## **1** The human gut virome database

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#### 10 ABSTRACT

11 The gut microbiome profoundly impacts human health and disease, but viruses that infect these microbes 12 are likely also important. Problematically, viral sequences are often missed due to insufficient reference 13 viral genomes. Here we (i) built a human gut virome database, GVD, from 648 viral particle 14 metagenomes or microbial metagenomes from 572 individuals previously searched for viruses, (ii) 15 assessed its effectiveness, and (iii) conducted meta-analyses. GVD contains 13,203 unique viral 16 populations (approximately species-level taxa) organized into 702 novel genera, which roughly doubles 17 known phage genera and improves viral detection rates over NCBI viral RefSeq nearly 60-fold. Applying 18 GVD, we assessed and rejected the idea of a 'core' gut virome in healthy individuals, and found through 19 meta-analyses that technical artifacts are more impactful than any 'treatment' effect across the entire 20 meta-study dataset. Together, this foundational resource and these findings will help human microbiome 21 researchers better identify viral roles in health and disease.

# 22 Main text

The human gut microbiome is now thought to play an integral role in health and disease <sup>1-4</sup>.
Persistent alterations in the structure, diversity and function of gut microbial communities—dysbiosis—
are increasingly recognized as key contributors in the establishment and maintenance of a growing
number of disease states <sup>5-7</sup>, including obesity <sup>8</sup> and cancer <sup>9</sup>. Gut dysbiosis can develop from complex

interplays between host, cognate microbiota and external environmental factors <sup>10,11</sup>. Within the gut
microbial consortium, the bacteriome has been the most extensively studied, where significant shifts in
population dynamics have been observed between healthy and diseased individuals <sup>12</sup>. However,
emerging views <sup>10,13,14</sup> suggest that the gut virome plays an important role in homeostatic regulation and
disease progression through multiple interaction paths with the co-occurring bacteriome, and even
directly with human immune system components <sup>15</sup>.

33 The first step in studying viruses in complex communities is to "see" them. Problematically, 34 identifying viral sequences in large datasets is notoriously challenging. Because viruses lack a universal 35 viral marker <sup>16</sup>, as opposed to bacterial 16S rRNA for example, researchers often resort to sequence 36 homology searches against reference databases (e.g. NCBI viral RefSeq). Such searches are variably 37 successful with anywhere from 14% to 87% of the observed gut viral genomes having detectable similarity to viruses in such databases <sup>10</sup>. This large range stems from several factors that are not 38 39 mutually-exclusive including the following: (i) broad under-representation of viral genome space in 40 databases, (ii) non-standardized database usage per study, (iii) overrepresentation of certain virus groups 41 due to sample preparation and cultured host availability, and (iv) natural sample variation. In addition, although viral reference datasets are being generated at unprecedented rates <sup>17</sup>, these new data are rarely 42 43 incorporated for cross-comparisons, which inflates virus novelty in new datasets and/or leaves many virus 44 sequences undetected. Therefore, given the rapid accrual of so many studies, there is a need to aggregate 45 their findings into a central gut-specific database to improve gut virome inference capabilities.

Here we collected and curated 648 gut metagenomes from 21 datasets (i.e., any metagenomic
dataset that looked at gut viruses published before 2018), consistently processed them to map known and
unknown viral populations, and used this in multiple meta-analyses to assess improvement and reveal
new biology. The resulting Gut Virome Database (GVD) was born by (i) collecting 648 gut metagenomes
from 572 individuals, (ii) extensive metadata curation through literature mining and, as needed, direct
communication with the original researchers, and (iii) re-analysis of the virome data to establish
consistent processing and extensive virus identification. The value of GVD was assessed for performance

53 against the best currently available databases (NCBI viral RefSeq and IMG/VR<sup>18</sup>), and then used to re-

54 evaluate global diversity patterns and the relationship between gut virome diversity and diet.

