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# 3 Frog nest foam as a drug delivery system

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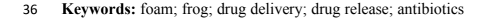
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### 20 Abstract

21 Foams have frequently been used as systems for the delivery of cosmetic and therapeutic molecules; however, there is high variability in the foamability and long-term stability of 22 synthetic foams. The development of pharmaceutical foams that exhibit desirable foaming 23 24 properties, delivering appropriate amounts of the active pharmaceutical ingredient (API) and that have excellent biocompatibility is of great interest. The production of stable foams is rare 25 in the natural world; however, certain species of frogs have adopted foam production as a means 26 of providing a protective environment for their eggs and larvae from a predators and parasites, 27 to prevent desiccation, to control gaseous exchange, temperature extremes, and to reduce UV 28 damage. These foams show great stability (up to 10 days in tropical environments) and are 29 highly biocompatible due to the sensitive nature of amphibian skin. This work demonstrates for 30 the first time, that nests of the Túngara frog (Engystomops pustulosus) is stable ex situ with 31 32 useful physiochemical and biocompatible properties and is capable of encapsulating a range of compounds, including antibiotics. These protein foam mixtures may find utility as a topical 33 drug delivery system (DDS). 34

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### 38 Introduction

Foams have been used as delivery systems or vehicles to deliver cosmetic and therapeutic 39 molecules to normal and injured skin since the 1970's [1-5]. Yet the long-term stability of liquid 40 aqueous foams has been a challenge, with some formulations offering useful foamability 41 42 properties (e.g. foam expansion time), but poor stability[6]. There has been some progress made through the combination of various foam and surfactant components to create high foamability 43 and long-term foam stability[6–9]. However, the development of biocompatible, liquid foams 44 with high foamability and long-term stability remains a challenge in materials science[6]. A 45 range of foams is already in use for topical treatments, such as Ibuprofen foams (Biatain® Ib) 46 used to relieve a wide range of exuding wounds, urea-containing foams (KerraFoam) to help 47 alleviate the symptoms of psoriasis, and antibiotic foams containing clindamycin and other 48 antimicrobials such as silver sulfacetamide[2,4]. A major advantage of medicated foams is their 49 ability to cover large surface areas, while containing highly concentrated drugs for topical 50 treatment[1]. One major difficulty can be the delivery of an adequate concentration of active 51 52 pharmaceutical ingredients (APIs) for treatment over a sustained period, therefore often necessitating repeated, regular applications. In the case of open wounds and burns, regular 53 removal of dressings may lead to increased infection risk and damage to healing surfaces, aid 54 the emergence of antimicrobial resistance through the delivery of sub-minimum inhibitory 55 concentrations of antibiotics, ultimately resulting in reduced infection control and wound 56 healing[10]. Liposomes have proposed for dermatological applications; however, are exhibit 57 major stability issues. There is therefore a need for the development of biomaterials that allow 58 extended times between application combined with high stability and improved 59 biocompatibility; natural foams can provide these benefits. 60

Anurans (frogs) exhibit enormous diversity in reproductive strategies and styles[11], and many 61 species of tropical and subtropical frog lay their eggs in stable proteinaceous foams that differ 62 in composition between species. Foam-nesting behaviour, thought to have evolved as a means 63 to avoid aquatic predators, prevent desiccation of eggs, control gaseous exchange, buffer 64 temperature extremes, reduce solar radiation damage, and protect eggs from microbial 65 colonisation[12]. Stable biological foams and foam-producing surfactants are rare in nature, 66 presumably due to the requirement for high-energy input for their generation, and the potential 67 of surface-active components to negatively affect cell and membrane function[13,14]. Frog nest 68 foams are remarkable for their strong surfactant activity combined with harmlessness to naked 69 eggs and sperm. The leptodactylid frogs of the neotropics are one such anuran lineage that has 70

