1	Biology and taxonomy of crAss-like bacteriophages, the most abundant virus in the
2	human gut
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17 Abstract

CrAssphage is yet to be cultured even though it represents the most abundant virus in 18 the gut microbiota of humans. Recently, sequence based classification was performed on 19 20 distantly related crAss-like phages from multiple environments, leading to the proposal of a familial level taxonomic group [Yutin N, et al. (2018) Discovery of an expansive 21 22 bacteriophage family that includes the most abundant viruses from the human gut. Nat 23 Microbiol 3(1):38–46]. Here, we assembled the metagenomic sequencing reads from 702 24 human faecal virome/phageome samples and obtained 98 complete circular crAss-like phage 25 genomes and 145 contigs \geq 70kb. In silico comparative genomics and taxonomic analysis was performed, resulting in a classification scheme of crAss-like phages from human faecal 26 27 microbiomes into 4 candidate subfamilies composed of 10 candidate genera. Moreover, 28 laboratory analysis was performed on faecal samples from an individual harbouring 7 distinct 29 crAss-like phages. We achieved propagation of crAss-like phages in ex vivo human faecal 30 fermentations and visualised Podoviridae virions by electron microscopy. Furthermore, 31 detection of a crAss-like phage capsid protein could be linked to metagenomic sequencing 32 data confirming crAss-like phage structural annotations. 33

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42 Significance

CrAssphage is the most abundant biological entity in the human gut, but it remains 43 uncultured in the laboratory and its host(s) is unknown. CrAssphage was not identified in 44 45 metagenomic studies for many years as its sequence is so different from anything present in databases. To this day, it can only be detected from sequences assembled from metagenomics 46 47 or viromic datasets (crAss – cross Assembly). In this study, we identified 243 new crAss-like phages from human faecal metagenomic studies. Taxonomic analysis of these crAss-like 48 phages highlighted their extensive diversity within the human microbiome. We also present 49 50 the first propagation of crAssphage in faecal fermentations and provide the first electron micrographs of this extraordinary bacteriophage. 51

53 Introduction

In recent years, increasing numbers of bacteria, archaea, fungi, protists and viruses 54 residing on and within the human body have been associated with various states of human 55 56 health and disease, including diet, age, weight, inflammatory bowel disease (IBD), diabetes, and cognition (1-7). A relatively small number of eukaryote viruses present in the 57 gastrointestinal tract can target the human host, however, much larger and much more 58 59 complex populations of viruses that target bacteria (bacteriophages) also reside there. The role of phages in the gut has been a subject of increased interest as initial investigations have 60 revealed substantial differences in bacteriophage populations between healthy and diseased 61 cohorts (7–11). It is likely that phages have an important role in shaping our gut microbiome, 62 but their precise role remains poorly understood. 63

64 In 2014, metagenomic studies of the viral fraction of the human gut microbiota 65 identified a DNA phage, crAssphage, detectable in approximately 50% of individuals from specific human populations and reaching up to 90% of the total viral DNA load in faeces of 66 67 certain individuals (12). Dutilh and colleagues noted that crAssphage had been overlooked in previous metagenomic studies as the vast majority of its genes do not match known 68 sequences present in databases. It has been predicted, based on indirect evidence using host 69 co-occurrence profiling, that prototypical crAssphage infects *Bacteroides*, an abundant genus 70 71 of bacteria important for the normal gut function of humans. However, since crAssphage has 72 never been isolated in culture, its host range, replication strategy, virion morphology and impact on the human gastrointestinal microbiota remains unknown. Thus, a better 73 understanding of crAssphage is crucial to understanding phage host dynamics in the human 74 75 gut microbiota.

Originally crAssphage was published as an individual phage following crossassembly of several metagenomic samples (12). Analysis by Manrique *et al.*, of the healthy

78 human gut phageome identified 4 circular crAssphage genomes and several related 79 incomplete contigs (10). PCR amplification and sequencing of the crAssphage polymerase gene by Liang and colleagues similarly demonstrated diversity amongst crAssphage-80 81 positive faecal samples (13). Recently, Cinek et al. described updated PCR primer sequences for the detection and evaluation of crAssphage diversity, while Stachler et al. 82 developed their own primers targeting conserved genomic regions to evaluate the 83 84 abundance of crAssphage as an indicator of human faecal pollution (14, 15). Finally, an epidemiological survey of crAssphages conducted by Dutilh, Edwards and colleagues has 85 86 suggested crAssphage is associated with humans and primates globally with significant diversity (manuscript currently in preparation). 87

A recent study provided the first detailed sequence-based taxonomic categorisation 88 89 of crAss-like phages, proposing a novel familial level taxonomic group that would include 90 crAssphage itself, as well as various related bacteriophages, from multiple environments (16). However, the authors noted that this classification is in contrast with the classical 91 92 viral taxonomy scheme currently in use. Such taxonomy strictly categorises crAssphage as a member of the *Podoviridae* family. Previous attempts to reconcile sequence-based and 93 classical viral taxonomy have proposed Podoviridae sharing >40% orthologous protein-94 coding genes be grouped at the taxonomic rank of genus, while phages sharing only 20-95 96 40% orthologous protein-coding genes should be grouped at the higher taxonomic rank of 97 subfamily (17). Other reports describe a phage genus as a cohesive group of viruses sharing >50% nucleotide sequence similarity (18). As crAssphage is not a single entity, but 98 rather a group of crAss-like phages that share similarity with the prototypical crAssphage 99 100 at various levels, a comparative analysis of crass-like phage sequences is required to enable detailed taxonomic characterisation. 101

102 In this study, we combine several *in silico* and *in vitro* approaches to further explore the diversity of crAss-like phages in the human gut, and better understand their biological 103 properties. We performed an in-depth analysis of crAss-like sequences from a number of 104 105 previously published and unpublished human faecal virome datasets (1, 7, 9, 10, 19). Subsequent to the assembly of metagenomic sequencing reads, crAss-like phage contigs were 106 107 identified using conserved genetic signatures. In total, 98 complete circular and 145 nearcomplete (≥70kb) linear contigs of crAss-like phages were identified for genomic and 108 109 taxonomic analyses. Laboratory analysis of crAss-like phages was focused on a human donor 110 identified as a stable carrier of several highly predominant crAssphage-like DNA sequences, including one closely related to the prototypical crAssphage. Ex vivo faecal fermentations 111 112 enabled the amplification of a virus highly related to the prototypical crAssphage, with 113 electron micrographs supporting the proposal that crAss-like phages are members of the *Podoviridae* family. These results represent the first example of biological characterisation of 114 this highly prevalent and, potentially, very important human microbiome virus. 115

116 **Results**

Detection of crAss-like phage contigs. Following the assembly of 702 human faecal 117 virome/phageome metagenomic samples listed in Supplementary Table 1, contigs were 118 119 screened for relatedness to the prototypical crAssphage virus, henceforth referred to as crAssphage sensu stricto. Initially, the polymerase of crAssphage sensu stricto (UGP_018, 120 NC_024711.1) was used for crAss-like phage detection due to its use in several studies as a 121 122 genetic signature to determine diversity of crAss-like phages (13, 20, 21). However, we 123 extended our criteria in order to include partial genomes (\geq 70kb) that may not have included 124 the polymerase gene in the assembly. Therefore, after an initial detection of crAss-like phages 125 using the polymerase sequence, we identified the most conserved crAss-like phage protein in 126 our dataset as the terminase protein, encoded by crAssphage sensu stricto UGP_092. The 127 terminase was subsequently used as a second genetic signature for identifying crAss-like 128 phage contigs.

Initially, 239 contigs \geq 70kb were detected with similarity to crAssphage *sensu stricto* 129 130 polymerase sequence. An additional 59 contigs \geq 70kb were subsequently detected with relatedness to crAssphage sensu stricto terminase sequence. Following an initial examination 131 of the contig sequences retrieved, more stringent parameters were implemented. Only contigs 132 whose polymerase and/or terminase sequence(s) aligned with greater than 350bp were 133 134 considered for further analysis as crAss-like phages. This reduced the total number of crAss-135 like phages to 256. In addition, as several assembled metagenomic samples were from the 136 same person sequenced at multiple time points, redundant contigs were removed from further analysis. When two or more contigs aligned with 100 percent identity, the longer contig or the 137 138 contig with the highest coverage was retained. This resulted in a total of 244 crAss-like contigs (including crAssphage *sensu stricto*), with 143 contigs containing both a polymerase 139 140 and terminase, 60 a polymerase only and 40 a terminase only. Of the 244 crAss-like phage

141 contigs, metadata was available for the majority of their originating faecal samples. CrAss142 like phages were detected in healthy individuals across a wide age range (including infants 1
143 year of age and individuals ≥65 years of age) and individuals suffering from Crohn's disease,
144 ulcerative colitis, cystic fibrosis, kwashiorkor and marasmus.

145 Taxonomy of crAss-like phages. In order to compare the phylogeny of the more distantly related phages proposed to be included into a crAss-like familial level taxon by 146 Yutin et al. (16) with those identified in this study, a phylogenetic tree of conserved crAss-147 like phage terminase sequences was constructed (Supplementary Figure 1). Amino acid 148 149 terminase sequences were used to generate mid-point rooted phylogenetic trees. Predominantly, the terminase sequences of very distant crAss-like phage relatives identified 150 by Yutin et al. from various environmental sources were distinct from the various candidate 151 152 genera of crAss-like phages observed in the phylogram. However, the human gut microbiome phage, IAS virus (16), characterised by Yutin et al. as crAss-like, clustered closely with 153 candidate genus VI crAss-like phages identified in this study. 154

Previously, studies have used the percentage of shared homologous proteins as a means 155 of defining phage taxonomic ranks (17). Therefore, clusters of phages sharing between 20-156 40% of their protein-coding genes were categorised as related at the subfamily level, while 157 phages sharing >40% protein-coding genes were grouped at the genus level. A heatmap based 158 159 on the percentages of shared orthologous proteins suggests that crAss-like phages form 4 160 candidate subfamilies. The four subfamilies were assigned the nomenclature alphacrAssvirinae (which contains crAssphage sensu stricto), betacrAssvirinae (which 161 contains IAS virus), gammacrAssvirinae and deltacrAssvirinae (Figure 1). These subfamilies 162 163 can be further subdivided into 10 candidate genera, with Candidate Genus I containing crAssphage sensu stricto and Candidate Genus VI containing the IAS virus. Metadata of all 164

165 crAss-like phages analysed in this study, including their categorisation into the various
166 taxonomic divisions, is available in Supplementary Table 2.