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## 56 RESULTS AND DISCUSSION

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58 GVD contains 13,204 viral populations, dominated by phages

59 To build a collection of the commensal human gut virome, 648 metagenomic samples from 572 60 individuals were processed from all datasets publicly available as of December 2017 (n=19), along with 2 61 unpublished datasets where access was granted prior to publication. These studies represented a total of 62 1.28 Tbp of sequence data derived from a spectrum of gut virome study areas including: (i) healthy gut viromes of infants  $^{19,20}$  and adults  $^{21-26}$ , as well as individuals experiencing (ii) fecal matter transplant, or 63 FMT<sup>27-31</sup>, (iii) inflammatory bowel disease, or IBD<sup>32,33</sup>, (iv) HIV infection<sup>34</sup>, (v) Type I diabetes<sup>35,36</sup>, 64 (vi) malnutrition <sup>37</sup>, or (vii) chronic fatigue syndrome <sup>38</sup> (see **Supplementary Table 1**). Datasets had a 65 66 worldwide distribution, though most originated from the United States (48.4%; Fig. 1a). All reads were 67 processed consistently, assembled into contigs and viral-like sequence were identified using three 68 independent methods and validated by cross-comparisons between methods (Fig. 1b, see Methods). To 69 avoid duplicate viral fragments/partial virus genomes across the datasets, contigs were de-replicated by 70 clustering sequences according to percentage of average nucleotide identity (ANI) and sequence length. Multiple reports  $^{17,39-43}$  have revealed that > 95% ANI was a suitable threshold for defining a set of 71 72 closely-related discrete 'viral populations', with follow-on studies suggesting that this cut-off establishes populations that are largely concordant with a biologically relevant viral species definition<sup>39,41,44</sup>. Using 73 74 this clustering strategy, we identified highly variable numbers of unique viral populations per study 75 (range: 0 - 3596; mean = 670) (Supplementary Fig. 1a). GVD comprises 13,203 viral populations (N50 = 34.220 bp ; L50 = 2,066 bp). For context, NCBI's viral RefSeq v88 (released May 2018) database holds 76 77 8,013 viruses of eukaryotes, bacteria and archaea from all environments, combined. Moreover, if only 78 comparing phage genomes to the same database. GVD contains 7 times more phages compared to the

entire set of cultured phage isolates in viral RefSeq to date. Thus, GVD greatly augments the repertoire ofknown viruses in the human gut.

81 Taxonomically, 96.1% of GVD viral populations are bacterial viruses (i.e., phages), with a 82 minority of GVD viral populations more likely to represent eukaryotic viruses (3.8%) and archaeal 83 viruses (0.1%) (Fig. 2a). Though in the minority, the 505 eukaryotic viruses were taxonomically diverse 84 (14 families), dominated by ssDNA families Anelloviridae (72%), Genomoviridae (10%) and 85 Circoviridae (8%). All, with the exception of Genomoviruses, have been reported previously in the datasets underlying GVD<sup>34</sup>. Among the phages, 82% did not have ICTV classification, with the 86 87 remaining fraction comprised of dsDNA tailed phage families (Siphoviridae, Myoviridae and 88 Podoviridae), Microviridae and Inoviridae (see Supplementary Table 2). Twelve unknown archaeal 89 viral populations were detected, with no close genome/gene homology to any of the classified archaeal 90 viruses. The high number of unclassified phages likely results from underrepresentation of gut phages in 91 the database, coupled to unresolved and/or missing taxonomic assignments for ~ 60% of reference phage 92 genomes in RefSeq, with the currently classified fraction organized into ~250 genera<sup>45</sup>. To fill this phage 93 and archaeal virus taxonomic classification gap, we used a genome-based, gene-sharing network strategy 94 <sup>46,47</sup> that *de novo* predicts genus-level groupings ('viral clusters' or 'VCs') from viral population data. A network was computed from 6,373 GVD phage genomes (only those  $\geq 10$  kb in length; 48% of GVD), 95 96 combined with 2,304 curated reference phage genomes from NCBI Viral RefSeq (version 88). The 97 resulting gene-sharing network (Fig. 2b) revealed 957 VCs, 702 of which were novel and exclusively 98 composed of GVD genomes  $(3,220 \text{ viral genomes or } \sim 51\% \text{ of GVD genomes})$ . This would roughly 99 double the current number of ICTV-recognized phage genera. Though not explored here, as our goals 100 focused on taxonomic classification, the shared protein content within and between VCs calculated in our 101 network analyses could be used to guide qPCR assays for NGS validation <sup>48</sup> and/or tracking of viruses at 102 either the viral population- or genera- level under changing conditions<sup>35</sup>.