evolved stable foams as an offspring protection mechanism. The nests of the Túngara frog 71 (Engystomops pustulosus) are remarkable in structures that act as incubation chambers for 72 eggs[15] (Supp. Fig. 1A & B; https://doi.org/10.6084/m9.figshare.13281416.v1), allowing 73 rapid growth and development of embryos, offering a protective environment against predation, 74 whilst providing temperature regulation and oxygen transfer for optimal growth conditions. 75 These nests, there are not destroyed by microbes during larval development despite construction 76 within highly-microbe-rich water[16]. The foam nest structure is highly stable, remaining 77 assembled for as much as 10 days in the tropical environment[17,18], yet the surfactant activity 78 79 of the foam does not cause damage to the sperm, eggs or developing embryos[13]. The main surfactant protein within these nests is an 11kDa protein, Ranaspumin-2 (RSN-2), which does 80 81 not disrupt biological membranes or cells, but it still provides sufficient surfactant activity at 82 the air-water interface to allow foam formation[19]. The RSN-2 protein appears to form a 83 clamshell-like structure, which can undergo an unfolding conformational change to expose nonpolar patches on the protein surface to the air, while highly polar regions remain in contact 84 85 with the water interface to provide the surfactant activity. RSN-2 has been successfully used in industry as a surfactant in nanoparticle production[20], but the use of the whole nest foam 86 87 protein composition in drug delivery or for tissue engineering applications has not been explored. 88

Here we show that the unseparated, total protein mixture of *E. pustulosus* nest foam is stable *ex* situ for extended periods with useful physiochemical and biocompatible properties and does not the need of the addition of oxygen as with other drug delivery systems ([21]; DDS) and may be used to encapsulate a range of hydrophobic and hydrophilic model compounds. These data suggest that anuran-derived protein foams may have potential applications as DDSs.

### 95 **Results & Discussion**

## 96 Biophysical properties of Tungara frog foam preparation

The composition and properties of DDSs can determine the effectiveness of drug release, and 97 while several factors may affect drug release, the thermodynamics driving the passive diffusion 98 process is key[22]. Thus, increasing and stabilising the local concentration of the permeant is 99 the simplest strategy to facilitate bioavailability[3]. The protein composition of *E. pustulosus* 100 foam fluid **SDS-PAGE** Fig. 101 was analysed by (Supp. 1C; https://doi.org/10.6084/m9.figshare.13281416.v1), confirming previous work that foam from 102 this species contains six major proteins ranging between 10 and 40kDa in size[18] and that the 103 foam nests used in this study were of typical composition. CD spectra of the samples showed a 104 negative maximum at 215 nm and a positive maximum at 194 nm, (Fig. 1A) indicating that, 105 cumulatively, the protein mixture in the foam fluid comprises predominately  $\beta$ -sheet structures. 106 Further insight into secondary structure of the foam mixture was obtained by FTIR, which also 107 exhibited spectra consistent with overall dominance of  $\beta$ -sheet structures (Fig 1B;[23]). For 108 both foam solutions, transitions were observed at ~1680 cm<sup>-1</sup>, which are frequencies 109 characteristic of Amide I band, signifying C=O bond stretches typically engaged in β-sheet 110 bonded network structures. These data support previous observations that average secondary 111 structure content of the proteins is predominantly  $\beta$ -sheet[18] and indicates that the 112 centrifugation steps in the preparation of the foam for these experiments does not alter the 113 114 overall structure and composition of the foam from the wild *E. pustulosus* nests.

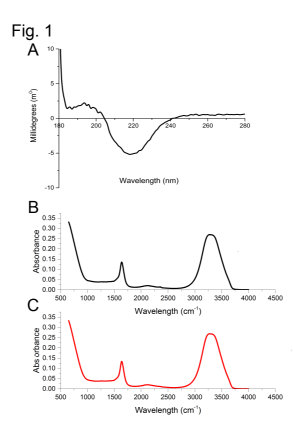




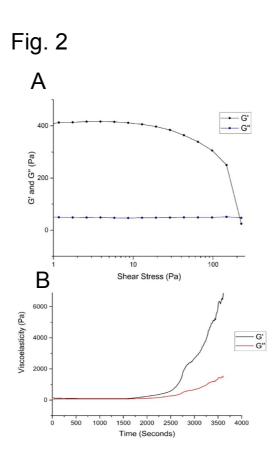
Figure. 1. Structural characterisation of *E. pustulosus* nest foam proteins. 1A: Circular Dichroism of foam fluid using 0.1mm pathlength cuvettes containing 1mg/ml protein foam fluid solution. 1B: Fourier Transform Infrared Spectroscopy (FTIR) foam fluid (B) and whole foam (C). For both CD and FTIR spectra were corrected for baseline and buffer effects each measurement was carried out in triplicate and the mean of the data is presented.