An alternative approach for characterising the encoded proteome of crAss-like phages 167 was performed by visualisation of genome clusters using the t-SNE machine learning 168 algorithm with Euclidean distances of orthologous genes distribution between genomes as an 169 input. Applying the previously determined 10 crAss-like phage candidate genera 170 171 classifications to the t-SNE two-dimensional ordination demonstrated that some clusters showed uniformity while others groups were quite dispersed, such as Candidate Genus II and 172 173 VII, respectively (Figure 2A). In addition, no single cluster of crAss-like phages is exclusively associated with healthy or diseased individuals. 174

Groups of crAss-like phages with a similar G+C nucleotide content would be expected 175 176 to infect related bacteria, since phage G+C content often aligns to that of its host (22, 23). Therefore, several groups of crAss-like phages, such as candidate genera II, IV, V, VII and X, 177 are likely infect closely related bacterial taxa within the human microbiome (Figure 2B). 178 Candidate genus I is the most homogenous group of crAss-like phages containing crAssphage 179 sensu stricto and 30 additional complete circular genomes and 29 linear contigs \geq 70kb with a 180 distinct G+C nucleotide content (29.11 \pm 0.14%). Candidate genera III and VI display the 181 greatest heterogeneity, with G+C contents of 28.94 $\pm 3.03\%$ and 35.81 $\pm 2.56\%$, respectively. 182

Nucleotide comparison of crAss-like phages. To further investigate the relatedness of crAss-like phages, a more detailed comparison at the nucleotide level was performed by calculating their average nucleotide identity (Figure 3). Candidate genera III and VI of crAsslike phages, as defined by the percentage of their shared encoded proteins, also do not cluster into clearly definable groups based on nucleotide composition. Candidate Genus I, containing crAssphage *sensu stricto*, forms a well-defined homogenous taxonomic group even when analysed at the higher resolution of nucleotide composition. This is to be expected as

crAssphage *sensu stricto* was the starting point for finding all crAss-like phages examined in
this study and thus has the most sequences available for analysis.

Interestingly, the majority of crAss-like candidate genera demonstrate the same type of 192 193 genomic organization (Supplementary Figure 2). Prominent features were shared between 194 candidate genera I – V, IX, and X. These include; circular genomes with size ranging from 92 to 104kb, two clearly separated genome regions with opposite gene orientation and inversed 195 196 G+C skew (the smaller region encodes proteins involved in replication, the bigger region coding for proteins involved in transcription and virion assembly, as suggested by Yutin et 197 198 al.), the presence of giant open reading frames with sizes up to 15kb (UGP_052, UGP_053, 199 UGP 052 in the genome of crAssphage *sensu stricto*), possibly coding for fused subunits of 200 RNA polymerase (16), as well as an absence or scarcity of tRNA genes. By contrast, 201 members of candidate genus VI had two genome regions of approximately equal size with 202 opposite gene orientation and G+C skew and large sets of tRNA genes (up to 27; Supplementary Table 2). A prominent common feature of the members of candidate genera 203 204 VII and VIII was absence of the giant open reading frames.

In order to further demonstrate the homogeneity of the candidate genus I of crAss-like 205 206 phages, comparative genomic analysis was performed on complete genomes. We characterised crAss-like phages as having pac-type circularly permuted genomes (24, 25); 207 208 therefore, only genomes determined as circular were considered for this analysis. The 209 genomic start coordinates of circular Candidate Genus I crAss-like phages were altered to 210 match that of the published prototypical crAssphage sensu stricto. Candidate Genus I crAss-211 like phages showed high levels of synteny and strong homology across their entire genomes. 212 However, the most notable area of diversity is observed in the crAss-like phage putative receptor binding protein (UGP_074), which likely targets the different crAssphage strains 213 214 towards their specific bacterial hosts (Supplementary Figure 3).

Prevalence of crAss-like phages in human faecal virome samples. To get insights into relative abundance of different crAss-like phages in various human populations we aligned quality filtered reads, representing 532 human faecal samples from the same datasets as used for assembly of crAss-like genomes, to a database of 93 nonredundant crAss-like phage genomic sequences (with <90% of homology and/or <90% overlap between them) representing all 10 candidate genera.

Crass-like phage colonization rates varied from 51-58% in Malawian infants to 98-221 222 100% of healthy individuals of various ages in the Western cohorts. While relative crAss-like 223 phage content ranged from 0 to 87% of the reads per sample, and depended significantly on the country of residence (p = 6.5E-09 in Kruskal-Wallis test) and age group of the donor (p =224 225 1.6E-10). In ~8% of all virome samples, >50% of reads aligned to crAss-like phage genomes. 226 Lowest overall crAss-like phage counts were seen in healthy Irish and Malawian infants and 227 in USA adults with IBD (Figure 4A). On a global scale, crAss-like candidate genera I, III, and VIII seem to be the most prevalent ones (Figure 4B). 228

229 The specific composition of crAss-like phages in faeces partly separated a cohort of 230 healthy and malnourished infants living in rural areas of Malawi from the healthy and diseased urban Western cohorts (Figure 4C). PERMANOVA analysis suggested that crAss-231 like phage composition was mostly driven by place of residence ($R^2 = 0.24$, p = 0.001) with 232 condition and age group also having significant impact ($R^2 = 0.05$ and 0.01 respectively, p =233 234 (0.001). This observation is further supported by a clear difference in the distribution of specific crAss-like candidate genera across different populations (Figure 5). Specifically, 235 236 Candidate Genus I, which includes crAssphage *sensu stricto* is by far the most prevalent type 237 of crAss-like phages in Western population regardless of age. At the same time, same genus was extremely scarce in Malawian cohort where Candidate Genus III and VIII were the most 238 239 common (p = 6.7E-03 and 1.4E-06, respectively).

240 Faecal fermentations of a crAssphage rich sample. During an ongoing longitudinal study of faecal viromes in healthy adults we identified one individual (subject ID 924), in 241 which crAssphage *sensu stricto* was consistently contributing >30% of virome metagenomic 242 243 reads over a 12 month period. Thus, this donor was selected in order to investigate if crAssphage sensu stricto could be propagated in a batch faecal fermentation system. 244 Quantitative PCR (qPCR) detection of a conserved fragment of the crAssphage sensu stricto 245 DNA polymerase gene in the viral nucleic acid fractions throughout the fermentation revealed 246 that crAssphage sensu stricto was effectively propagated. CrAssphage sensu stricto was 247

found to increase in titre by 89 fold for up to 21 hours into the fermentation (Figure 6A).

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Interestingly, shotgun metagenomic sequencing of the viral enriched DNA from the 249 250 fermentation supernatants showed the presence of six other crAss-like phages in the study 251 subject, in addition to crAssphage sensu stricto (Supplementary Table 2). These crAss-like phage contigs were all \geq 70kb and grouped into five of the candidate genera (Figure 6B), four 252 of which contributed to $\geq 1\%$ of the reads per sample. The most abundant crAss-like contig of 253 254 subject ID 924, designated as Fferm ms 6 (linear, 90.4kb), is a member of proposed Candidate Genus I and closely related to crAssphage sensu stricto. Contig Fferm ms 2 255 256 (linear, 88.8 kb) is the second most abundant in the sample and belongs to Candidate Genus V. Other crAss-like phages showed varying degrees of similarity at the amino acid level to 257 258 different crAss-like phage at the genus-level taxonomic groups. Analysis of bacterial 259 microbiota in the fermentation vessel using compositional 16S rRNA gene amplicon sequencing revealed a concomitant increase in the course of fermentation of a number of 260 Bacteroides species, including; B. dorei, B. uniformis, B. fragilis, B. xylanisolvens, B. nordii, 261 262 Parabacteroides distasonis and Parabacteroides chinchillae (Supplementary Figure 4).

263 **Biological characterisation of crAss-like phages.** Transmission electron microscopy 264 (TEM) of a crAssphage *sensu stricto* rich faecal filtrate showed a significant presence of

265 short-tailed or non-tailed viral particles with icosahedral or isometric heads (53% of *Podoviridae* type and 29% of *Microviridae* or a smaller type of *Podoviridae*), with lower 266 levels of tailed bacteriophages of the family Siphoviridae (15%; Figure 7A). Podoviridae-267 268 type virions could be further classified into two types: type I, with head diameters of ~76.5 nm and short tails; and type II, with a similar head size but head-tail collar structures and 269 slightly longer tails (Figure 7B). Sequencing of the same fraction as used for the TEM 270 271 showed that approximately 40% of reads aligned to crAss-like genomic contigs (Figure 7D). Based on the size of crAss-like genomic contigs assembled from subject ID 924 samples 272 273 (88.8-97.3 kb), it seems likely that the predominant Podoviridae morphology observed corresponds to the crAss-like group of bacteriophages. For comparison, *Microviridae* phages 274 have genomes 4.4-6.1 kb and icosahedral capsids of approx. 15-30 nm in diameter (26, 27). 275

276 The same CsCl fraction that was subjected to metagenomic sequencing and TEM visualisation was also analysed by SDS-PAGE followed by identification of major bands 277 using MALDI-TOF mass spectrometry. A major structural protein of a crAss-like phage, 278 279 denoted as Fferm ms 2 MCP, was detected following MALDI-TOF analysis of a band excised from the ~55kDa area on a SDS-PAGE gel (Figure 7C). The obtained peptide profile 280 corresponded to a protein of 490 amino acids and 55.4 kDa, encoded by Fferm_ms_2. Further 281 analyses using BLASTp showed the protein to have 37% identity with UGP_086, predicted 282 283 as the major capsid protein of the prototypical crAssphage (16).