103 Next, we sought to link phage populations to their hosts using *in silico* strategies (see Methods).
104 The most common identifiable phage hosts (Fig. 2c) in GVD belonged the bacterial phylum Firmicutes

105	(38%), about 2-fold more than the next most abundantly identified host phyla (Bacteroidetes and
106	Proteobacteria; see Supplementary Table 2). Though Firmicutes and Bacteroides are the most prominent
107	bacterial phyla in the human gastrointestinal tract <sup>49</sup> , Firmicutes typically outnumber Bacteroidetes in
108	unhealthy individuals with metabolic and digestive disorders <sup>50–52</sup> . GVD metagenomes originated from
109	~16% healthy individuals and ~84% unhealthy individuals, many of which have metabolic and digestive
110	disorders. Thus, it is perhaps not surprising that most of the annotated viral populations were linked to the
111	phylum Firmicutes.

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#### 113 GVD significantly improves virus detection in all gut datasets

114 We then quantitatively evaluated virus identification sensitivity (through read mapping) between 115 multiple databases by comparing the number of identified viral populations in each study detected by 116 GVD, viral RefSeq v88, IMG/VR 1.1 (2018 release) and the individual virome datasets ('IV') from each 117 study (Fig. 3). For the latter, IV reads were mapped against viral populations (predicted in this study) 118 derived exclusively from its matching IV. In all datasets, GVD surpassed viral RefSeq (mean increase: 119 59-fold  $\pm$  95-fold) and IVs (mean increase: 3.2-fold  $\pm$  6.6-fold). In 5 of 18 studies (28%), GVD 120 outperformed IMG/VR (mean increase: 1.1-fold  $\pm 2$ -fold), with the remaining studies finding no 121 significant difference between or too low of a sample size to compare GVD and IMG/VR. After GVD, 122 IMG/VR was the next best performing database for viral detection in the gut, as our tests showed an 123 average of 49-fold (± 87-fold) increase over viral RefSeq. IMG/VR was expected to surpass viral RefSeq, 124 as it aggregates both cultivated reference virus genomes, >12,000 prophages and >700,000 uncultivated virus genomes/fragments from many environments, including multiple human body sites<sup>53</sup>. Moreover, 125 126 given the high performance of IMG/VR in our tests, we wondered about the extent of viral population 127 overlap with GVD (Fig. 3b). There were 1,730 viral populations shared between the two databases, but 128 still each database is overwhelmingly unique (82% and 69% unique to GVD and IMG/VR, respectively). 129 This is because IMG/VR includes human gut studies that did not explore the viral fraction as well. 130 Overall, the significant increase in virus detection by GVD over other databases (two-tailed Mann-

131	Whitney U-tests; $p$ -value < 0.05) highlights the low representation of gut viruses recorded in RefSeq and
132	thus demonstrates the value of GVD for sequence-based virus identification in human gut microbiome
133	datasets. Because the datasets used to compile GVD were originally analyzed most often (55% of the
134	studies) using viral RefSeq as the primary source to identify viruses (Supplementary Table 1), we
135	wondered whether significant fractions of viruses could have been missed, and whether a possibly
136	reduced viral "signal" would influence previous conclusions.

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# 138 MDA amplification skews diversity and prohibits quantitative analysis of gut viromes

139 To evaluate this possible reduced viral "signal", we first examined the role of methodological 140 approaches in influencing inferences about ssDNA viruses. This is because we noticed that the bulk of 141 ssDNA eukarvotic viruses (Anelloviruses, Circoviruses, Genomoviruses, Geminiviruses) and phages 142 (Microviruses) originated from only 4 of the 21 studies gathered in this work (Fig. 4 a,b). These studies evaluated 2 infant gut viromes <sup>19,37</sup> and 2 adult inflammatory bowel disease viromes <sup>31,32</sup>, and they 143 144 reported relative abundance shifts of ssDNA and dsDNA phages within these viromes. From this 145 observation, these studies concluded that such shifts could discriminate between healthy and disease 146 states associated with virome development in early life.