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To investigate the viscoelastic properties of the foam, oscillation sweep experiments were 122 employed by rheology. The whole foam fluid was found to tolerate up to 100 Pa of shear stress 123 force before it reaches a breaking point, at which the elastic modulus of the foam decreases and 124 foam structure and stability is lost (Fig. 2A). Time sweep experiments indicated that up to 1500 125 s the foam preparation moduli are unchanged by stress and frequency. After 1500 s both elastic 126 and viscosity moduli increase demonstrating that water is being lost from the foam (Fig. 2B). 127 It has been suggested that stress increases the chances of water loss from the foam followed by 128 coarsening, which leads to an increase in viscoelasticity [24]. The *E. pustulosus* whole foam is 129 able to withstand shear stress and pressure before breaking down, demonstrating the long-130 lasting stability that may be observed in nature. Pharmaceutical foams are typically required to 131 remain stable in order to be properly manipulated while being applied, but have low shear 132 allowing them to break down shortly thereafter [25,26]. The foam derived from E. pustulosus 133

has exhibits long-term stability in harsh tropical environments (e.g. heat, high level exposure
to ultraviolet light, and physical disruption) and behaves differently from typical
pharmaceutical foams[2]. The *E. pustulosus* foam is stable enough to be manipulated and able
to withstand shear forces, giving the potential to deliver drugs over prolonged periods.

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Figure 2. Viscoelastic properties of *E. pustulosus* nest foam. 2A: Time sweep rheology data for foam, showing both elastic (G') and viscous (G'') moduli. Stress was set at 100 Pa, and carried out over 1 hour. 2B: Oscillation sweep rheology data for *E. pustulosus* foam, showing both elastic (G') and viscous (G'') moduli. Shear stress was increased from 1 Pa to 200 Pa. Each measurement was taken in triplicate at 20<sup>o</sup>C.

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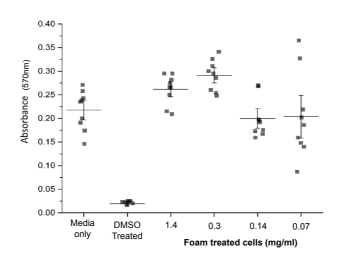
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## 150 Frog foam biocompatibility with human epithelial cells

To investigate the susceptibility of mammalian cells to any possible toxic effects of the foam, 151 cells were cultured in the presence of foam fluid and potential toxicity was assayed using an 152 MTT-based cell viability assay. Exposing HaCaT keratinocyte cells to a range of *E. pustulosus* 153 154 foam concentrations did not affect the overall cell viability and multiplication of the cells (Fig. 3). The higher foam fluid concentrations to which the cells were exposed are representative of 155 foam concentration present in *E. pustulosus* nests (1-2mg/ml protein[17]). Cells exposed to the 156 foam behave in the same manner as untreated control cells, demonstrating that the foam proteins 157 from *E. pustulosus* are non-toxic to epithelial cells and are therefore unlikely to cause damage 158 to the skin or underlaying tissues if used as a topical drug delivery system. The foam and its 159 protein components are already known to be harmless to human erythrocytes [27]. This high 160 degree of biocompatibility is consistent with the foam and its precursor components being 161 harmless to naked amphibian sperm, eggs, and oviduct surfaces of the frogs[27]. 162





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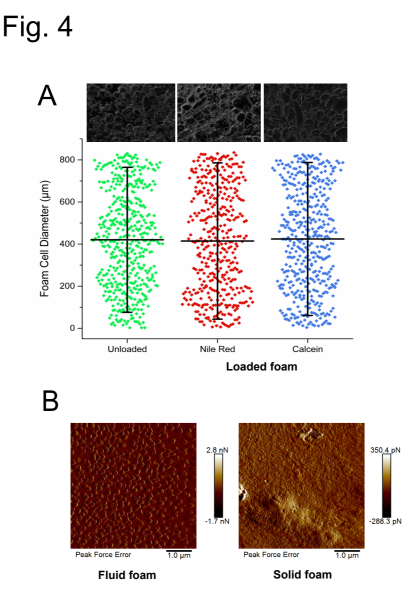
**Fig. 3. Biocompatibility of** *E. pustulosus* **nest foam with human epithelial cells.** MTT assay of HaCaT cell viability following exposure to a range of fluid foam concentrations over 24 hours at 37<sup>o</sup>C. Each treatment was performed in triplicate, and media alone was used as normal viability control, and cells were treated with DMSO for non-viable control. Treatments 1 to 4 were a dilution of fluid foam protein concentrations - 1.4 mg/ml, 0.3 mg/ml, 0.14 mg/ml and 0.07 mg/ml respectively. Error bars represent the standard deviation of the data about the mean (horizontal line).