In addition, we attempted to independently establish the size of crAss-like phage virions by passing faecal filtrates through a series of filters with gradually decreasing pore sizes (Supplementary Figure 5). Filtration through 0.1 µm pores (equivalent to 100 nm) resulted in partial retention of crAss-like phages while pores of 0.02 µm size completely removed crAssphage from the filtrate, as judged by the qPCR assay.

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290 Discussion

The overall objective of this study was to gain a more in depth insight into one of the 291 most enigmatic phages discovered to date, crAssphage. This phage is highly abundant in the 292 293 human microbiome on a global scale; however, it remains poorly understood. One reason 294 why crAssphage has remained such a mystery is due to the lack of available genome sequences for comparison. When crAssphage was assigned a specific nomenclature and 295 296 uploaded to a public repository by Dutilh and colleagues (12), it became a template for other studies to compare against. This highlights the need for researchers to upload both the 297 298 sequencing reads and assembled contigs following metagenomic studies.

CrAssphage is a representative of an expanding group of human gut-associated 299 300 bacteriophages. While previous studies have proposed a sequence-based classification of 301 crAss-like viruses at the familial level (16), our in silico analysis fits within classical familial 302 taxonomic assignments whereby crAss-like phages are categorised as Podoviridae. In this study, we present 243 new crAss-like phage genomes from various metagenomic studies. 303 304 Comparative genomics of the 244 available crAss-like phages demonstrates an extensive degree of diversity among these phages, including the potential identification of four crAss-305 306 like phage subfamilies. While the *alphacrAssvirinae* subfamily is currently the largest of the 4 subfamilies, future studies looking for additional homologues of betacrAssvirinae, 307 308 gammacrAssvirinae and deltacrAssvirinae members will refine these taxonomic categories.

Assigning phage taxonomy, in the absence of a universal genetic marker such as 16S rRNA, is a difficult and potentially erroneous process. In our study, we adopted a method previously employed to assign taxonomic ranks to *Podoviridae* based on the percentage of shared homologous proteins (17). This categorisation strategy identified 10 candidate genera, with crAss-like phages in each genera originating from the faeces of putatively healthy individuals and people suffering from various diet and bowel-related disorders. Alternative

315 proposed methods for defining phage genera include grouping phages with >50% nucleotide 316 similarity identity together (18). Noteworthy, the 10 proposed crAss-like phage genera as 317 determined by percentage of shared homologous proteins closely resembles that observed for 318 crAss-like phage groups when characterised by >50% shared average nucleotide identity.

Several crAss-like phage genera proposed in this study have distinct nucleotide G+C compositions. The nucleotide composition of obligate parasites, such as phages, likely evolves in close association with the host bacterium (23, 28–30). Thus, Candidate Genera III and VI with diverse G+C compositions are either heterogeneous groups of crAss-like phages that require further sequences to refine their taxonomic structure, or they are potentially capable of infecting across a broad host range.

Quantitative analysis of crAss-like phage content in several cohorts revealed that in 325 326 agreement with the previous studies the vast majority of faecal viral metagenomic samples 327 contained varied amounts of crAssphage DNA. CrAssphage sensu stricto (Candidate Genus I) is by far most predominant type in Western populations, co-existing with other crAss-like 328 phages in the majority of samples. By contrast, in the cohort of malnourished and healthy 329 Malawian infants (9, 31), other candidate genera such as III, VIII and IX seem to play the 330 leading role. It is well known that non-Western rural populations, which mostly consume high 331 fibre, low fat and low animal protein diet are predominantly associated with high 332 333 Prevotella/low Bacteroides type of gut microbiota (known as enterotype II (32)), as opposed 334 to Bacteroides/Clostridia-dominated microbiota (enterotype I) in urban populations consuming western diet (33, 34). Indeed, our analysis of the Reyes et al. (2015) 16S rRNA 335 336 gene sequencing data confirmed high prevalence of Prevotella in Malawian samples 337 (Supplementary Figure 6). One can hypothesize that members of candidate genera III, VIII and IX might be associated with Prevotella or other members of the order Bacteroidales apart 338 339 from Bacteroides sensu stricto.

340 The *in vitro* analysis of samples obtained from subject ID 924 was particularly intriguing. By mapping metagenomic sequencing reads against crAssphage sensu stricto, it 341 was initially thought that this donor only carried the prototypical crAssphage at levels 342 exceeding 30% of total viral reads for a 1 year period. A subsequent mining for phages 343 344 related to crAssphage sensu stricto using metagenomic sequencing at later time points, with and without multiple displacement amplification resulted in 5 additional crAss-like phages 345 346 being simultaneously detected from a single donor. However, the initial screening and inclusion criteria for bioinformatic detection of crAss-like phages resulted in a fragmented 347 348 crAss-like phage contig being missed. The overlooked crAss-like phage, Fferm_ms_2 (Candidate Genus V), turned out to be extremely important during the *in vitro* biological 349 characterisation experiment. Therefore, it is possible many additional crAss-like phage 350 351 genomes could be present within the metagenomic datasets that were examined in this study, 352 but they were not included in our analysis because of the inclusion criteria chosen or even the choice of assembly program. 353

In total, subject ID 924 consistently carried 7 crAss-like phages, which resolved in our 354 taxonomic analysis into 5 candidate genera. Three of the crAss-like phages were identified in 355 Candidate Genus VI, supporting the notion this is a heterogeneous group and not simply 356 composed of broad host range infecting phages. It is possible that there are potentially more 357 358 than 7 crAss-like phages within subject ID 924. However, we believe that only a single 359 representative of each candidate crAss-like phage genus (with the exception of the heterogeneous candidate genus VI) could assemble correctly, with two or more highly 360 361 identical phages amalgamating their single nucleotide polymorphisms into a single consensus 362 representative sequence (Supplementary Figure 7).

This study demonstrates the proliferation of crAss-like phages in a faecal fermenter, the first evidence of crAss-like phage propagation in the laboratory. Furthermore, following our

365 ability to propagate faecal crAss-like phages, we conducted the first transmission electron micrographs (TEMs) of these phages. Indeed, the most abundant faecal viruses present in 366 samples used to inoculate faecal fermentation were *Podoviridae*. This is in agreement with 367 368 the predictions made by Yutin et al., following their detailed genome annotation of two crAss-like phages (16). Interestingly, however, our TEMs suggest presence of two types of 369 virions with short non-contractile tails (Figure 7C). Presumably, the more abundant type I 370 371 virions with shorter tail can belong to members of Candidate Genus I, also found as the most abundant crAss-like phage group in subject ID 924 by means of metagenomic sequencing 372 373 (Figure 6B). Whereas type II virions with slightly longer tails and visible head-tail collar structures may correspond to Candidate Genus VI, found as the second most abundant crAss-374 375 like phage subfamily in shotgun metagenomics. But without isolating these phages in pure 376 culture, it is not possible to accurately assign which *Podoviridae* tail corresponds to which specific crAss-like phage subfamily or genera. 377

This work provides the first *in vitro* evidence confirming that crAss-like phages are 378 379 members of the *Podoviridae* family. This is shown from three levels of experimentation using the same CsCl fraction purified from crAssphage rich faeces of a healthy human donor. The 380 TEM images produced from the CsCl fraction showed an abundance of the signature 381 Podoviridae morphology. Other phage capsids present, predominantly Microviridae, would 382 383 typically be associated with smaller genome sizes than that of crAss-like phages (26). 384 Sequencing of the same fraction identified that almost 40% of the reads aligned to crAss-like phages. This is consistent with the percentage of *Podoviridae* identified in the TEM images. 385 Furthermore, a highly predominant protein denoted as Fferm_ms_2_MCP, was isolated from 386 387 the fraction and was found to have significant similarity to crAss-like phages of (Candidate Genera V) as well as a moderate degree of similarity to crAssphage sensu stricto (Candidate 388

Genera I). This *in vitro ev*idence, in line with the taxonomic analysis performed by Yutin *et al.*, proves that crAss-like phages do indeed belong to the *Podoviridae* family.

Identifying a means of propagating crAss-like phages is of particular importance. 391 392 However, it was also observed that the primers applied in the qPCR analyses of viral nucleic acids were not suitable for targeting crAss-like phages associated with the various 393 subfamilies and candidate genera that differed significantly from crAssphage sensu stricto. 394 395 With the availability of more crAss-like phage sequences, broad and narrow spectrum primers can now be designed and applied in the analysis of these phages. The choice of 396 397 primers for detecting crAss-like phages was also discussed in the recent work of Cinek et al. (14). This will be an important part of further work. 398

It also has to be considered that human gut crAssphage is not one single entity, but 399 400 rather a group of diverse viruses, sharing certain signature genomic traits. It is most likely 401 that these diverse phages target multiple bacterial taxa. Previously, a member of the *Bacteroides* genus was hypothesised as being the host for crAssphage (12). In a study prior to 402 403 the discovery of crAssphage (35), a 95.9kb contig corresponding to a putative virus φ HSC05 was shown to be stably engrafted after transplantation of human faecal virus fraction into 404 405 germ-free mice colonized with an artificial defined community of 15 bacterial species. The artificial bacterial community, among others, included: Bacteroides thetaiotaomicron (2 406 407 strains), B. caccae, B. ovatus, B. vulgatus, B. cellulosilyticus and B. uniformis. One might 408 conclude that one of the above mentioned 7 strains of the genus Bacteroides, more likely than the remaining 8 strains of Gram-positive anaerobic bacteria used in that study, must have 409 served as a host for crAssphage propagation. The retrospective analysis of contigs from that 410 411 study conducted by ourselves showed that the oHSC05 contig was 91.73% identical by its 412 nucleotide sequence to crAssphage sensu stricto. Since crAssphage had not been described at the time the article was published, this very interesting observation was never made by theauthors of the original work.

