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1 Characterization of virus-like particles associated with the human faecal and caecal

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24 ABSTRACT

25	This work represents an investigation into the presence, abundance and diversity of
26	virus-like particles (VLPs) associated with human faecal and caecal samples. Various
27	methodologies for the recovery of VLPs from faeces were tested and optimized, including
28	successful down-stream processing of such samples for the purpose of an in-depth electron
29	microscopic analysis, pulsed-field gel electrophoresis and efficient DNA recovery. The
30	applicability of the developed VLP characterization method beyond the use of faecal samples
31	was then verified using samples obtained from human caecal fluid.
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33	Keywords: bacteriophages; microbial ecology; gastrointestinal tract; transmission electron
34	microscopy; caecum; faeces.
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37 1. INTRODUCTION

³⁸ Viruses are the most numerous biological entities within the biosphere of our planet ³⁹ (being present at an estimated number of $\sim 10^{31}$), with bacteriophages representing the most ⁴⁰ abundant group of environmental viruses [1, 2]. Bacteriophages are ubiquitous viruses that ⁴¹ infect bacterial cells and disrupt their metabolism. Multiple bacteriophage types can infect a ⁴² specific microbial isolate, with most bacteriophages infecting only certain species or even ⁴³ strains of bacteria [3].

Although they are abundant and potentially important to microbial populations 44 indigenous to different ecological niches within the human gastrointestinal tract and to host 45 health, little attention has been paid to bacterial virus-like particle (VLP) assemblages and 46 their interactions with the gastrointestinal microbiota and/or human host until recently. 47 Various publications have highlighted the potential importance of bacteriophages in 48 inflammation states, including Crohn's disease [4, 5, 6], and as therapeutic agents [7, 8]. 49 Breitbart et al. [9] conducted the first metagenomic study on dsDNA-containing VLPs 50 associated with the human faecal virome using a fresh sample from a healthy adult male. On 51 the basis of previous estimates of gut microbial diversity, Breitbart et al. [9] predicted that 52 there are two to five times more viral genotypes (~ 1.200 viral genotypes) present in the 53 human gastrointestinal microbiota than the number of bacterial species, with the vast majority 54 of these VLPs representing bacteriophages and prophages. RNA viruses present in human 55 faeces have been found to be mostly associated with plant viruses, such as Pepper Mild 56 Mottle Virus, with RNA bacteriophages making a minimal contribution to the diversity of the 57 intestinal virome [10]. Using frozen samples from four adult female monozygotic twins and 58 their mothers. Reves et al. [11] demonstrated that each individual harbours a unique virome 59 (ssDNA and dsDNA) regardless of their genetic relatedness to another individual, and that 60 intrapersonal virome diversity is very low, with >95 % of virotypes retained over a one-year 61

period (between 71 and 2,773 viral genotypes identified among the samples). More recently,
Minot *et al.* [12] confirmed inter-individual variation in the virome and, similar to
modulation of the faecal bacteriome by foodstuffs, demonstrated that diet influences host
faecal virome structure. Minot *et al.* [13] also confirmed the relative stability of an
individual's virome, monitoring 24 faecal samples from one individual over a 2.5-year
period.

The murine virome has been used to demonstrate enrichment of bacteriophageencoded, antibiotic-resistance genes (related and unrelated to the drug used) after antibiotic
therapy, with the adaptive capacity of the virome (specifically its bacteriophage component)
suggested to protect gut bacteria, thereby preserving the microbiota's robustness during
antibiotic stress [14].

It is clear from the aforementioned studies that the virome-bacteriome community in 73 the gut is governed by complex and dynamic interactions in health, and that its balance may 74 be disturbed when under stress, e.g. during antibiotic intervention [14]. Although 75 metagenomic studies have greatly improved our understanding of the virome associated with 76 the human and murine gut microbiomes, it has been notable how little bacteriophage 77 material, and consequently DNA, has been isolated from samples when this information was 78 included in a publication. Reves et al. [11] reported the isolation of ~500 ng DNA from 2 to 5 79 g of frozen faeces, whereas Thurber et al. [15] stated that between 500 and 3000 ng of DNA 80 could be isolated from 500 g of human faeces, though the publication they cite [9] provides 81 no information regarding the amount of DNA isolated from the 500 g faecal sample 82 examined in the original study. In addition, to the best of our knowledge, no attempts have 83 been made to enumerate bacteriophages in faecal filtrates prior to CsCl purification. The aims 84 of this study were to develop reliable and effective methods for the recovery and 85

- 86 characterization of VLPs in human samples, and to apply these methods to human faecal and
- 87 caecal samples to demonstrate the methods' efficiency.