## 172 In vitro release of drug-loaded frog foam

A single foam cell is defined as a bubble of gas enclosed in a liquid film that can be polyhedral or circular, heterogeneous or homogeneous, and usually range between 0.1 and 3mm in diameter[25]. The cell structure of *E. pustulosus* nest-foam was evaluated microscopically and the foam cell sizes measured (**Fig. 4A**). The foam cells in all samples were found to be a heterogeneous mixture of uneven, spherical and polyhedral cells with a Feret diameter ranging from 10-800  $\mu$ m, falling in the normal range of foam cell size of foams that have been used previously in pharmaceutical applications[25].

180 The relative density of the foam was found to be 0.25 g of protein/ml. AFM PeakForce analysis 181 of the fluid foam and gel foam indicated consistency of the individual adhesion force ( $F_{ad}$ ) 182 measurements in each foam forms (**Fig. 4B**) indicating that the foam surface forces are 183 homogeneous across the surface of each form. Moreover, in the case of the fluid foam, the  $F_{ad}$ 184 is higher and the AFM images shows the formation of ~200 nm droplets as the foam was dried 185 on to mica surfaces. This combination of low density and high structural stability is unusual 186 and could suggest the utility of anuran foam-nest proteins as a pharmaceutical foam.

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202 Fig. 4. Drug loading of *E. pustulosus* nest foam does not alter the structure of the foam. 4A. Foam cell diameter measurements scatter plot. Feret diameter of each foam cell/bubble was 203 measured using Fiji software. Bars on the scatter encompass 10-90% of the data points, with 204 205 the central horizontal line representing mean values. Above the scatter graph are representative images of unloaded foam, foam loaded with 1 mg/ml NR and foam loaded with 1mg/ml calcein. 206 All images were taken using freshly defrosted foam. 4B. Atomic Force Microscopy PeakForce 207 analysis of foam fluid and gel foam to investigate the consistency of the adhesion force (Fad) in 208 the foam. 209

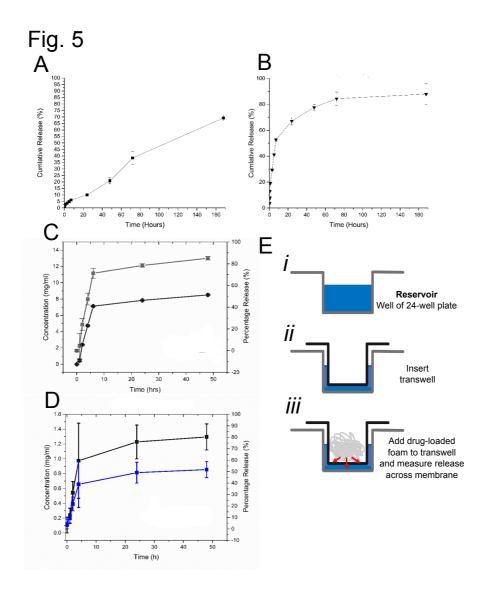
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To evaluate the drug release behaviour of E. pustulosus foam, experiments were performed 212 using a dialysis-based method where two model compounds (one hydrophobic and one 213 hydrophilic), NR and calcein, were encapsulated in the foams. Calcein exhibits a "burst-type" 214 release profile from the foam, whereas NR is discharged at a linear rate over 168 hours (Fig. 215 5A & B). These data indicate that the whole foam can absorb and release both hydrophobic 216 (NR) and hydrophilic (calcein) molecules and release these at different rates over a prolonged 217 218 period, with up to 85% of the loaded dye released over a period of 168 hours. Moreover, the 219 loading of the nest foam with NR or calcein did not alter the cell size or shape of the foam (Fig. 220 4).

To test the ability of the foam to release a clinically relevant drug molecule, the whole foam loaded in the same manner as for the previous two molecules with the red-pigmented antibiotic rifampicin. The dialysis method of release showed that rifampicin was released at a steady rate over the first five hours with a release of around 80% of loaded antibiotic, followed by a second, slower phase of release (**Fig. 5C**).