147 However, the abundance of ssDNA viruses can also be enriched from methodologies used in 148 making the viromes, even if all samples are processed consistently. Specifically, early virome studies 149 where limiting viral nucleic acids were obtained, often used whole genome amplification kits that 150 leverage a DNA polymerase from the phi29 ssDNA virus to obtain many-fold increases in DNA via multiple displacement amplification or MDA <sup>54</sup>. Though attractive at first, MDA is now known to have 151 stochastic biases (e.g., 100s –10,000s-fold biases in coverage, <sup>55,56</sup>), which result from randomized initial 152 153 template interactions and can induce chimera formation and uneven amplification of linear genomic 154 sections (whether ssDNA or dsDNA templates), as well as systematic biases resulting from preferential amplification of small, circular and ssDNA genomes <sup>57-61</sup>. Taken together, MDA-associated artifacts skew 155 156 the taxonomic representation of a community in non-repeatable ways and preclude quantitative analysis

of viromes <sup>57</sup>. Although non-quantitative, MDA-amplified viromes do still have value enriching for
ssDNA viruses, as well as estimating presence of viruses.

159 Consistent with the idea that these ssDNA viruses are methodologically enriched in the MDA 160 libraries, we found that non-MDA amplified gut viromes contained significantly less ssDNA viruses than 161 MDA amplified gut viromes (range: 0% - 4% versus 0-42%; Mann-Whitney U-test; *p*-value = 0.0083), though sample size was quite low. Further, while we see a strong linear relationship ( $R^2 = 0.86$ ) between 162 163 sequencing depth and the number of viral populations sequenced in non-MDA viromes, this relationship is weak in MDA viromes ( $R^2 = 0.39$ ), suggesting that MDA can skew the number of assembled viral 164 165 contigs in datasets (Supplementary Fig. 1b). Critically, 14 of the 21 studies gathered in this work 166 employed MDA, which calls into question the quantitative nature of these datasets. Fortunately, viral 167 nucleic acid extraction from feces often yield sufficient quantities for high throughput sequencing <sup>26</sup>, and 168 in cases where they do not there are now several viable alternative methods to more quantitatively 169 establish viromes with as little as 1pg of DNA <sup>61,62</sup>. Problematically, current established gut virome 170 protocols recommend an MDA step  $^{48,63}$ . If a researcher's goal is to provide quantitative datasets, then we strongly advocate against this recommendation and instead suggest that alternative methods<sup>61,62</sup> be used to 171 172 generate gut viromes.

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174 *Human gut virome study conclusions are more impacted by methodology than disease state* 

175 Given a systematically processed GVD, we next sought to determine whether global clustering patterns

176 would emerge between study themes between all dataset used to build GVD. To this end, viral

177 populations identified in this study were matched back to their respective datasets, and used in a co-

178 occurrence network analysis (see **Methods**) to assess co-variation at two levels: between study datasets

179 (Fig. 4c), and between viromes across all datasets (Fig. 4d). Between datasets, the fraction of shared viral

180 populations was low (mean:  $3\% \pm 3\%$ ; Fig. 4c), except for 6 datasets that clustered together (hierarchical

- 181 clustering bootstrap = 100%; Fig. 4c) and had a higher level of shared viral populations (>4-fold
- 182 increase). Presumably, these elevated similarities across the 6 datasets may be due to deeper sequencing