With more divergent sequences, we could assume that different members of the 415 416 Bacteroides genus, or even Bacteroidetes phylum for example, may serve as hosts for different crAss-like phages. One host that has been hypothesised for prototypical crAss-like 417 phages is B. dorei. This was inferred following the analysis of a dataset generated from 418 419 infants and toddlers with islet autoimmunity. It was correlated that crAssphage was only present when B. dorei also was detected within the samples. This was not true for other 420 421 Bacteroides members tested, including B. vulgatus which is highly related to B. dorei. This 422 correlation is compelling; however, it should be noted that there was no confirmation that 423 crAssphage has any role in causing bacteriome alterations that lead to islet autoimmunity 424 (36). Interestingly, one of the key Bacteroides species detected from our faecal fermentation 425 16S rRNA analysis was B. dorei. Its levels were inversely proportional to that of crAssphage. Therefore, this possible phage-host pair should be investigated further. 426

427 CrAss-like phages have also been defined as a part of the core human gut phageome (10). This emphasises the importance of identifying hosts for diverse crAss-like phages 428 429 belonging to different candidate genera proposed in this study. Such knowledge along with the ability to propagate crAss-like phages in vitro will provide an insight into its biological 430 431 significance including their possible role in shaping the bacterial composition of the human 432 gut microbiome in a positive or negative manner, in context of various disease states, such as 433 inflammatory bowel disease, cancer, and obesity among others. Thus far, only a few studies has attempted to correlate crAss-like phages with a gastrointestinal disorder (7, 13, 36). 434 435 Exploring this aspect of crAss-like phages further will be a key part of future work.

In conclusion, our results expand the repertoire of known crAss-like phagessignificantly, providing a path towards the identification of further crass-like phages and their

438 hosts. This will lead to a better understanding of their role, if any, in human health and disease. Our work also provides an interesting insight into the diversity of these human gut-439 associated phages in various populations through in silico and in vitro methods. In addition, 440 441 we also demonstrate that these enigmatic phages can be efficiently propagated *in vitro* in a mixed culture as well as present the first TEMs of crAss-like phages, giving an insight into 442 their morphology. CrAss-like phages appear to be universally present in human populations, 443 including various disease states. Due to the specificity of phage-host interactions, the 444 diversity of crAss-like phages suggests they infect multiple diverse bacteria of the human 445 446 gastrointestinal microbiota. However, more studies will be required to determine the biological significance and role of crAss-like phages in the human gut and determine if its 447 448 presence positively negatively impacts human gastrointestinal health. or

449 Methods

Metagenomic datasets and contig assemblies. Sequencing reads from publicly 450 available metagenomic datasets were downloaded from NCBI Sequence Read Archive (SRA) 451 452 database. All published and unpublished metagenomic datasets that yielded crAss-like phage 453 contigs, the DNA preparation protocol, the sequencing technology, the assembly program, and information related to contig nomenclature, are briefly described in Supplementary Table 454 455 1. All reads were processed using Trimmomatic v0.32 to remove adaptor sequences and to trim reads when the Phred quality score dropped below 30 for a 4bp sliding window. 456 457 Trimmed reads were assembled using either SPAdes v3.6.2 (37) or metaSPAdes v3.10.0 (38). Contigs from the assembly of 702 metagenomic samples were assigned a specific 458 nomenclature, representing: [1] study/sample description, [2] SPAdes or metaSPAdes 459 460 assembly, and [3] numerical rank of largest-to-smallest assembled contigs. The full list of 461 contigs assembled in this study, the available associated metadata, and contig accession numbers, are detailed in Supplementary Table 2. 462

Detection and curation of crAss-like phages. The detection of crAss-like phage 463 contigs was performed as follows. The amino acid polymerase sequence of prototypical 464 crAssphage (UGP_018, NC_024711.1) was queried using BLAST v2.2.28+ (39) against a 465 translated nucleotide database consisting of assembled metagenome contig sequences. The 466 467 most conserved orthologous protein group detected in our initial putative crAss-like phage 468 screening included prototypical crAssphage protein UGP_092, which was annotated through the HHPred homology and structural prediction web server (40) as a phage terminase. This 469 was then used as a second genetic signature of crAss-like phages and used in an additional 470 471 BLAST search. All putative crAss-like phages selected for analysis met the following criteria: [1] a BLAST hit against either prototypical crAssphage polymerase or terminase 472

with an e-value less than 1e-05, [2] a BLAST query alignment length \geq 350bp, and [3] a minimum contig length of 70kb (representing near-complete crAss-like phage contigs).

Identification of crAss-like phage orthologous proteins and clusters. The encoded 475 476 proteins of crAss-like phages were predicted using Prodigal v2.6.3 (41). Orthologous proteins 477 shared between crAss-like phages were detected using OrthoMCL v2.0 using default parameters (42). The presence/absence of orthologous proteins between crass-like phages was 478 479 initially converted into a binary count matrix where the percentage of shared orthologous proteins was calculated (Figure 1B). The optimum number of phage clusters was calculated 480 481 using the percentage of shared homologous proteins using the NbClust v3.0 package for R (43). Hierarchical clustering was performed on the count matrix of percentage shared crAss-482 like phage orthologous proteins using Ward's minimum variance method ['Ward.D2' 483 484 algorithm in R (44)]. The resulting dendrogram was cut at k = 10 based on the estimation of 485 the number of crAss-like phage clusters (Figure 1A).

As a verification of the 10 predicted crAss-like phage clusters, the original abundance matrix of crass-like phage orthologous proteins was used to calculate Euclidean distances between samples. These distance variations were calculated using the t-SNE machine learning algorithm ['tsne' v0.1-3 for R; (45)] and plotted using ggplot v2.2.1 (Figure 2). The presence or absence of orthologous protein groups was used to determine the core proteome of crAss-like phage clusters (Supplementary Figure 8).

Phylogeny of crAss-like phage terminase sequences. Following the work of Yutin *et al.*, (16) all publically available crAss-like phage terminase sequences were included in an
additional phylogenetic analysis (Supplementary Figure 2). The terminase amino acid
sequences of crAss-like phages were aligned using Muscle v3.8.31 (46). The resultant
alignment was converted to Phylip format and phylogeny was determined by PhyML using a
JTT amino acid substitution model (47). The phylogenetic tree was visualised using FigTree

498 v1.4.3. The phylogenetic tree is coloured based on the crAss-like phage clustering analysis499 with node support values displayed.

Genomic comparisons of crAss-like phages. The average nucleotide identity between 500 501 crAss-like phage contigs was calculated using Pyani v0.2.3 by the ANIm method with a 502 500bp fragment size. Pairwise comparisons of complete crAss-like phage genomes belonging to Candidate Genera I was performed using Easyfig v2.2.2. Genomic start coordinates and 503 504 contig orientations were altered to match the published GenBank sequence of prototypical crAssphage NC 024711.1. The order of crAss-like phages in the Easyfig image was adjusted 505 506 to match to the order they appear in the average nucleotide identity analysis (Figure 3). The 507 Easyfig image was generated using tBLASTx comparisons, with a minimum BLAST length 508 of 50bp and identity of 30bp (Supplementary Figure 3). The presence of crAss-like phage 509 tRNA-encoding sequences were detected using ARAGORN v1.2.36 (48). To determine the 510 genomic packaging mechanism of crAss-like phages, metagenomic sequencing reads from a TruSeq (Illumina) manually fragmented DNA library were analysed using PhageTerm (25). 511 512 Single nucleotide polymorphisms (SNPs) of crAss-like phages were observed by aligning metagenomic sequencing reads to the consensus assembled contig sequence using Bowtie2 513 514 and Samtools, and visualising SNPs using Tablet v1.17.08.17 (49).

Alignment of virome metagenomic reads to crAss-like contigs. The quality filtered 515 516 reads from 532 human faecal viromes (as subset of 701 viromes selected based on availability 517 of sufficient metadata) were then aligned to the set of 93 nonredundant crAss-like phage genomic (with <90% of homology and/or <90% overlap between them) using Bowtie2 v2.3.0 518 (50) using the end-to-end alignment mode. A count table was generated with Samtools 519 520 v0.1.19 which was then imported into R v3.3.1 for statistical analysis. β-diversity of crAsslike viral populations in human cohorts was visualized using PCoA plot based on Spearman 521 522 rank distances (D = $1 - \rho$, where ρ is Spearman rank correlation coefficient of relative

abundance of different crAss-like contigs between samples). Statistical analysis was
performed using permutational multivariate analysis of variance (PERMANOVA)
implemented in Vegan v2.4.3 package for R (51) and non-parametric Kruskal-Wallis test.