To further investigate the release of drug molecules from E. pustulosus foam, a novel trans-226 well-based release assay was developed, which emulates delivery of the foam preparation 227 across a more complex protein coated surface. Foam was loaded in to a collagen-coated 228 229 transwell sat in a 24-well tissue culture plate containing 1 ml of PBS and release of drug molecule was determined through the assay of the amount of rifampicin that passed from the 230 foam through the transwell membrane. In the transwell assay, around 50% of rifampicin was 231 also released over a 48 hrs (Fig. 5D). This novel transwell assay for investigating compound 232 release from foams offers a simpler route to assaying drug release from foams and other 233 aqueous-based materials that does not require the manipulation of dialysis tubing. Moreover, 234 the release mimics clinical applications through release from a single face of the transwell (A 235 schematic of the Transwell-based assay is provided in Fig. 5E) with the advantage that 236 transwells are available in a number of sizes and pore diameters to suit a range of needs. 237

These data indicate that the foam can be loaded with drug molecules and can be used as an extended release system. The foam from *E. pustulosus* is stable and the API release is relatively slow with the foam potentially acting as a barrier in the local environment. The *E. pustulosus* foam compares well with release of rifampicin from loaded nanoparticles where 80-90% of release occurs, but within an 8 h time window[28].



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244 Fig. 5. *E. pustulosus* nest foam can take up and release model compounds and drug molecules. 5A: 245 Cumulative release of the dye NR from loaded whole foam over 168 hrs using the dialysis method. 246 5B: Cumulative release of the dye calcien from loaded whole foam over 168 hrs using the dialysis method. 5C: Cumulative release of the antibiotic rifampicin from loaded whole foam over 168 hrs 247 248 using the dialysis method;  $-\bullet$ - concentration (mg/ml);  $-\blacksquare$ - percentage release of the dye or drug. **5D**: Release of rifampicin using novel transwell method; -■- concentration (mg/ml of protein); -■-249 250 percentage release of rifampicin. Each point represents the mean of data collected in triplicate and error bars indicate the standard deviation of the data. Sink conditions satisfied by replacing sample 251 252 volume with fresh buffer at each sample point. Dye release concentrations were calculated using 253 spectrophotometric standard curves for respective dye or drug. 5E: Schematic representing the 254 experimental set-up using the novel transwell release assay, with reservoirs of 24-well plate filled with buffer (i), the insertion of the permeable transwell (ii), and the addition of drug-loaded foam to 255 256 the transwell to allow drug release across the transwell membrane which can be quantified in the 257 reservoir.

Many nanoparticle-based systems release their drug-load rapidly resulting in ineffective long-258 term treatment possibilities[29] and have shown to exhibit some toxic properties[30]. The 259 anuran foam-based preparations extend this release period to around 48 hrs depending on the 260 261 nature of the compounds used, expanding the possible utility of these foam systems. While there has been multiple antibiotic loaded liposomes[31-33](AmBisome, Lambin, Doxil) 262 brought to market, little have transferred the applicability to topical conditions, and their 263 efficiency is still lacking. Further, tetracycline loaded nanocomposite hydrogels have been 264 presented as for extended release delivery through the skin, released of maximum of 15% of 265 266 the loaded antibiotic[32]. Foams are considered more popular with patients than gels[2], and a stable foam may provide a solution to both these issues. Previous studies of pharmaceutical 267 268 foams have investigated the immediate delivery of drugs through to the dermis by exploiting the fast breakdown of foam preparations, but rarely have they been used for long-term drug 269 270 release[2]. The foam derived from *E. pustulosus* provides a material with intermediate release properties, where uptake is efficient, its high stability and slow release properties enable 271 272 continuous release over a clinically useful period of time.

#### 273 Conclusions

There is an increased interest in drug delivery systems to refine the use of antimicrobial drugs 274 currently available on the market, which may enable novel delivery systems help combat the 275 rise of antimicrobial resistance in the clinic. Anuran foams from reproductive nests may provide 276 a novel area for investigation in the area of controlled release. Anuran foam nests are highly 277 biocompatible, durable and stable, and have excellent drug release properties. They exhibit 278 none of the issues associated with fabric-based drug release such as instability, rapid release 279 characteristics or toxicity. These advantages suggest that anuran foam may provide a novel 280 topical drug delivery system for controlled release of drugs. 281

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### 288 Materials and Methods

### 289 Materials

Nile red (NR), calcein, ethanol 97% (v/v), phosphate buffered saline (PBS) tablets (pH 7.2)
were all purchased from Sigma and MTT (3-(4,5-dimethylthiazol-2-yl)-2,5diphenyltetrazolium was purchased from Thermofisher.