183 (Fig. 4d, top panel) that allowed deeper sequencing into the rare tail of viral populations among samples. 184 A similar trend was observed when looking at the level of individuals within each study (Fig. 4d), where 185 the co-occurrence network revealed close clustering between individuals derived from the same study, 186 irrespective of geographical origin, health status and/or diet. This per study clustering implies that, taken 187 together, these studies are not comparable likely due inconsistent sampling and extraction methodologies. 188 We then investigated the prevalence of gut viral populations amongst all samples, so as to establish whether any viral populations were detected in all samples (i.e., a 'core' gut virome<sup>22</sup>). On average,  $138\pm$ 189 190 170 (average  $\pm$  SD; range: 0 to 849) viral populations were detected per sample, but not one viral 191 population was found across all samples. We then explored deeper to detect whether subsets of the 192 samples would reveal shared viral populations. We found that only 28 viral populations occurred in over 193 20% of the GVD samples. Most viral populations were detected in very few samples. In fact, >40% of 194 the viral populations occurred in <0.5% of the samples and 98% of the viral populations occurred in 195 <0.1% of the samples in GVD (Fig. 5 a, b and Supplementary Table 3). Further, we specifically looked 196 at the prevalence of crAssphages, a well-recognized, multi-genera group of phages known to be 197 widespread in gut viromes<sup>64</sup> (Fig.5 b, c). While crAssphages are ubiquitous across the GVD samples. 198 there was not one crAssphage viral population found universally, with the most widespread crAssphage 199 population occurring in only 38% of samples. Importantly, when we looked at all healthy samples and 200 healthy western samples specifically, still no shared viral populations were identified in all samples. 201 (Supplementary Fig. 2a, b). Assuming samples were sufficiently sequenced, this may be indicative that 202 individuals carry a unique 'gut virome fingerprint', even between twins, which is perhaps not surprising 203 given recent suggestions of a similar 'fingerprint' for gut microbes (the 'personal' microbial microbiomes 204  $^{65}$ ). This apparent lack of core gut virome among individuals contrasts with a recent report  $^{22}$ , in which 205 overlapping patterns of phage genomes between 2 unrelated healthy individuals, as well as within a re-206 analyzed larger cohort  $^{66}$  revealed three levels of sharing patterns: (i) core (phage found in >50% of 207 samples, (ii) common (phage found in >20-50% of samples), and (iii) unique (phage found in <20% of 208 samples). Our analyses showed no viral populations shared above >50% of samples, thus bringing into

question the presence of a 'core' virome as previously defined<sup>22</sup>, as well as a very limited 'common' 209 210 virome (20-50% sharing across samples), in which we observed either 1% (all healthy; n=132) or 0.1% 211 (all healthy Westerners; n=18) of GVD viral populations, similar to the 3% previously reported<sup>22</sup> (see 212 Supplementary Table 4). Likely, this discrepancy with our results could be attributed to how viruses were identified through read mapping. In the initial study reporting a core virome<sup>22</sup>, a virus was 213 214 considered present if a single read mapped to a genome, a very permissive cut-off which does not take 215 into account shared homologous regions between distinct viral populations. In this study, we considered 216 a virus present if reads mapped 70% of the genome length (if genome is <5kb) or reads mapped at least 217 5kb of the genome (for genome >5kb in length) (see Methods). While our cut-off is more conservative, it 218 better ensures that we are detecting the same viral population. Nonetheless, the idea of a core virome 219 might still be an open question.

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221 *Re-evaluation of a previous study: the virome across different geographic regions and lifestyles* 

222 Due to the high level of sample clustering per study (Fig. 4c), we were unable to conduct cross-223 study analyses. Instead, we sought to assess if the virome community patterns between populations of 224 varying lifestyles (industrialized versus semi-industrialized versus hunter-gatherer) would vary between the initial study <sup>26</sup> or GVD-based, to test whether there were geographic biases around GVD viral 225 226 populations, and how well sampled are the different geographic regions. This initial study encompassed a 227 globally-distributed dataset (USA, Italy, Tanzania and two Peruvian populations: Tunapuco and Matses; 228 Fig. 6a), and explored the impact of geography and diet on eukaryotic gut viruses (but did not include 229 phages) and found that the hunter-gatherers (Hadza in Tanzania and the Matses in Peru) had the highest eukaryotic viral richness<sup>26</sup>. 230

In this re-analysis, however, we included phages in addition to eukaryotic viruses, and focused on how the virome diversity varied along the dataset. We first evaluated whether per-region GVD-mediated detection of viruses would incur biases, potentially stemming from underrepresented viral populations from less-sampled geographical regions. This did not appear to be the case, as significant increases in