88 2. MATERIALS AND METHODS

89 2.1 Processing of and isolation of VLPs from faecal and caecal samples

90 Faecal samples were obtained from six healthy adult (2 male, 4 female) donors of between 23 and 52 years of age, all of whom were members of the Department of Food and 91 Nutritional Sciences, University of Reading. None of the donors had taken antibiotics, 92 prebiotics and/or probiotics in the 3 months prior to sampling, and none had any history of 93 gastrointestinal disorder. All donors gave informed oral consent for their faecal samples to be 94 used for microbiological analyses. Ethical approval for the collection of caecal effluent from 95 96 patients was obtained from St Thomas' Hospital Research Ethics Committee (06/Q0702/74) covering Guy's and St Thomas' Hospitals and transferred by agreement to London Bridge 97 Hospital. Where available, clinical information for the colonoscopy patients is given in the 98 text. Samples were collected at Reading (faeces) or transported from St Thomas' Hospital 99 (caecum), and maintained under anaerobic conditions (faeces, MACS1000 anaerobic 100 workstation, Don Whitley Scientific, gas composition 80 % N₂, 10 % H₂, 10 % CO₂; caecum, 101 on ice in a gas jar with an anaerobic gas-generating pack; Oxoid Ltd) for a maximum of 2 h 102 before processing. Caecal samples were collected during routine colonoscopy following 103 preparation of the bowel with sodium picosulphate and a reduced fibre diet for 3 days. Liquid 104 residue in the caecum was aspirated via the colonoscope suction channel into a standard 30 105 ml trap specimen container and immediately transferred to a gas jar. 106

Faecal homogenates (20 %, w/v, from 25 g of faecal material) were prepared in sterile
TBT [0.1 μm filtered prior to autoclaving; 100 mM Tris/HCl, pH 8.0; 100 mM NaCl; 10 mM
MgCl₂.6H₂O] or sterile 0.5 % 'Lab-Lemco'/6 % NaCl (LL [16]). Caecal homogenates (20 %,
v/v, from 10 to 30 ml of caecal effluent) were prepared in LL. Faecal and caecal samples
were placed into a filter stomacher bag and homogenized in a Stomacher 400 Lab System
(Seward) for 120 s at low speed. The bag was removed from the stomacher and massaged

manually to further disperse any large particles remaining in the sample; the sample was then 113 stomached for a further 120 s. The homogenate was kept on ice for 2 h to allow desorption of 114 VLPs from solid material, then centrifuged at 11,180 g for 30 min at 10°C. The supernatant 115 was transferred to a fresh tube and centrifuged again at 11,180 g for 30 min at 10°C. 116 Supernatant [herein referred to as faecal filtrate (FF) for both faecal and caecal samples] was 117 passed through sterile 0.45 µm cellulose acetate filters (Millipore) and the FF was then 118 collected in a sterile container. An aliquot (10 µl) of each of the FFs was examined by 119 epifluorescence microscopy (EFM) after viral particles had been stained with SYBR Gold 120 (see below) to confirm that the samples were free of bacteria. 121 FF prepared in LL was used for enumeration of VLPs in faeces via EFM (see below) 122 and 1 ml aliquots of FF prepared in TBT were for examination by transmission electron 123 microscopy (TEM; see below). 124 Aliquots of FF and LL (two sets of three aliquots of 100 µl each: one for aerobic 125 cultivation, one for anaerobic cultivation) were spread onto Columbia blood agar containing 126 127 5 % laked horse blood (Oxoid), and incubated aerobically and anaerobically. Sterility of the filtrates was confirmed by the absence of microbial growth on plates following incubation for 128 2 (aerobic) and 5 (anaerobic) days. Sterile brain-heart infusion broth was inoculated with 100 129 μl of FF and LL, and incubated aerobically for 2 days: aliquots (100 μl) were spread in 130 triplicate on plates to confirm sterility of the broth cultures. In addition, 10-ml aliquots of FF 131 were stored at 4°C for 6 months after collection, and were found to be free of bacteria when 132 checked (by plating) at monthly intervals. 133

Poly(ethylene glycol) (PEG; BioUltra, 8000; Sigma) was added to 2×20 ml aliquots of the LL-prepared 0.45 µm-filtered FFs (final concentration of PEG, 10 %, w/v). The samples were shaken gently to dissolve the PEG, then left at 4°C for 16 h ([16] used 16 h as they observed an improved recovery of RNA bacteriophage compared to shorter incubations;

[17] used 10 h, as they determined this to be longer than the minimum time required to obtain 138 a constant and stable white layer of viral precipitate). PEG-precipitated VLPs were then 139 harvested by centrifugation at 4,500 g for 30 min in a swing-out-bucket rotor at 4°C. The 140 supernatant was removed and the pellet resuspended in 1 ml TBT for pulsed-field gel 141 electrophoresis (PFGE) or 5 ml TBT for purification of VLPs on a CsCl gradient [15] with 142 ultracentrifugation performed at 100,000 g for 2 h at 10°C in a fixed-angle Type 50 Ti rotor 143 (Beckman Coulter). VLPs recovered by CsCl (1.35 and 1.5 g/cm³ fractions) were dialysed 144 (12,000 Da cut-off) twice against 400 ml sterile TBT, passed through a sterile 0.45 µm 145 cellulose acetate filter and stored at 4°C until DNA was extracted. An aliquot (5 µl) of each 146 of the CsCl-purified samples was viewed using EFM after viral particles had been stained 147 with SYBR Gold (see below) to confirm that the samples were free of bacteria. 148

