged dsDNA with a 6 nt overhang

337 (green) as well as dsDNA with no cholesterol tag (yellow) and ssDNA with no cholesterol tag

338 (orange) and rhodamine-labelled DOPE/DOPC liposomes (a) or DPhPC liposomes (b)

339 prepared from lipid stocks containing 0, 10, 20, 30 or 40% cholesterol.

341 Discussion

342 The effect of solution composition on DNA-liposome colocalistion

343 Monovalent ions, divalent ions and pH known to modulate biophysical properties of membrane 344 bilayers (Sachs et al., 2004) and the structural stability of DNA nanostructures (Douglas et al., 345 2009). Divalent cations such as Mg^{2+} are of particular interest to DNA nanotechnology applications to stabilise DNA duplexes and are required to facilitate the formation higher-order 346 nucleic acids structures (Misra and Draper, 1998; Williams et al., 1989). Divalent cations also 347 348 maintain the stability of large DNA nanostructures by inhibiting the electrostatic repulsion 349 between DNA strands and are therefore considered essential in the assembly process (Douglas 350 et al., 2009; Kielar et al., 2018). The sensitivity of DNA to salt and pH has been harnessed to 351 design DNA nanostructures able to switch configuration in response to changes in ion 352 concentration, allowing the development of DNA-based nanosensors (Singh et al., 353 2018). Despite these recent advances, the tolerance of DNA nanostructures to low-salt 354 environments remains variable and design-dependent thus necessitating regular 355 characterization and optimization (Hahn et al., 2014).

356

357 We sought to test whether changes to membrane density and diffusivity due to external buffer 358 would affect cholesterol binding to the non-polar tail-group region of a membrane. Monovalent cations such as Na⁺ and divalent cations such as Mg²⁺ have been shown via modelling to 359 promote lipid-lipid binding interactions within a bilayer (Böckmann et al., 2003) and affect the 360 361 behaviour of water molecules at the membrane-water interface (Velikonja et al., 2013). The 362 interaction between surface charges on membrane bilayers and the phosphate groups of DNA 363 molecules is also affected by divalent cations (Antipina and Gurtovenko, 2016; Binder and Zschörnig, 2002). This interaction is dependent on the ratio of monovalent to divalent salts 364 365 (Budker et al., 1980). In our tests we saw no membrane interactions between non-cholesterol 366 tagged DNA at any MgCl₂ concentration, suggesting that divalent cation-mediated binding was negligible under the conditions tested. Coarse-grained and atomistic simulations of DNA-lipid 367 368 interactions could lead to a more detailed understanding of the effect of cations on the binding 369 of DNA (Uusitalo et al., 2015; Yoo and Aksimentiev, 2015).

370

371 Binding was inhibited in acidic conditions (< pH 4). However, otherwise, there was no optimal 372 pH for liposomes formed from either neutral DPhPC, or DOPE/DOPC, which contains 373 zwitterionic phosphatidylethanolamine headgroups that are positively charged below pH 3.5 374 and negatively charged above pH 8 (Tsui et al., 1986). This suggests lipid ionisation does not 375 play a significant role and variation in pH around physiological pH are not likely to affect 376 membrane binding of cholesterol-tagged DNA. Hydronium ions (H₃O⁺) have a similar effect on lipid bilayers as Na⁺ and Mg²⁺, and may explain the inhibition of binding in strongly acidic 377 conditions (Deplazes et al., 2018). 378

For both liposome compositions, the colocalisation of ssDNA and liposomes was significantly
reduced in comparison with double stranded configurations. This suggests the duplex dsDNA
binds better to liposomes. Membrane-bound ssDNA has been observed through Förster
Resonance Energy Transfer (FRET) (Roy et al., 2008) to lie close to the surface of lipid bilayer

membranes, while dsDNA remains in a stable position protruding normal to the membrane
surface (Ma et al., 2019). Thus, this improved binding may be due to the greater rigidity of
dsDNA vs ssDNA.

The addition of a 6 nt overhang on cholesterol-tagged DNA strands has been postulated to assist during nanostructure assembly by inhibiting strand aggregation (Ohmann et al., 2019). We included a 6 nt overhang next to the cholesterol group on our dsDNA strand (dsDNA-6 nt) and observed a significant decrease in binding only on DPhPC liposomes. Lipid composition should, therefore, be considered when incorporating overhangs, but there would be no large penalty from routine incorporation on membrane-targetting nanostructures.

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The effect of membrane cholesterol on DNA-liposome binding differed between phospholipid compositions. Cholesterol has been shown to affect the strength, fluidity and permeability of bilayers (Róg et al., 2009) as well as the organisation of ions and water molecules at the membrane-bilayer interface (Magarkar et al., 2014). The different response that we observed between different phospholipid types may be explained by differences in tail-group structure. Branched chain lipids such as DPhPC occupy a greater area per molecule within a bilayer compared to linear-chain lipids such as DOPE and DOPC (Tristram-Nagle et al., 2010).

400

Increasing the cholesterol content of lipid mixtures above 20% promoted the binding of DNA
on DOPE/DOPC liposomes but inhibited binding on DPhPC liposomes. This was possibly due
to the lower cholesterol saturation limit of DPhPC compared with DOPE (Huang et al., 1999).
Here we controlled the cholesterol content during liposome preparation but did not quantify
the exact cholesterol content in liposome membranes. Future work using fluorescent markers
or high performance liquid chromatography analysis of liposome samples (Christie, 1985)
could accurately quantify the membrane cholesterol content to better benchmark the role of

- 408 cholesterol in DNA-lipid binding, and to account for any loss in cholesterol during liposome409 preparation.
- 410

411 Conclusion

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In this work we have characterised the role of salt and pH during the assembly of DNAliposome complexes. We tested different lipid species and DNA configurations to screen for optimal conditions to promote binding of DNA to liposomes. Our results suggest that lipid type, pH and DNA configuration are the most important parameters to consider when optimising for the binding of DNA nanostructures to liposomes, whereas mono- and divalentsalt concentration plays a minor role. These results will be helpful in experimental design and reagant choice for future experiments combining DNA and lipid nanotechnologies.

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