526 Recruitment of a crAssphage faecal donor and faecal fermentations. Human faecal viromes from a number of ongoing studies sequenced using Illumina HiSeq and MiSeq 527 platforms were screened for crAss-like phages by aligning the obtained sequencing reads 528 against prototypical crAssphage NC_024711.1 using Bowtie2 v2.3.0. One individual (subject 529 530 ID 924) was found to carry crAssphage consistently at levels exceeding 30% of the total 531 number of reads over a one year period. The recruited individual is an adult female that suffers from gastritis and is vitamin B12 deficient. A frozen standard inoculum (FSI) sample 532 was processed as described by (52) with the following modification: the sample was 533 534 resuspended in 1X phosphate buffered saline (37 mM NaCl, 2.7 mM KCl, 8 mM Na₂HPO₄, 535 and 2 mM KH₂PO₄.), 0.05% (w/v) L-cysteine (Sigma Aldrich, Ireland) and (1 mg/L) resazurin (Sigma Aldrich, Ireland). The crAssphage-rich FSI was inoculated into 400 ml 536 537 YCFA-GSCM broth in a 500 ml fermenter vessel at 5% (v/v). Fermentation media was prepared exactly as described by (53) with the addition of glucose (2 g/L), soluble starch (2 538 g/L), cellobiose (2 g/L) and maltose (2 g/L). Fermentation was performed in batch format at 539 approximately 37°C for 51 hours. Dissolved oxygen was sustained at <0.1% by constantly 540 541 sparging the vessel with anaerobic gas mix (80% (v/v) N_2 , 10% (v/v) CO_2 , 10% (v/v) H_2) and 542 stirring at 200 rpm. Both 2M NaOH and HCl solutions were used to maintain pH at ~7. Samples were collected at the following time points; 0, 4, 21, 28, 45 and 51 hours. Collected 543 samples were centrifuged at 4,700 rpm at +4°C for 10 minutes. The resulting supernatants 544 545 were filtered once through a 0.45 μ M pore syringe filter and stored at +4°C. Resultant pellets were stored at -80°C. 546

547 Extraction of viral nucleic acids and sequencing library preparation. Total virome extractions were performed on 0.45 μ M pore filtered fermentation supernatants. Solid NaCl 548 and polyethylene glycol 8000 were added to the filtrates to give a final concentration of 0.5M 549 and 10% (w/v), respectively. After overnight incubation at +4°C samples were centrifuged at 550 551 4,700 rpm and +4°C for 20 minutes. The pellets were then resuspended in 400µl of SM buffer (1M Tris-HCl pH 7.5, 5M NaCl, 1M MgSO₄) and briefly vortexed with an equal volume of 552 553 chloroform. This mixture was then centrifuged at 2,500g for 5 minutes using a standard desktop centrifuge. The resultant aqueous phase was then transferred into an Eppendorf to 554 555 which 40µl DNase buffer (10mM CaCl₂ and 50mM MgCl₂) and 8U and 4U TURBO DNase (Ambion/ThermoFisher Scientific) and RNase I (ThermoFisher Scientific) were added, 556 respectively. This was incubated at 37°C for 1 hour followed by an enzyme inactivation step 557 558 at 70°C for 10 minutes. This was followed by the addition of 2µl proteinase K and 10% SDS 559 and further incubation at 56°C for 20 minutes. Lastly, 100µl phage lysis buffer (4.5 M guanidinium isothiocyanate, 44 mM sodium citrate pH 7.0, 0.88% sarkosyl, 0.72% 2-560 mercaptoethanol) was added to lyse the viral particles. The final incubation was carried out at 561 65°C for 10 minutes. The resulting lysates were lightly vortexed with an equal volume of 562 phenol/chloroform/isoamyl alcohol 25:24:1 (Fisher Scientific) and were centrifuged at room 563 temperature for 5 minutes at 8,000g. This was again repeated with the resulting aqueous 564 phase. Following the second extraction, the aqueous phase was passed through a DNeasy 565 566 Blood and Tissue Kit (Qiagen) for final lysate purification. The wash steps were each repeated twice and the final elution was carried out in 50µl elution buffer. Viral DNA 567 quantification was carried out with the Qubit HS DNA Assay Kit (Invitrogen/ThermoFisher 568 569 Scientific) in a Qubit 3.0 Flurometer (Life Technologies). The viral nucleic acids were then subjected to reverse transcription using SuperScript IV Reverse Transcriptase (RT) kit 570 571 (Invitrogen/ThermoFisher Scientific). The protocol was carried out exactly as described in

572 the manufacturer's protocol for random hexamer primers. Following this, 1µl of the reversed transcribed viral DNA was subjected to GenomiPhi V2 (GE Healthcare) Multiple 573 Displacement Amplification (MDA). Finally, MDA and non-MDA viral DNA was prepared 574 for sequencing using TruSeq DNA Library Preparation Kit (Illumina, Ireland). All steps were 575 performed as per the manufacturer's instructions. Prepared libraries were sequenced on an 576 Illumina HiSeq platform (Illumina, San Diego, California) with 2x300bp paired-end 577 chemistry at GATC Biotech AG, Germany. Reads were filtered, trimmed and assembled into 578 contigs as described above. A count matrix was created by aligning quality-filtered reads back 579 580 to contigs using Bowtie2 and Samtools.

CrAssphage PCR detection. Two oligonucleotide primer pairs were designed based 581 on the prototypical crAssphage DNA polymerase sequence UGP_018 (1) using PerlPrimer 582 583 software (54). Primer sequences as follows: CrAss-Pol-F5 5'are GCCTATTGTTGCTCAAGCTATTGAA-3', CrAss-Pol-R5 5'-584 5'-ACAACAGAACCAGCTGCCAT-3', CrAss-Pol-F6 585

586 AGTGGTCTTGCTCCNGAACAATGG-3' and CrAss-Pol-R6 5'-

AACCTCCAGTTGCAACAGTATAAGT-3'. PCR products were cloned into pCR2.1-TOPO 587 TA vector (ThermoFisher Scientific) and obtained plasmids at known concentrations were 588 used to establish calibration curves through serial two-fold dilutions. Subsequently, qPCR 589 590 were run in 15µl reaction volumes using SensiFAST SYBR No-ROX mastermix and 591 LightCycler 480 thermocycler with the following conditions: initial denaturation at 95°C for 5 minutes, then 35 cycles of 94°C for 20 seconds, 55°C for 20 seconds and 72°C for 20 592 seconds, with a final extension at 72°C for 7 minutes. All samples were run in triplicate and 593 594 the standard error was determined following calculation of DNA concentration based on the above standard curve. 595

596 Electron microscopy and detection of crAssphage proteins. A virus-enriched fraction of the crAssphage positive faecal sample, collected from subject ID 924, was 597 prepared for electron microscopy imaging as follows. A 1:20 suspension (w/v) of faeces was 598 599 prepared in SM buffer followed by vigorous vortexing until homogenised. The homogenised 600 sample was chilled on ice for 5 minutes prior to centrifugation twice at 4,700 rpm for 10 minutes at $+4^{\circ}$ C. The resulting supernatant was then filtered twice through a 0.45 μ M pore 601 602 syringe filters. The filtrate was ultra-centrifuged at 120,000g for 3 hours using a F65L-6x13.5 rotor (ThermoScientific). The resulting pellets were resuspended in 5 ml SM buffer. The viral 603 604 suspensions were ultracentrifuged again by overlaying them onto a caesium chloride (CsCl) step gradient of 5M and 3M, followed by centrifugation at 105,000g for 2.5 hours. A band of 605 606 viral particles visible under side illumination was collected and buffer-exchanged using 3 607 sequential rounds of 10-fold diluting and concentrating to the original volume by ultra-608 filtration using Amicon Centifugal Filter Units 10,000 MWCO (Merck). The purified fraction was then analysed by qPCR for the presence of crAssphage as described above. Following 609 610 this, 5µl aliquots of the viral fraction were applied to Formvar/Carbon 200 Mesh, Cu grids (Electron Microscopy Sciences) with subsequent removal of excess sample by blotting. Grids 611 were then negatively contrasted with 0.5% (w/v) uranyl acetate and examined at UCD 612 Conway Imaging Core Facility (University College Dublin, Dublin, Ireland) by transmission 613 614 electron microscope. The faecal viral fraction from subject ID 924 was further concentrated 615 using Amicon Ultra-0.5 Centrifugal Filter Unit with 3 kDa MWCO membrane (Merck, Ireland). This concentrated fraction was loaded onto a premade Bolt 4-12% Bis-Tris Plus 616 reducing SDS-PAGE gel (Invitrogen) and separated at 200 V for 30 minutes using 1X 617 618 NuPAGE MOPS SDS Running Buffer. Six brightest bands with approximate molecular weights of 28, 35, 45, 55, 120 and 200 kDa were excised and subjected to MALDI-TOF/TOF 619

620 (Bruker ultraflex III) protein identification following in-gel trypsinization, at Metabolomics
621 & Proteomics Technology Facility (University of York, York, UK).

16S rRNA gene library preparations. Total DNA was extracted from the pellets 622 623 formed following centrifugation of fermentation samples. This was carried out using the QIAamp Fast DNA Stool Mini Kit (Qiagen, Hilden, Germany). All steps were carried out as 624 per the manufacturer's protocol with the addition of a bead-beating step to aid total DNA 625 626 extraction from the bacterial cells. Approximately 200mg of each pellet was placed in a 2ml screw-cap tube containing a mixture of one 3.5 mm glass bead, a 200µl scoop of 1mm 627 628 zirconium beads and a 200µl scoop of 0.1mm zirconium beads (ThistleScientific) with 1ml of InhibitEX Buffer. Bead-beating was carried out three times for 30 seconds using the 629 FastPrep-24 benchtop homogeniser (MP Biomedicals). Between each bead-beating the 630 631 samples were cooled on ice for 30 seconds. The samples were then lysed at 95°C for 5 632 minutes. All other steps were carried out as per the manufacturer's protocol. Following extraction of total bacterial DNA, the hypervariable regions of V3 and V4 16S ribosomal 633 634 RNA genes were amplified from 15ng of the DNA using Phusion High-Fidelity PCR Master Mix (ThermoFisher Scientific) and 0.2µM of each of the following primers, containing 635 Illumina-compatible overhang adapter 16S-FP: 5'-636 sequences: TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCCTACGGGNGGCWGCAG-3' and 637 5'-638 16S-RP:

639 GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGACTACHVGGGTATCTAATCC-

3'. The PCR program was run as follows: 98°C for 30 seconds, 25 cycles of 98°C for 10
seconds, 55°C for 15 seconds and 72°C for 20 seconds, with a final extension of 72°C for 5
minutes. The amplicons were then purified using Agencourt AMPure XP magnetic beads
(Beckman-Coulter) followed by a second PCR to attach dual Illumina Nextera indices using
the Nextera XT index kit v2 (Illumina). Purification was performed once again and the