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2.2 Epifluorescence microscopy (EFM)

A combination of the methods of Thurber and Patel [15, 18] was used to prepare FFs 151 (faecal only) for EFM. VLPs present in FF were not fixed in paraformaldehyde prior to 152 enumeration, as Wen et al. [19] showed that aldehyde fixation leads to a rapid loss in viral 153 abundance. The SYBR Gold concentration of 1-5× recommended by [15] did not work with 154 FFs; therefore, the 400× concentration recommended by [18] for use with planktonic aquatic 155 samples was used in this study. 156

The filtration system (see [15] for specifics of the set-up employed), including a 157 glass-graduated column, for collecting VLPs on filters was cleaned using 5 ml of 0.1 µm-158 filtered, sterile H₂O and 5 ml of 0.1 µm-filtered ethanol. A sterile pair of flat-tipped forceps 159 was used to remove a 0.02-µm white Anodisc 25 membrane (Whatman) from its box, and the 160 filter was fitted to the glass frit of the filtration system under a low vacuum [<10 psi (~62 161 kPa)]. Duplicate 2 ml aliquots of 0.1 µm filtered, sterile H₂O and 10 µl of sterile LL in 2 ml 162

 $0.1 \,\mu\text{m}$ filtered, sterile H₂O were used as negative controls (to assess whether there was any 163 contamination in the water, the LL, the filter tower or the filter surface) and passed through 164 filters under low vacuum before any of the FF aliquots were filtered in duplicate. For each 165 sample, a 10 µl aliquot of 0.45 µm-filtered FF was added to 2 ml of the 0.1 µm filtered, 166 sterile H₂O and the suspension gently mixed. The diluted sample was then introduced into the 167 filter system with the low vacuum maintained. The liquid was passed through the filter, and 168 VLPs were collected on the filter. The clip and glass column were carefully removed from 169 the filtration system and the filter was gently removed from the glass frit (still under 170 vacuum). While holding the filter with a forceps, the back of the filter was gently blotted dry 171 on a clean Kimwipe; the filter was then left to dry on a Kimwipe in a sterile Petri dish for 10 172 min in a dark box. The filtration system was cleaned with 5 ml of 0.1 µm-filtered, sterile H₂O 173 and 5 ml of 0.1 µm-filtered ethanol, and the next sample processed. 174

Filters were stained with 100 μ l droplets of 400× SYBR Gold and dried as described by [18], prior to being applied in pairs to sterile glass slides. The mountant (20 μ l per coverslip) ProLong[®] Gold antifade reagent (Invitrogen) was added to 25 mm glass coverslips. Coverslips were picked up with sterile forceps, inverted and placed on the Anodisc filters on the microscope slides. Slides were then left at room temperature in the dark for 16 h to allow the antifade to cure. Slides prepared in this manner can be stored at room temperature for 1 week or at 4°C for 2 months.

Slides were viewed at 1000× magnification under a Nikon Microphot-SA microscope
fitted with a B-2A (blue excitation) filter and attached to a CoolSNAP-Pro MONOCHROME
(Media Cybernetics Inc.) camera. Images were captured using Image-Pro PLUS version

4.5.0.19 (Media Cybernetics Inc.), and VLPs in 25 fields of view were enumerated by eye.

186 The number of VLPs per millilitre of FF was calculated using the following equation: mean

number of VLPs in 25 fields of view $\times 100 \times 25760.205$; where 100 represents taking the

sample back to per millilitre FF and 25760.205 represents the number of fields of view on a filter. To give an estimate of the number of VLPs per gram of faeces, results were multiplied by a factor of 5. The detection limit of the method was 1.03×10^5 VLPs/ml FF or 5.15×10^5 VLPs/g faeces.

- 192
- 193 **2.3 TEM of FFs (faecal and caecal)**

Aliquots (1 ml) of FF prepared in TBT (faeces) or LL (caecum) were used for TEM 194 195 analysis within 1 week of collection: these filtrates had not been concentrated using PEG. Carbon films (~3×3 mm in size) were floated from mica-sheets into a drop of filtrate (100 196 μ l). After an adsorption time of 5–10 min, samples were transferred into a drop of 1 % (v/v) 197 of EM-grade glutaraldehyde (20 min) and subsequently into a drop of 2 % (w/v) uranyl 198 acetate for negative staining (1-2 min). After two washes for a few seconds in drops of 199 distilled water, samples were picked up with 400-mesh copper grids (Plano, Wetzlar, D). 200 Electron micrographs were taken in a Tecnai 10 transmission electron microscope (FEI 201 202 Company, Eindhoven, the Netherlands) at an accelerating voltage of 80 kV. Digital micrographs were taken with a Megaview G2 CCD camera (Olympus SIS, Münster, 203 Germany). For estimation of VLP titers on the EM grids, a freshly prepared Lactococcus 204 lactis bacteriophage preparation with defined titer and unique morphology (i.e. prolate-205 headed bacteriophage P001 [20]) was added in concentrations of 10⁷ and 10⁸ plaque-forming 206 units (pfu) per ml to the faecal sample of donor 2 (with highest bacteriophage titer according 207 to TEM analysis). 208