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### 294 Collection of *Engystomops pustulosus* foam

Freshly laid foam nests or adult frogs in amplexus (allowed to lay in captivity, before release) 295 were collected from a number of sites in northern Trinidad in June and July of 2014, 2015 and 296 2016. Nests were removed from the surface of the water in which they were produced and the 297 eggs were removed manually before being stored at -20°C for transfer to the Glasgow, where 298 they were stored at -80°C. Foam was freshly defrosted for all experiments. Soluble foam 299 300 material ('foam fluid') was produced by centrifuging whole foam for 10 min (16.000 x g). 301 providing a solution with approximately ~2 mg/ml protein. This also yields a supernatant/pellicle residual of semi-solid compressed foam ('gel foam') on top of the foam 302 fluid layer. 303

## 304 Microscopy

Optical microscopy: Whole foam was defrosted at room temperature before use. All foam
 images were taken using transmitted light on a Nikon SMZ1500 stereomicroscope with images
 acquired using a DFK 33UX264 CMOS camera (The Imaging Source Europe GmbH,
 Germany) using NIS-Elements AR.3.2 software. Fiji software (https://fiji.sc/) from the ImageJ
 (https://imagej.net) package was used for image analysis.

Atomic force microscopy (AFM): Samples (5 µl) of foam were deposited onto a freshly cleaved 310 311 mica surface (1.5 cm x 1.5 cm; G250-2 Mica sheets 25 mm x 25 mm x 0.15 mm; Agar Scientific Ltd, Essex, UK) and left to dry at room temperature for 1h before imaging. The images were 312 obtained by scanning the mica surface in air under ambient conditions using a Scanning Probe 313 Microscope (MultiMode® 8, Digital Instruments, Santa Barbara, CA, USA; Bruker Nanoscope 314 analysis software Version 1.40), operating using the PeakForce QNM mode. The AFM 315 measurements were obtained using ScanAsyst-air probes, for which the spring constant (0.58 316 N/m; Nominal 0.4 N/m) and deflection sensitivity had been calibrated, but not the tip radius 317 (the nominal value used was 2 nm). 318

### **319** Sodium Dodecyl Sulfate Poly-Acrylamide Gel Electrophoresis (SDS-PAGE)

- 320 Solid and liquid foam samples were electrophoresed on precast NuPAGE 15% polyacrylamide
- 321 Bis-Tris gels (Invitrogen) at 120V using 4 X SDS reducing loading buffer (Invitrogen). Each
- 322 gel was stained with InstaBlue Coomassie protein stain for ~45 min.

### 323 Circular Dichroism Spectroscopy (CD)

CD was used to investigate the overall secondary structure content of the proteins. Spectra were 324 acquired using a Chirascan Plus (Applied Photophysics) instrument using a 0.1mm quartz 325 cuvette (Hellma) at 20°C. All samples (10 mg/ml protein) were measured in the far-UV in a 326 wavelength range of 180 nm to 280 nm range, with step size of 1 nm, bandwidth of 1 nm, and 327 reading time of 1 s per nm. Triplicate measurements were taken for each sample run, baseline 328 peak, PBS control and foam sample spectra, with triplicate spectra then averaged. Baseline and 329 PBS traces where subtracted from the sample spectra before secondary structure predictions 330 were made. All data analysis was performed using Global3 software and Excel. 331

## 332 Fourier transform infrared spectroscopy

Fourier transform infrared (FTIR) spectroscopy was carried out using a Nicolet iS10 Smart iTR spectrophotometer (Thermo Scientific). Solid and liquid foam spectra were recorded in the range on 4000 cm<sup>-1</sup> and 500 cm<sup>-1</sup>, over 128 scans at a resolution of 4 cm<sup>-1</sup> and an interval of 1 cm<sup>-1</sup>. Background spectra were measured and the foam spectra were corrected accordingly.

### 337 Rheology

Rheology measurements were determined using a HAAKE MARS Rotational Rheometer (Thermo Scientific). Foam samples were subjected to oscillation sweeps and time sweeps. All experiments were carried out using P20 upper plate and TM20 lower plate. The oscillation sweeps were completed with a 1 mm gap and 0.1 Pa to 200 Pa range. Time sweep experiments were run for 1 h, at 100 Pa and 3 Hz using a 0.5 mm gap. Data points were collected in triplicate and averaged before analysis was carried out.