645 libraries were quantified using a Qubit dsDNA HS Assay Kit. The libraries were then pooled in equimolar concentration and sent for sequencing on an Illumina MiSeq platform (Illumina, 646 San Diego, California) at GATC Biotech AG, Germany. The quality of the raw reads were 647 648 assessed with FastQC (v11.5) and initial quality filtering was performed using Trimmomatic 649 v0.36. Filtered reads were imported into R (v3.4.3) for analysis with DADA2 v1.6.0. (55) 650 Further quality filtering and trimming (maxN of 0 and a maxEE of 2) was carried out on both 651 the forward and reverse reads with only retention in cases of pairs being of sufficient high 652 quality. Error correction was performed on forward and reverse reads separately and 653 following this, reads were merged. The resulting unique Ribosomal Variant Sequences (RSVs) were subjected to further chimera filtering using USEACH v8.1 (56) with the 654 655 Chimera-Slayer gold database v20110519. The retained, high quality, chimera-free, RSVs 656 were classified with the RDP-classifier in mothur v1.34.4 (57) against the RDP database 657 v11.4 (phylum to genus) and SPINGO (58) for species assignment. Plots were generated using the R package ggplot2 v2.2.1. 658

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667 Author contributions:

EG and SRS performed the laboratory and bioinformatic work, respectively. AS assisted in
both the laboratory and bioinformatic analyses. AGC performed the 16S analysis. FJR, LAD
and EGT assisted in the design, implementation and interpretation of experiments. EG, AS
and SRS wrote the paper and generated the figures. AGC, FJR, LAD and EGT reviewed
drafts of the manuscript and provided constructive criticism for its improvement. PR and CH
secured the funding and wrote the paper. All authors contributed to the analysis of the data.

674

675 **Conflict of interest:**

The authors declare no conflict of interest.

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678 **Data deposition:**

- The 244 crAss-like phage contigs analysed in this study have been submitted to GenBank and
- are currently under revision. Contigs are currently accessible at:
- 681 https://figshare.com/articles/crAss-like_contigs_fasta_tar_gz/6098321

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817 **Figure Legends**

Figure 1. Determination of crAssphage candidate subfamilies and genera based on the 818 percentage of shared protein-encoding genes. (A) The 4 red lines cut the hierarchical 819 820 clustering dendrogram of crAss-like phage contigs, with Euclidean distances calculated between the percentages of shared protein-encoding genes, into the 4 proposed candidate 821 subfamilies of crAss-like phages. The histogram insert (top-right) represents the calculated 822 823 optimal number of crAss-like phage clusters. The 10 optimal crAss-like phage clusters represent the putative candidate genera, and are assigned specific colours. (B) Heatmap 824 825 showing the percentage of shared protein-coding genes between crAss-like phage genomes. CrAss-like phages with 20-40% shared protein encoding genes are considered related at the 826 827 subfamily level while phages with >40% similarity are believed to be related at the genus 828 level, consistent with the calculated number of crAss-like phage clusters.

Figure 2. Two-dimensional ordination of crAss-like phages based on the abundance of their protein-encoded orthologous sequences was performed using t-SNE machine learning algorithm. (A) CrAss-like phages are coloured by candidate genus annotations and shape is determined by their origin. CrAss-like phages originating from individuals with kwashiorkor and marasmus, or lacking metadata, are grouped together as 'Other/Unknown'. (B) CrAsslike phages are coloured by the percentage G+C nucleotide composition of their contig, while shape represents complete (circular) or partial (linear) genomes.

Figure 3. Average nucleotide identity of crAss-like phage contigs. The column annotation colour scheme highlights the predicted crAss-like phage candidate genus annotations, while the coloured row annotation represents the origin of the respective crAss-like phage contig.

Figure 4. Prevalence of crAss-like phage in human faecal viromes. (A) Relative abundance of total crAss-like phage in several cohorts differing in age, health status and country of origin, based on the fraction of metagenomic reads aligned. Bars represent median relative

abundances, the values within boxes represent percentage of positive samples. (B) Relative
abundance of specific crAss-like candidate genera in total human populations analysed. (C)
PCoA plot of crAss-like phages based on Spearman rank distances.

Figure 5. Relative abundance of the ten candidate genera of crAss-like phages in six different
human cohorts based on the fraction of metagenomic reads aligned. Bars represent median
relative abundances, while values within boxes represent percentage of positive samples.

848 Figure 6. Analysis of crAss-like phage dynamics in a faecal fermenter. (A) Evidence of crAssphage *sensu stricto* propagation following *in vitro* fermentations (standard error, n=3). 849 850 The level of crAssphage sensu stricto propagation was determined by qPCR analysis of viralenriched DNA, respectively, using primers specific to a segment of the crAssphage sensu 851 852 stricto DNA polymerase gene. (B) Six additional crAss-like phages, that group into five of 853 the candidate genera, were identified following sequencing of the same viral-enriched DNA 854 from the fermenter. The relative abundance of each of these crAss-like phages is skewed due to the biased amplification of other components of the viral-enriched DNA fraction that is 855 856 associated with multiple displacement amplification.

Figure 7. CrAss-like phage morphology was examined using a CsCl fraction purified from a 857 crAssphage rich faecal filtrate of donor subject ID 924. (A) Analysis of the fraction through 858 transmission electron microscopy (TEM) was performed. The TEM images are largely 859 860 dominated by Podovirdae (53%), Microviridae (29%), Siphoviridae (15%) and other phage 861 morphologies (3%). (B) Further examination of the observed *Podoviridae* identifies two variants with differing tail morphologies. Both variants have head diameters of ~76.5 nm. (C) 862 863 SDS-PAGE gel of the CsCl fraction. Six bands containing possible crAssphage proteins were 864 excised and analysed by mass spectrometry. A protein, denoted as Fferm_ms_2_MCP, isolated from the ~55 kDa (*) band was found to have high sequence similarity with 865 Candidate Genus V crAss-like phages. (D) Sequencing of the CsCl purified viral fraction, 866

without multiple displacement amplification, showed that approximately 40% the readsaligned to crAss-like phages.

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870 Supplementary Figure Legends

Supplementary Figure 1. Phylogeny of crAss-like phage terminase protein sequences, including publically available terminase sequences from the Yutin *et al.* (2017) characterisation of familial-related crAss-like phages. The figure legend insert corresponds to the colour scheme of the 10 proposed candidate genera groupings. NC_024711 crAssphage and IAS virus, discussed in the main text, are highlighted in red. Bootstrapping node support values are shown.

Supplementary Figure 2. Comparison of general structural feature of representative
complete circular genomes of the 10 proposed genera of crAss-like bacteriophages.
Innermost circle (green/blue), G+C skew; middle circle, G+C content deviation from mean
value; outermost circle, protein-coding genes (CDS) located on positive (red) and negative
(blue) DNA strands, respectively; and tRNA genes (orange).

Supplementary Figure 3. Comparison of circular Candidate Genus I crAss-like phage genomes. Start co-ordinates of crAss-like phage genomes were adjusted to match crAssphage *sensu stricto*. The order of crAss-like phage genomes was determined by the average nucleotide identity comparisons. Open reading frames corresponding to specific predicted phage structural proteins are highlighted.

Supplementary Figure 4. The relative abundance of 16S rRNA throughout the crAssphagerich frozen standard inoculum initiated faecal fermentation. (A) The relative abundance of the major genera detected throughout the fermentation. *Bacteroides* (*), the genus hypothesised to be associated with crAssphage, can be seen to decrease between time points 0 and 4 of the fermentation after which levels gradually begin to increase again. (B) The relative abundance

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892	of total <i>Bacteroides</i> at each time point. (C) Abundances of individual <i>Bacteroides</i> species
893	detected. B. dorei is found to be particularly abundant and seemingly inversely proportional
894	to the detected crAssphage levels.

- 895 **Supplementary Figure 5.** Quantitative PCR analysis of filtrates obtained with different pore
- sizes from a crAssphage-rich faecal sample collected from subject ID 924.

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897 Supplementary Figure 6. Comparison of 16S rRNA Prevotella abundances in healthy Irish

adults and infants with Malawian infants.

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899 Supplementary Figure 7. Visualisation of an example of metagenomic read-specific single

nucleotide polymorphisms within the assembled of crAss-like phage contig, Fferm_ms_2,

- highlighting within sample species and/or strain level diversity of crAss-like phages are not
- 902 resolved.
- 903 Supplementary Figure 8. Visualisation of the core proteome of the 10 crAss-like phage904 candidate genera.



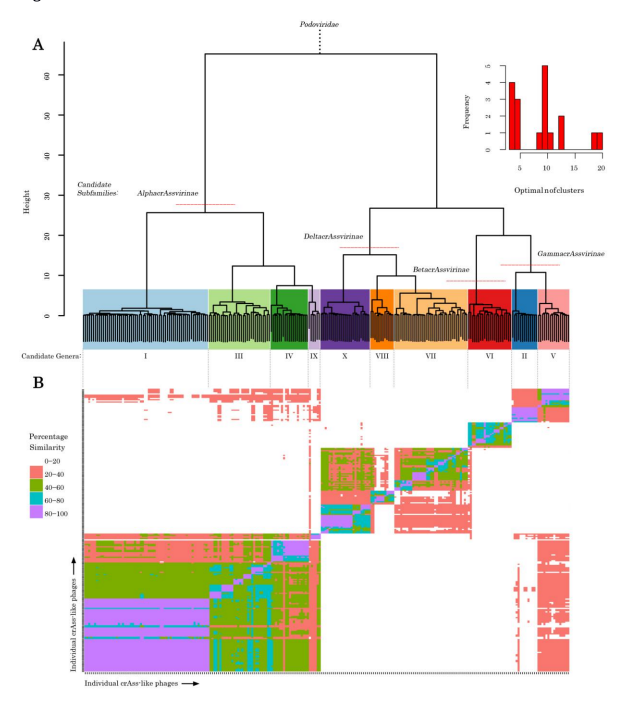


Figure 2.