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210 **2.4 PFGE**

Aliquots (40 μl) of PEG-precipitated samples resuspended in 1 ml TBT were used for
PFGE. These were heated at 60 or 75°C as appropriate. An aliquot (25 μl) of each heated

(viscous) suspension and loading dye (5 μ l) were combined and loaded into wells, which 213 were sealed with molten agarose. Run conditions for PFGE were as described by Fuhrman et 214 al. [21]: a 1 % agarose gel (SeaKem LE agarose) was made in 0.5× TBE [250 ml of 5× TBE 215 buffer (27 g Sigma 7-9, 13.75 g boric acid, 5 ml 0.5 M EDTA, 500 ml of H₂O) added to 2.25 216 1 H₂O] and a CHEF DR II apparatus (Bio-Rad) was run for 18 h at 6 V and 14°C in 0.5× 217 TBE, with a 1–10 s switch time. Gels were stained with ethidium bromide (5 μ g/ml) for 20 218 min and destained in distilled H₂O for 10 min, or with 1× SYBR Gold (Molecular Probes) 219 220 made in 0.5× TBE. Bands of DNA were visualized under UV light (ethidium bromide) or using a Dark Reader DR89X Transilluminator (Integrated Scientific Solutions Inc.) (SYBR 221 222 Gold).

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224 2.5 Isolation of DNA from CsCl-purified VLPs

DNA was extracted from 500 µl portions of CsCl-purified VLP samples according to 225 [15]. Prior to extracting DNA from purified VLPs, samples were treated with 20 µl of 1 226 mg/mL DNAse I (from bovine pancreas, 552 Kunitz/mg protein; made in water passed 227 through a 0.1 µm filter prior to autoclaving) for 1 h at 37°C, and were then heated at 80°C for 228 10 min to inactivate the DNAse prior to DNA extraction. DNA was air-dried and 229 resuspended in 50 µl of TE buffer. Sterility of samples (i.e. absence of bacterial DNA) was 230 confirmed by negative PCR from samples (10 µl) with universal primers [22]. DNA from an 231 232 in-house strain of *Bifidobacterium longum* (4-FAA1; [23]) was used as a positive control; sterile water was used as the negative control. 233

3. RESULTS AND DISCUSSION

235 **3.1 Examination of VLPs in FFs (faeces) by using EFM**

Previous studies examining the faecal virome have used 0.22 µm filters to prepare 236 samples [9, 11, 12, 13]. Preliminary work performed with 0.45 µm-filtered faecal samples 237 and 0.45 µm-filtered samples originating from *in vitro* fermentation systems inoculated with 238 faeces stained with 4',6-diamidino-2-phenylindole (DAPI) and viewed under an 239 epifluorescence microscope suggested that the abundance of dsDNA VLPs in intestinal 240 samples was high, and that these particles can be enumerated (unpublished observations). 241 Contamination by bacteria was not observed in any of these samples, and confirmed by an 242 absence of cultivable bacteria and no bacteria in TEM analyses, so we decided to continue 243 using larger-pore filters for processing samples. Klieve & Swain [24] had previously used 244 245 0.45 µm-filtered samples to characterize VLPs present in rumen contents, and reported no problems with contamination by bacteria. 246

To examine the VLPs present in FFs (faeces) by EFM, the method described by [9] 247 and [15] was used initially. However, it was found that we were unable to visualize many, if 248 any, VLPs present in the samples using 1-5× SYBR Gold as the DNA/RNA stain. Increasing 249 the concentration of SYBR Gold to 400×, as used by [18] for planktonic aquatic samples, 250 allowed us to enumerate and detect VLPs present in the FFs prepared from faecal samples of 251 six donors (Fig. 1 and Fig. 2). The samples were extracted in LL, though the method worked 252 as well with samples extracted in TBT (data not shown). To determine the relative abundance 253 of the predominant VLP morphologies in FFs, samples were not concentrated by CsCl 254 centrifugation (Fig. 1). All FFs were found to be free of bacteria by EFM (Fig. 2), with the 255 number of VLPs present in samples ranging from 2.4×10^8 to 1.12×10^9 VLPs/ml FF (mean 256 5.58×10^8 VLPs/ml FF) (equivalent to $\sim 1.2 \times 10^9$ to 5.58×10^9 VLPs/g faeces, mean 2.94×10^9). 257 Lepage et al. [4] enumerated VLPs in gut mucosal samples from 14 healthy individuals and 258