### 344 MTT cell viability assay

HaCaT cells (CLS, Eppelheim, Germany), a model human keratinocyte cell line were cultured in Dulbecco's modified Eagle's medium (DMEM) containing 4.5 g/l glucose supplemented with 10% (v/v) fetal bovine serum, 2 mM L-glutamine and 50 units/ml penicillin/streptomycin (cDMEM; Lonza, Slough, UK). Soluble foam proteins were prepared as above, with the supernatant being passed the a 0.22  $\mu$ m filter (Millex 33 mm) and subsequently concentrated

using an Amicon 10 kDa spin filter. The protein concentration was determined by Bradford 350 assay (BioRad). The HaCaT cells were plated onto 96 well plates (~1x10<sup>3</sup> cells per well and 351 grown to 80% confluence) and were treated with buffer containing foam proteins 352 (concentrations indicated in the figures) prior to incubation at 37 °C for 24 hours. After 24 hours 353 the media was removed from the cells, and replaced with 50 µl of fresh media and 50 µl of 354 MTT (5mg/ml) and incubated for 1 h at 37 °C. This was followed by replacing the media with 355 100 µl DMSO and further incubation in the dark at room temperature for 30 minutes prior to 356 reading the absorbance at 570nm[34]. Results were expressed as the % viability compared to 357 358 non-treated cells  $\pm$  SEM.

### 359 In vitro release of model compounds

Aliquots (500 mg) of whole foam were loaded with dve by mixing with either 400 µl of Nile 360 Red (NR; hydrophobic; 1 mg/ml in ethanol) or calcein (hydrophilic; 1 mg/ml in ethanol). The 361 mixture was the placed in dialysis tubing and sealed before being submerged in 10 ml PBS at 362 37 °C (pH 7; for NR-based release experiments, a 1:1 mixture of ethanol and PBS was used). 363 The release experiments were carried out at 37 °C, over 168 hours. To satisfy the perfect-sink 364 conditions, which allow for the determination of the diffusion parameters, the supernatant was 365 replaced with fresh PBS at 37 °C at each time point (indicated in the graphs). The concentration 366 of model compound in each sample was determined spectrophotometrically at 490 nm (calcein) 367 or 590 nm (NR) and the concentration determined with reference to standard control calibration 368 369 curves. Experiments were performed in triplicate.

### 370 In vitro Antibiotic release

371 Two *in vitro* techniques were used to investigate the release of the antibiotic rifampicin.

372 *Dialysis:* Aliquots (400 mg) of foam were mixed with 400  $\mu$ l of rifampicin (25 mg/ml). The 373 loaded foam was placed into dialysis tubing, sealed and submerged in 10 ml of PBS. This was 374 incubated at 37 °C for 48 hours. Samples (1 ml) were taken and fresh media added to maintain 375 sink conditions. Samples where measured by spectrophotometrically at 475 nm[35] against a 376 calibration curve.

377 *Transwell:* Aliquots of foam (100 mg) were loaded with 100 μl of rifampicin (25mg/ml).
378 Loaded foam was the placed into a transwell collagen-coated permeable support (0.4 μm;
379 Nunc). Each support was inserted into 24 well plate well containing 600ul of PBS. The plate

- 380 was then incubated for 48 hours at 37  $^{\circ}$ C. PBS (600µl) was collected from a well for each time
- point, and the absorbance measured at 475nm, in triplicate.

## 383 Supporting Information.

- 384 Supplementary Figure https://doi.org/10.6084/m9.figshare.13281416.v1; Fig. 1A. Adult
- 385 Túngara frog (*Engystomops pustulosus*). **B.** *In situ E. pustulosus* foam nest. **C.** SDS-PAGE gel
- 386 of *E. pustulosus* whole foam from a wild-collected nest electrophoresed

## 387 Acknowledgements

The authors would like to acknowledge the Engineering and Physical Science Research Council (EPSRC) via the Doctoral Training Centre (DTC) at the University of Strathclyde for the PhD studentship support to SB. We would also like to thank Prof. Roger Downie, University of Glasgow for his long-term assistance in the field and advice on Tungara Frogs. We acknowledge the support of the Microbiology Society and the Pauline Fitzpatrick Memorial Travel Fund to SB to support fieldwork in Trinidad. EMO was supported by a PhD studentship from the Psoriasis Association (ST3 15).