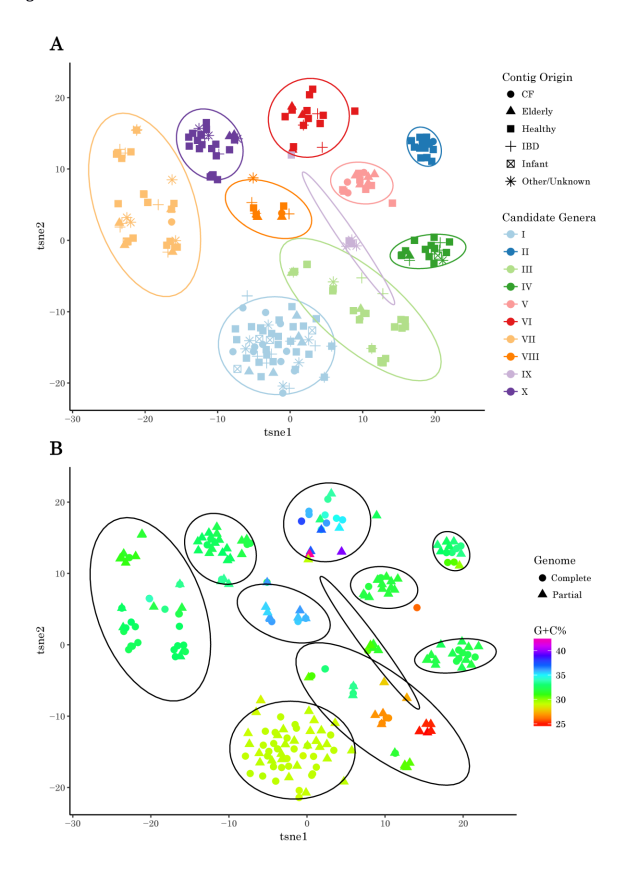


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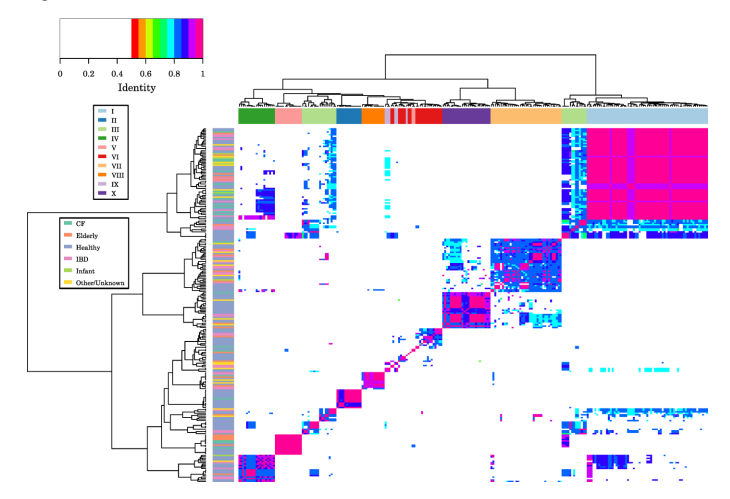


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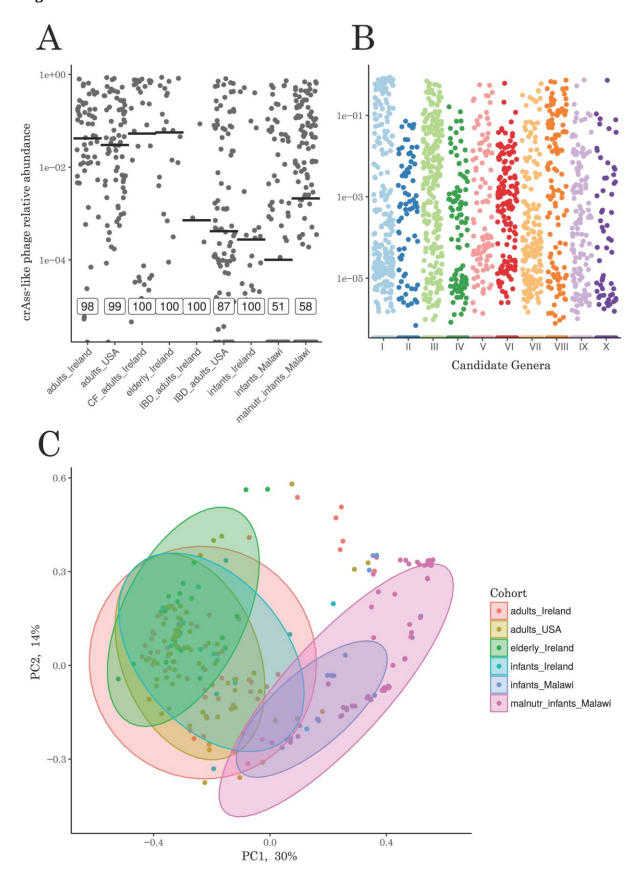


Figure 5

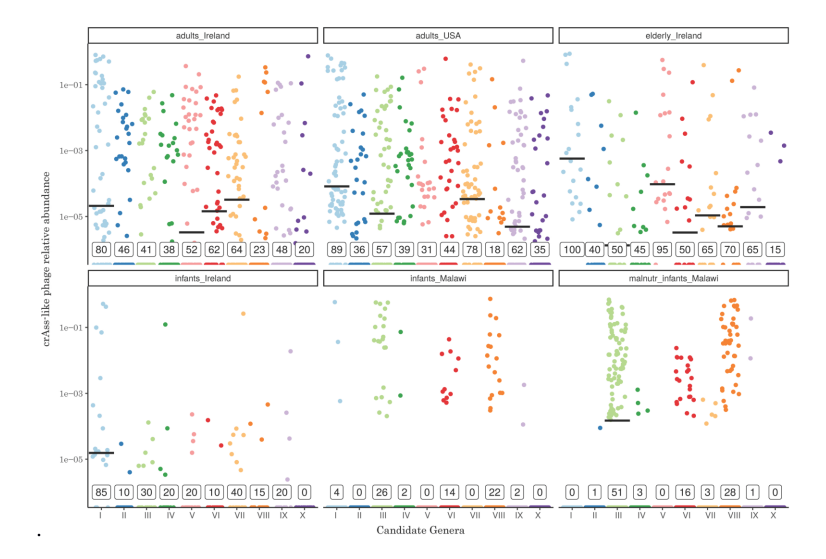


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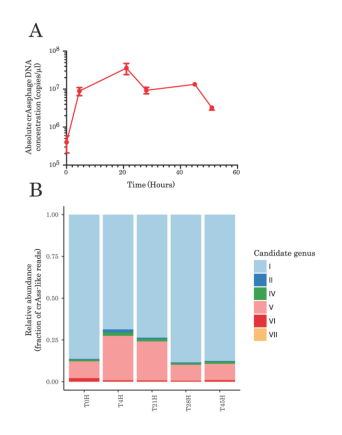
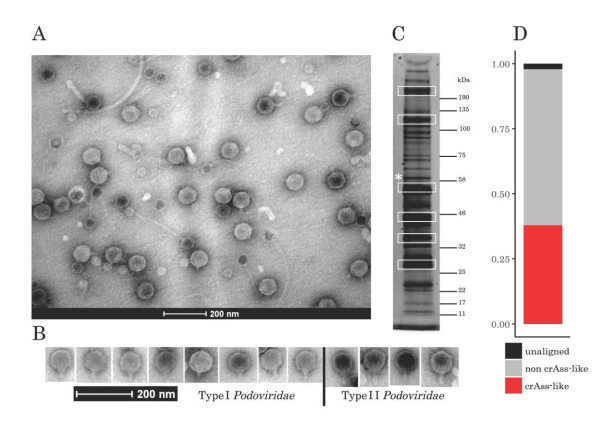
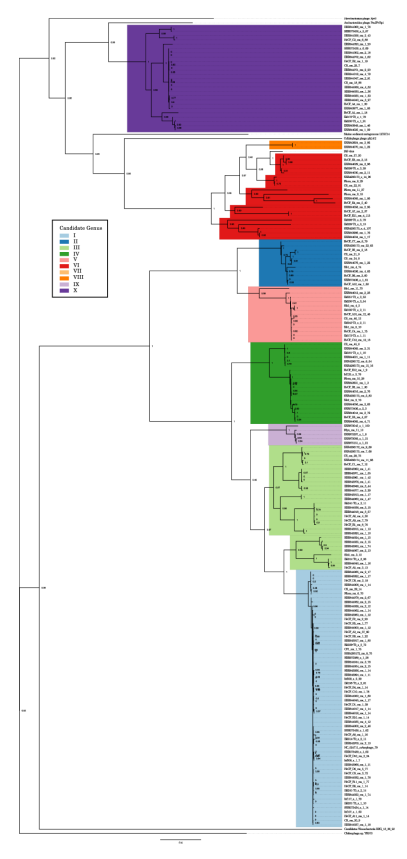


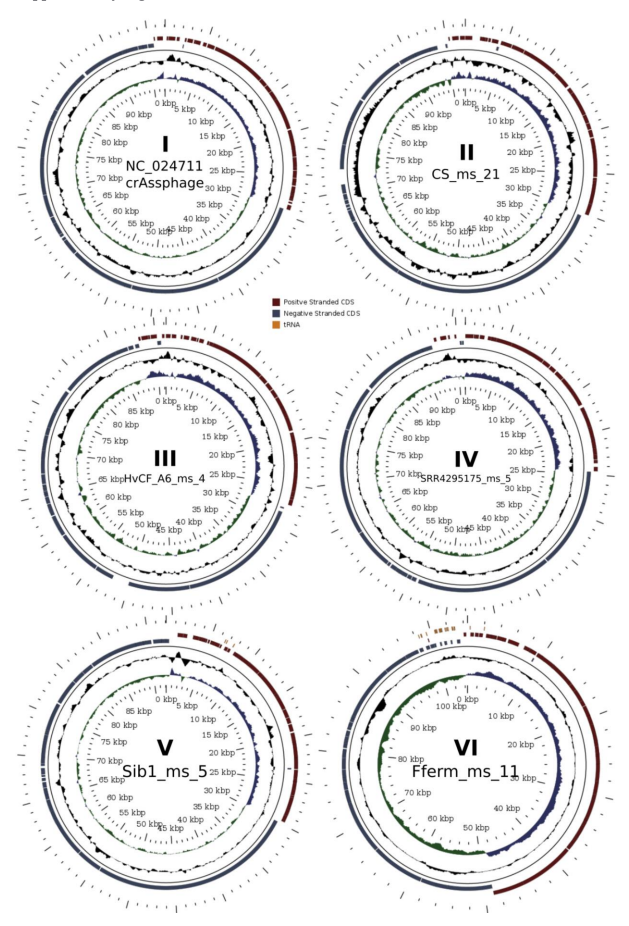
Figure 7.