19 Crohn's disease patients and found on average 1.2×10^9 VLPs/biopsy (range 4.4×10^7 – 259 1.7×10^{10}), in agreement with the mean value we present here for faecal VLPs. Of note, 260 Crohn's disease patients harboured significantly (P = 0.024) more VLPs than healthy 261 individuals $(2.9 \times 10^9 \text{ vs } 1.2 \times 10^8 \text{ VLPs/biopsy})$ in the study of Lepage *et al.* [4]. 262 It is generally accepted that there are around 10 bacteriophages for every microbial 263 cell in environmental samples investigated to date [25]. Extrapolating this figure to the gut 264 microbiota, from fluorescence in situ hybridization studies it is estimated that the faecal 265 microbiota harbours $\sim 10^{11}$ bacteria/g faeces in healthy adults [26]. Consequently, one would 266 expect the presence of at least 10¹² VLPs/g faeces. Enumerating VLPs in faeces via FFs (or 267 any liquid medium) by EFM is highly subjective as a dot of very intense fluorescence may in 268 fact represent a cluster of VLPs (Fig. 2), a phenomenon frequently encountered during this 269 270 study. Patel et al. [18] stated that, to accurately enumerate VLPs by EFM, micro-adjustments using the fine focus of the microscope have to be made to ensure that all viruses in a 271 particular grid-reticle box are counted. We agree with this statement, and furthermore add 272 that many VLPs have very likely been lost during the preparation and filtering of samples, 273 either by association with debris in the initial centrifugations or by being caught in the filters 274 because of clogging or because the VLPs are too big to pass through the pores (e.g. members 275 of the order *Megavirales* [27]). Even after centrifugation, the supernatants from several of the 276 samples, while appearing relatively translucent, were highly viscous (perhaps due to host 277 mucins) and clogged the 0.45 µm filters with less than 2 ml of sample being filtered. 278 Consequently, the values we provide for the numbers of VLPs in FFs and faecal samples are 279 a conservative estimate. We believe the true number of VLPs present in faeces to be higher, 280 possibly between 10^{10} and 10^{12} VLPs/g faeces. Lepage *et al.* [4] determined there to be 10^{10} 281 VLPs/mm³ tissue in their study of mucosal VLPs. 282

284 **3.2 Examination of VLP diversity in FFs by the use of TEM**

VLPs were readily detected in the TBT-extracted faecal samples from all donors by 285 TEM. Bacterial cells were never observed in any samples examined by TEM (limit of 286 detection 10⁶ per ml). It was striking how visibly different/distinct the VLP assemblages were 287 between the donors, with no two donors sharing the same VLPs, at least on the basis of 288 morphological appearance (Fig. 3 and Fig. 4; Supplementary Fig. 1). The vast majority of 289 290 VLPs present in the samples appear to represent bacteriophages. Donor 1's VLP assemblage was predominated by small and large isometric-headed Siphoviridae with various tail length 291 292 sizes (approx. 120 nm, 350 nm, 650 nm, or, in one extreme case, 1220 nm), with some small and large isometric-headed Mvoviridae also present (Fig. 3a). Notably, donor 2's VLP 293 assemblage was more diverse, predominated by numerous different morphotypes of 294 Myoviridae with Siphoviridae also present (Fig. 3b). Interestingly, two detached Myoviridae 295 tails of extreme length (480 nm) and thickness of the sheaths (40 nm) indicated the presence 296 of giant Myoviridae phages (Fig. 3b). Sime-Ngando et al. [28] have reported isolation of 297 bacteriophages with tails of 400 nm in length (heads 50-130 nm) from hypersaline lake 298 samples, though images of these large bacteriophages are unavailable for direct comparison 299 of the tails' structures with those found in this study. Donor 2's faecal sample contained 300 distinct small isometric-headed Myoviridae phages with uncommon radial fibers (approx 150 301 nm in length) attached to the capsids (~70 nm diameter) (Fig. 4). These fibers are clearly 302 extending the 110-nm tails of these phages, suggesting a primary role in bacteriophage 303 adsorption. This unique bacteriophage morphotype was also documented in low numbers in 304 Donor 4's faecal sample (not shown), and in caecal samples L16 (not shown) and L18 305 (Supplementary Fig. 2d). Fig. 4 also illustrates the apparent clustering of *Mvoviridae* phages 306 in samples. VLPs of various morphotypes derived from Donors 3, 4, 5, and 6 are shown in 307 Supplementary Figure 1a-d. "Zeppelin"-like VLPs of constant thickness (56 nm) but 308