- We would also like to thank the Wildlife Section, Forestry Division, of the Government of Trinidad and Tobago for issuing Special Game Licences under the Conservation of Wildlife Act, permitting us to collect *E. pustulosus* nests (Special Game Licences 2014-2016 and Wildlife Special Export Licence numbers: 001741, 001161 and 000646).
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## 400 **Conflict of interest**

401 All authors declare that they have no conflict of interest in relation to this work.

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### 504 Figure legends

**Fig. 1. Structural characterisation of** *E*, *pustulosus* **nest foam proteins. 1A:** Circular Dichroism of foam fluid using 0.1mm pathlength cuvettes containing 1mg/ml protein foam fluid solution. **1B:** Fourier Transform Infrared Spectroscopy (FTIR) foam fluid (**B**) and whole foam (**C**). For both CD and FTIR spectra were corrected for baseline and buffer effects each measurement was carried out in triplicate and the mean of the data is presented.

Fig. 2. Viscoelastic properties of *E. pustulosus* nest foam. 2A: Time sweep rheology data for foam, showing both elastic (G') and viscous (G'') moduli. Stress was set at 100 Pa, and carried out over 1 hour. 2B: Oscillation sweep rheology data for *E. pustulosus* foam, showing both elastic (G') and viscous (G'') moduli. Shear stress was increased from 1 Pa to 200 Pa. Each measurement was taken in triplicate at 20°C.

Fig. 3. Biocompatibility of *E. pustulosus* nest foam with human epithelial cells. MTT assay of HaCaT cell viability following exposure to a range of fluid foam concentrations over 24 hours at 37°C. Each treatment was performed in triplicate, and media alone was used as normal viability control, and cells were treated with DMSO for non-viable control. Treatments 1 to 4 were a dilution of fluid foam protein concentrations - 1.4 mg/ml, 0.3 mg/ml, 0.14 mg/ml and 0.07 mg/ml respectively. Error bars represent the standard deviation of the data about the mean (horizontal line).

522 Fig. 4. Drug loading of *E. pustulosus* nest foam does not alter the structure of the foam.

4A. Foam cell diameter measurements scatter plot. Feret diameter of each foam cell/bubble was measured using Fiji software. Bars on the scatter encompass 10-90% of the data points, with the central horizontal line representing mean values. Above the scatter graph are representative images of unloaded foam, foam loaded with 1 mg/ml NR and foam loaded with 1mg/ml calcein. All images were taken using freshly defrosted foam. **4B.** Atomic Force Microscopy PeakForce analysis of foam fluid and gel foam to investigate the consistency of the adhesion force ( $F_{ad}$ ) in the foam.

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Fig. 5. *E. pustulosus* nest foam can take up and release model compounds and drug molecules. 5A: Cumulative release of the dye NR from loaded whole foam over 168 hrs using the dialysis method. 5B: Cumulative release of the dye calcien from loaded whole foam over 168 hrs using the dialysis method. 5C: Cumulative release of the antibiotic rifampicin from loaded whole foam over 168 hrs using the dialysis method; -●- concentration (mg/ml); -■-

percentage release of the dye or drug. **5D:** Release of rifampicin using novel transwell method; 536 - concentration (mg/ml of protein); - percentage release of rifampicin. Each point 537 represents the mean of data collected in triplicate and error bars indicate the standard deviation 538 of the data. Sink conditions satisfied by replacing sample volume with fresh buffer at each 539 sample point. Dye release concentrations were calculated using spectrophotometric standard 540 curves for respective dye or drug. **5E:** Schematic representing the experimental set-up using 541 the novel transwell release assay, with reservoirs of 24-well plate filled with buffer (i), the 542 insertion of the permeable transwell (*ii*), and the addition of drug-loaded foam to the transwell 543 544 to allow drug release across the transwell membrane which can be quantified in the reservoir.

## 545 Supplementary Figure <u>https://doi.org/10.6084/m9.figshare.13281416.v1</u>

Supplementary Fig. 1A. Adult Túngara frog (*Engystomops pustulosus*). B. *In situ E. pustulosus* foam nest. C. SDS-PAGE gel of *E. pustulosus* whole foam from a wild-collected
nest electrophoresed through a 4-20% Tris-Glycine NuPAGE gel under reducing conditions.
Proteins in the 10-30 kDa range are the ranaspumins (RSN described by Fleming et al., 2009 highlighted by red boxes. RSN molecular mass based on amino acid sequence: RSN-1, 14kDa;
RSN-2, 11kDa; RSN-3, 18kDa; RSN-4, 21kDa; RSN-5, 18kDa; RSN-6, 27kDa) Marker: 10200kDa Broad Range marker (New England Biolabs; #P7704).

## 554 **Reference**

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