Supplementary Figure 1.

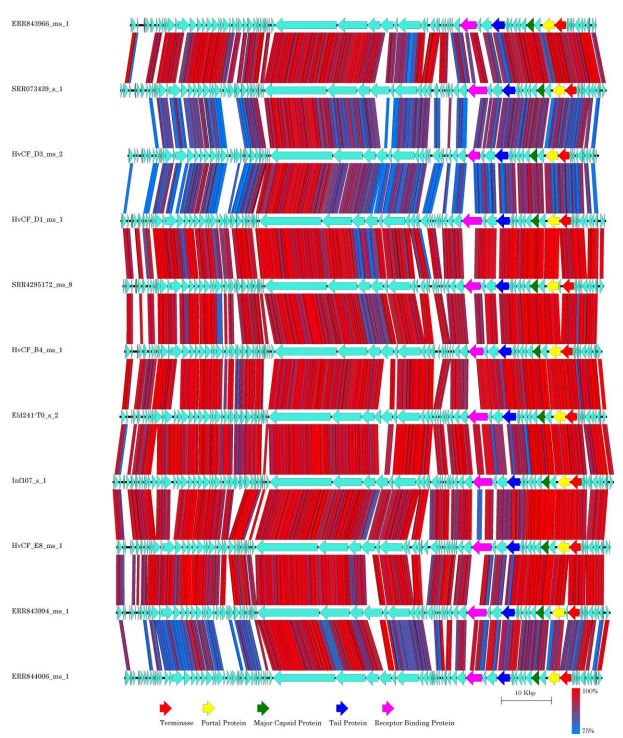


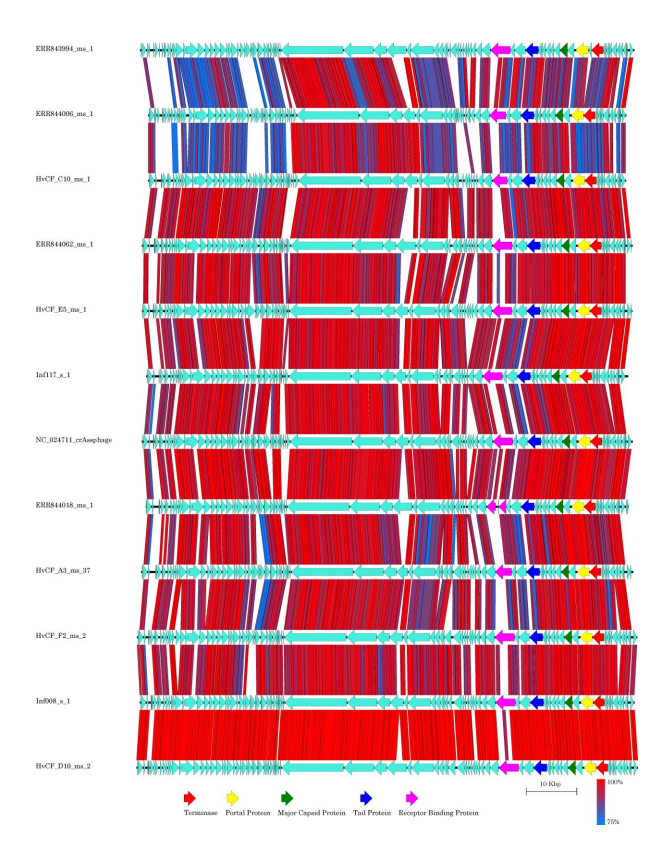
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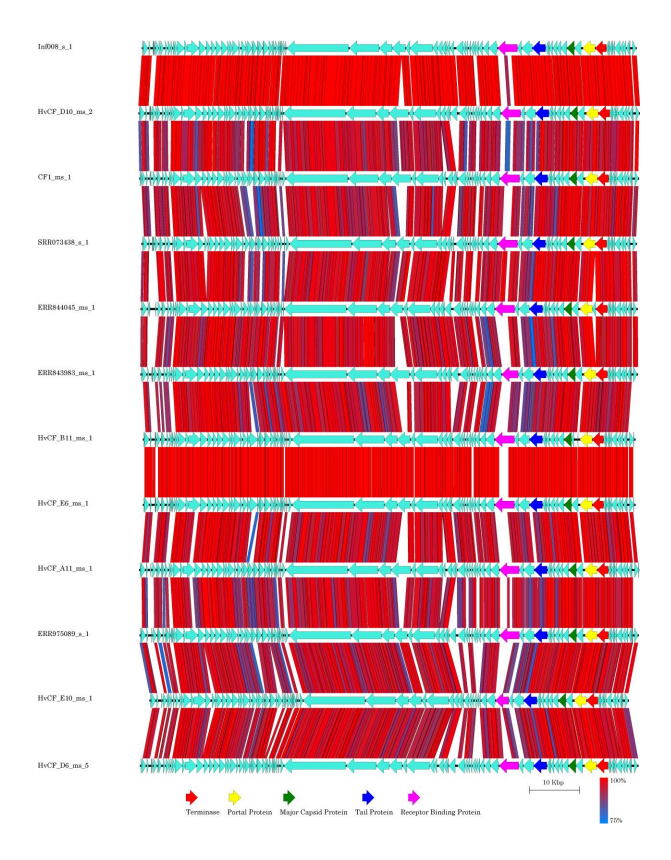




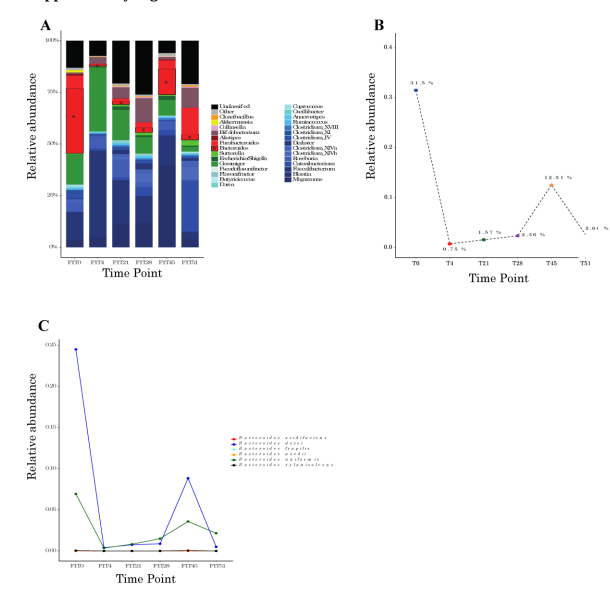
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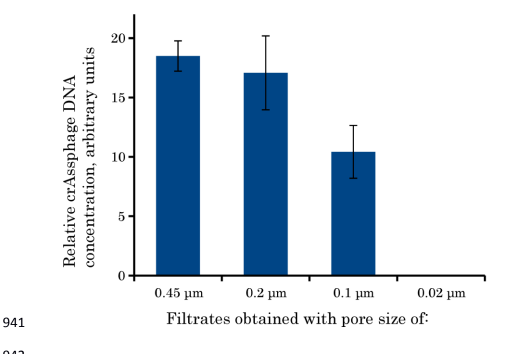


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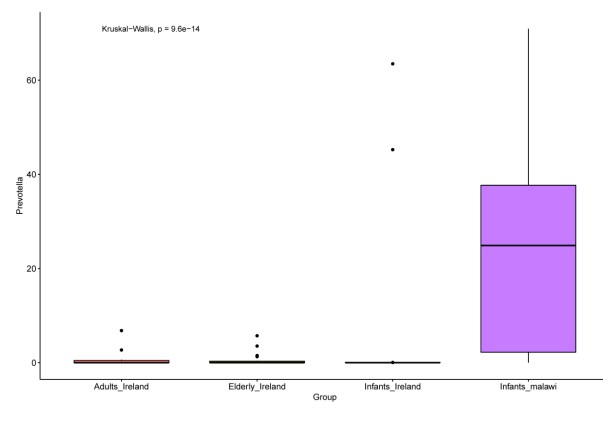
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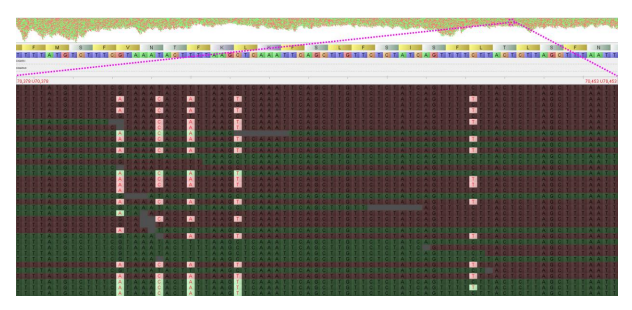


Supplementary Figure 6.





Supplementary Figure 7.



949 Supplementary Figure 8.