309	different lengths (370-630 nm) were detected in Donor 5 (Supplementary Fig. 1c) and Donor
310	6's samples (Supplementary Fig. 1d). Donor 6's sample was predominated by these
311	"zeppelin"-like VLPs, with only a few bacteriophages detected. Sime-Ngando et al. [28]
312	reported the presence of rods of up to $22-24 \times 1000$ nm: similar to the 'zeppelins' identified
313	in the present study, these rods had no visible internal or external structures. The largest
314	Myoviridae phages found in this study were documented for the sample of Donor 5
315	(Supplementary Fig. 1c), with the phages having prolate heads of 150×115 nm and tails of
316	400 nm in length. The sample from Donor 6 did not reveal the same level of diversity as seen
317	for the other faecal samples and the reason for this is not known (Supplementary Fig. 1d).
318	Enumeration of VLPs in the sample of Donor 2 (highest number of VLPs as detected by
319	TEM) demonstrated there to be approx. 5×10^7 VLPs/ml FF.
320	For the majority of the caecal samples, a lower extent of viral biodiversity was
321	documented. Sample L10 (no clinical data) was unique, containing exclusively small
322	isometric-headed <i>Podoviridae</i> (50 nm diameter) and ~20-nm long appendages (Fig. 5b).
323	Samples L08 (healthy) (Fig. 5a), and L02 and L03 (both IBD) (Suppl. Fig. 5b, c) appeared to
324	exclusively contain Myoviridae phages. The greatest extent of morphological variation and
325	the highest bacteriophage numbers were seen in caecal samples L16 (no clinical data) and
326	L17 (no clinical data) (Fig. 5c, d) and – to a lesser extent – in samples L01 (diverticulosis)
327	and L18 (no clinical data) (Supplementary Fig. 2a, d). A new type of a giant Myoviridae
328	bacteriophage was exclusively found in sample L16 (Fig. 5c) with isometric heads of 125 nm
329	in diameter, with tails of 340 nm in length and unique "curled" tail fibers of \sim 70-nm in
330	length.

Lepage *et al.* [4] demonstrated that gut mucosal samples were predominated by
morphotypes consistent with *Siphoviridae*, *Myoviridae* and *Podoviridae*, and that each

- individual appeared to be colonized by one dominant bacteriophage family. Our TEM resultsfor faecal and caecal VLP assemblages are consistent with these results.
- 335
- 336 3.3 PFGE of VLPs present in FFs

It was clear from the EFM and TEM analyses that, based on the number of VLPs 337 present in our samples, it should be possible to isolate large amounts of VLP-derived DNA 338 339 from faeces. Attempts to recover abundant VLP DNA from samples using the centrifugation method of Thurber et al. [15], and which was used by [9] and [11] to isolate VLP DNA from 340 341 faecal samples, proved disappointing in terms of the recovered DNA yield. However, increasing the centrifugation speed to 100,000 g, routinely used in the laboratory to purify 342 lactococcal bacteriophages for preparation of high-quality DNA and used by Kulikov et al. 343 [29] to recover bacteriophages from horse faeces, markedly improved recovery of faecal 344 VLPs. 345

To concentrate VLPs from 20-ml quantities of FF so that they could be applied to 346 CsCl gradients in 5 ml aliquots, it was decided to use PEG precipitation. This method of 347 recovery has previously been used with, for example, marine samples and faecal samples to 348 improve detection of F-specific coliphages in faecal material [16, 17], and allows large 349 starting volumes of sample to be used for recovering VLPs from human faeces. The method 350 can be scaled easily so that the VLPs from larger volumes of FF are precipitated for 351 collection by centrifugation: we used 20 ml of FF from each donor here to demonstrate the 352 efficacy of the method, but the entire FF derived for each donor's 25 g of starting material 353 could have been used for PEG precipitation of VLPs. 354

PEG-precipitated samples resuspended in 1 ml TBT were examined using PFGE.
Aliquots (40 µl) of the resuspended pellets were heated at 60°C [9] or 75°C ([24] used this
temperature to inactivate nucleases) before PFGE. Profiles containing one or more bands

were observed for samples from all donors (Fig. 6). Following treatment at 60°C, Donor 1's 358 sample produced a faint band at 105 kb, with this band being more prominent following 359 360 sample treatment at 75°C; at the higher temperature, bands were also observed at 135 kb and just below 48.5 kb, with the most prominent band at 79 kb. Donor 2's 60°C-treated sample 361 produced a profile with a faint smear and a thin band at 84 kb; with the 75°C-treated sample, 362 the smear was more pronounced, with a thick band observed between 75 and 84 kb. Similar 363 to Donor 2, the smear in Donor 4's sample became more pronounced following treatment at 364 the higher temperature. For Donor 3's 60°C-treated sample, two strong bands (at 66 and 97 365 366 kb) were observed; in the 75°C-treated samples, bands were observed at 57 kb, 75 kb and 97 kb. The profile of Donor 5 was most affected by changing the temperature at which samples 367 were treated prior to loading on the gel: the 60°C-treated sample had three bands visible 368 (most prominent at 100 kb, fainter bands at 48.5 kb and 66 kb), whereas the 75°C-treated 369 sample had these three bands with at least another 8 bands visible between 48.5 and 97.0 kb. 370 Donor 6's sample had a faint band just below 48.5 kb after heating at 60°C, whereas the 371 sample heated at the higher temperature produced two prominent bands that were smaller 372 than 48.5 kb. The results from the gel were reproducible (triplicate gels run; data not shown). 373 It is unsurprising that heating at 75°C produced more complex banding patterns, as we would 374 expect the higher temperature to disassemble (some of) the capsids of VLPs resistant to 375 heating at 60°C, thereby releasing packaged DNA. 376

Using this method, it was possible to visualize VLP DNA when stained with SYBR Gold (Fig. 6) or ethidium bromide (Supplementary Fig. 3). It was clear that increasing the temperature at which the samples were heated prior to loading onto the gel revealed greater diversity in the samples, and that each individual harboured a unique VLP assemblage (Fig. 4). This is in agreement with the findings of the metagenomic studies of [11] and [12]. The sample from Donor 6 did not reveal the same level of diversity as seen for the other donors, in agreement with the results shown by TEM (Supplementary Fig. 1d). This donor's sample
was predominated by "zeppelin"-like VLPs that may not have released their nucleid acids by
the conditions employed here.

Heating PEG-precipitated samples allowed us to generate PFGE images that 386 demonstrated that each individual harbours a unique VLP assemblage, and encouraged us to 387 continue with studies to improve recovery of VLP DNA from human faeces. It also showed 388 that our method of recovery of VLPs present in faeces was superior to that used previously 389 by [9] to generate a viral assemblage fingerprint. In that study, the authors stated that the 390 391 limited amount of DNA recovered using tangential flow filtration with a 500 g faecal sample from a 33-year-old made it necessary to enhance the sample bands on the gel relative to the 392 ladder. In contrast, no enhancement of the bands detected by SYBR Gold (Fig. 6) or ethidium 393 bromide (Supplementary Fig. 3) was required using the methodology described herein, 394 although SYBR Gold was, as expected, far more sensitive than ethidium bromide. PEG 395 precipitation of VLPs present in 20 ml of FF, as used in this study, equates to the extraction 396 of VLPs from ~4 g faeces (sample losses are seen during filtration, and vary from donor to 397 donor due to the differences in viscosity of faecal supernatant obtained after removal of most 398 bacteria and debris from samples after centrifugation), with 50 µl of the resuspended PEG 399 precipitate representing VLPs isolated from ~200 mg of faeces. 400

The current study used a crude method of extracting DNA for PFGE. Using the method of Rohozinski *et al.* [30], in which VLPs are embedded in agarose blocks, with CsClpurified VLPs prior to heating may allow better 'fingerprinting' of VLPs in faecal samples. This could, for example, be used as an inexpensive means of determining the effect of freeze-thawing faecal samples prior to the recovery of VLPs from faecal samples. All of the metagenomic studies conducted to date on the human faecal virome have used samples that have been frozen prior to recovery of VLPs from samples. 408

409 **3.4 Isolation of high-quality DNA from FFs**

Reves et al. [11] and Thurber et al. [15] (based on the work of [9]) have reported 410 recovery of ~500 ng (from 2–5 g of frozen faeces) and 500–3000 ng (from 500 g of fresh 411 faeces), respectively. The method of Reves et al. [11] is closest to that presented herein for 412 the recovery of VLPs from faeces [i.e. they hand-filtered samples, whereas Breitbart et al. [9] 413 used tangential flow filtration to process their sample]. As stated above, 20 ml of FF equates 414 to ~4 g faeces. We processed 500 µl portions of 3.5-4 ml CsCl-purified VLPs for DNA 415 extraction, and resuspended the DNA in 50 µl of TE (Fig. 7). If we had processed the entire 416 PEG-precipitated, CsCl-purified VLP sample for each donor, we would have recovered 417 between ~1000 and 1800 ng from ~4 g of faeces (theoretically ~6000–11000 ng from 25 g of 418 faeces). This increase in recovery of DNA is thought to be mainly due to the use of 0.45 µm 419 filters rather than 0.22 µm filters to process samples, supported by our observation that 420 421 passing FFs through 0.22 µm filters led to VLP counts by EFM that were approximately half of those of the 0.45 µm-filtered samples. Increasing filter size and using PEG precipitation to 422 increase the volume of sample that can be processed would allow greater sampling of the 423 faecal virome in future metagenomics studies. 424

In addition to demonstrating the utility of the method with faecal samples, we have 425 successfully isolated VLP DNA from human caecal effluent (Fig. 7). For each of the samples, 426 20 ml of 0.45 µm-filtered caecal filtrate prepared in LL was PEG-precipitated, with the VLPs 427 then purified on CsCl gradients. These 20-ml aliquots equated to ~4 ml caecal effluent from 428 each of the three samples examined. We started with 30, 30 and 10 ml, respectively, of 429 430 effluent from caecal samples L07 (healthy), L08 (no clinical data) and L10 (no clinical data). Therefore, if the filtrate from entire homogenates of these samples had been processed, we 431 would have expected to isolate between ~600 and 8000 ng VLP DNA from caecal effluent. 432

- 433 Consequently, it should be possible to conduct metagenomics studies of the VLP
 434 assemblages associated with the human caecum using the methods described herein.
- 435

436 **3.5 Conclusions**

We have presented a series of methods for enumerating and characterizing VLPs 437 present in human faecal and caecal samples. On the basis of enumeration of VLPs using 438 EFM, there are $\sim 10^8$ VLPs/ml FF, with faeces thought contain up to 10^{12} VLPs per gram of 439 sample. TEM analysis of faecal VLPs indicates that an individual is colonized by one 440 441 dominant bacteriophage family, with *Myoviridae* and *Siphoviridae* representing the main families of bacteriophage detected in faeces. PFGE analysis of PEG-concentrated samples 442 has demonstrated that each individual harbours a unique VLP population, and that the 443 temperature at which samples are treated greatly affects diversity observed. Using 0.45 µm 444 filters to prepare samples, it is possible to isolate twice as much DNA as reported previously 445 from ~4 g of faeces. The inclusion of PEG precipitation in the methodology means that VLPs 446 from large volumes of FF can be concentrated, allowing recovery of microgram quantities of 447 VLP DNA from faecal samples. In addition, PEG precipitation and CsCl purification can be 448 used to recover VLPs from human caecal samples, with nanogram quantities of VLP DNA 449 being recovered from the processed samples. 450

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457

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- **Fig. 1.** Number of VLPs detected in FFs by using EFM. Values are shown as mean + sD (n =
- 532 2 per donor). Passing samples through a 0.22 μm filter reduced the number of VLPs present
- 533 in samples by approximately half (data not shown). White bars, per ml FF (faeces); grey bars,
- 534 per g faeces.
- 535
- 536

Fig. 2. Image showing faecal VLPs as they appear under an epifluorescence microscope 536 (×1000 magnification). Bacteriophages appear as 'pinpricks' of light when stained with 537 SYBR Gold. The brightest 'pinpricks' (black arrows) represent clusters of up to 30 VLPs. 538 The black dot (shown in the white circle) on the images is due to a scratch on the microscope 539 lens. (a) Contamination on filter, which gives an idea of the size and appearance of a 540 bacterium in comparison with VLPs. (b) Negative control, 0.1 µm-filtered, autoclaved H₂O. 541 (c) Negative control, sterile 0.5 % 'Lab-Lemco'/6 % NaCl. (d) Donor 1, 0.22 µm-filtered 542 sample; (e) Donor 1 0.45 µm-filtered sample. (f) Donor 2, 0.22 µm-filtered sample; (g) 543 Donor 2, 0.45 µm-filtered sample. Scale bar, 10 µm. 544

- **Fig. 3.** Transmission electron micrographs revealing the diversity of VLPs found in faeces.
- 546 (a) Donor 1 (female, 41 years) and (b) Donor 2 (female, 36 years) after extraction of VLPs in
- 547 TBT buffer.
- 548

- 548 Fig. 4. Transmission electron micrographs of VLPs found in faeces of Donor 2 (female, 36
- 549 years) after extraction of VLPs in TBT buffer showing *Myoviridae* phages with radial
- whiskers attached to the capsids (top) and clusters of *Myoviridae* phages adsorbing to
- 551 membrane vesicle material (bottom).
- 552

- **Fig. 5.** Transmission electron micrographs revealing the diversity of VLPs found in caecal
- effluents. Sample (a) L08, (b) L10, (c) L16 and (d) L17 after extraction of VLPs in LL. No
- clinical data were available for these individuals.

- **Fig. 6.** Use of PFGE to examine VLP populations in PEG-precipitated FFs (faeces), and
- demonstration that increasing the temperature at which samples are heated prior to loading

onto the gel can affect the diversity uncovered. The gel was stained with SYBR Gold and

visualized as described in Methods. This is a crude (but inexpensive) method of examining

- 559 VLP populations in human faeces, and demonstrates that each individual harbours a unique
- 560 VLP profile. Ladder, lambda ladder (#340; New England Biolabs).

561	Fig. 7. Isolation of high-quality DNA from CsCl-purified samples of human gastrointestinal
562	VLPs. VLP preparations were CsCl-purified and DNA extracted as described in Methods.
563	Aliquots (10 μ l) of sample were run on a 0.8 % agarose gel at 90 V for 30 min. Lane 1,
564	molecular size ladder; lane 2, ϕ KPLN1 positive control (a bacteriophage isolated from the
565	human caecum that infects Klebsiella pneumoniae subsp. pneumoniae K2 strains; L. Hoyles,
566	unpublished data), 63 ng DNA/ μ l; lane 3, VLP extract from faeces of Donor 4, 3.1 ng
567	DNA/ μ l; lane 4, VLP extract from faeces of Donor 3, 4.5 ng DNA/ μ l; lane 5, VLP extract
568	from caecal sample L07 (healthy), 0.2 ng DNA/ μ l; lane 6, VLP extract from caecal sample
569	L08 (no clinical data), 2.7 ng DNA/ μ l; lane 7, VLP extract from caecal sample L10 (no
570	clinical data), 3.8 ng DNA/ μ l. The two black arrows highlight the presence of RNA in the
571	sample from Donor 3, confirmed by treating the CsCl-purified sample with 20 μ l of 1 mg/mL
572	RNAse A (bovine pancreas, \geq 70 Kunitz/mg protein, prepared in 0.1 µm-filtered, sterile H ₂ O)
573	with the DNAse in a second DNA extraction (not shown). VLPs and DNA were extracted
574	from human caecum samples (1:4 LL, v/v) as described in Methods as part of a study
575	examining the microbiota associated with the human caecum (L. Hoyles, unpublished data).
576	With the exception of the sample in lane 5, DNA visible to the naked eye was extracted from
577	all samples of gastrointestinal origin.