235 virus detection were observed across 4 out of the 5 regions sampled (Fig. 6b). We next calculated 236 diversity indices (Fig. 6c and Supplementary Fig. 3) for each regional dataset, and looked at the number 237 of viral populations mapped with GVD. Overall, we reached a similar conclusion to the initial study (even 238 when considering phages), in which the hunter-gatherers (Peru Matses) generally contained higher viral 239 richness (Fig. 6c - left) and biodiversity (Shannon's H, Fig. 6c - middle), but not higher evenness 240 (Peilou's J, Fig. 6c -right). Collector's curves revealed that we have not saturated the human gut viral 241 diversity among individuals globally (Supplementary Fig. 4) or even among just among American 242 samples (Supplementary Fig. 2, inset). Thus, it appears much more viral diversity remains to be 243 discovered across all geographic regions. 244 We next wondered whether the addition of phage in our analysis would reflect on overall viral 245 community similarities by using Bray-Curtis distances between individuals across these geographic and 246 lifestyle gradients (Fig. 6d). While unequal database representation can have an impact on alpha-247 diversity, beta-diversity is often less impacted <sup>67</sup>. Principal coordinate analyses (PCoA) of Bray-Curtis 248 distances derived from using the individual Rampelli et al., 2017 virome database (Fig. 6d, left panel) and 249 GVD (Fig. 6d, right panel) revealed no significant differences (Mantel's test; R = 0.95, p = 0.001).

250 However, analysis of the GVD-referenced PCoA revealed individuals with the same lifestyle and from 251 the same region clustered together (PERMANOVA;  $p \leq 0.001$ ) and provided better resolution of the 252 clustering in comparison to the IV-referenced PCoA. However, lifestyle alone may not account for the 253 observed clustering patterns. The viromes of the Hadza in Tanzania and semi-industrialized, agrarian 254 Tunapuco population in Peru strongly overlapped (hierarchical clustering bootstrap = 100%; Fig. 6d), most likely driven by their diets rich in root vegetables<sup>68–70</sup>. Nonetheless, when we look at differences 255 256 between dominant viral populations (found in >50% individuals) across these geographic and lifestyle 257 gradients, we see that there are key viruses missing from Western, industrialized gut viromes (Fig. 6e), specifically viruses that infect the genus Prevotella spp. This parallels the bacterial analyses that show 258 259 that Prevotella spp. are enriched in non-Western gut microbiomes and many species are missing from

Western, industrial gut microbiomes <sup>69–71</sup>. Overall, this suggests that lifestyle and diet has an impact not
only on the bacterial community, but also on the viral community in the gut.

## 262 CONCLUSIONS

263 The lack of a curated database for the detection of viral sequences in the human gut has been 264 identified as the most critical shortcoming of applying metagenomic approaches to studying the human gut virome <sup>72</sup>. Although GVD is geared towards filling this gap and performs well (increasing viral 265 266 detection 59-fold over the most commonly used database, NCBI viral RefSeq), there are limitations. First, 267 the geographic and ethnic representation across the dataset is not very broad. Meta-analyses will benefit 268 from more broadly representative datasets. Second, GVD was built using all datasets available by the end 269 of 2017. Since then, as of May 2019, there are 11 additional datasets that study the gut virome, 8 of which 270 use viral particle-enriched metagenomes (Supplementary Table 5). Further, there are many more human 271 gut microbial metagenomic datasets and these could be a rich source for virus reference genomes as found for soils<sup>73</sup> and the large-scale Earth Virome study<sup>74</sup>. To maintain significance as a resource, we will 272 273 update GVD annually by extracting the viral signal from such gut-related datasets, as well as monitoring 274 IMG/VR for gut-related viruses that should be integrated. *Third*, GVD is accessible through direct 275 download as a single fasta file containing all GVD viral populations (see link in the 'Data availability' 276 statement below), and is likely best paired with IMG/VR to maximize viral signal recovery. Future GVD 277 updates and development will be required to improve the user experience for those not comfortable at 278 command-line interfaces, but these are likely best integrated with large-scale standardizing efforts like the 279 National Microbiome Initiative.

Given the relatively minimal value added via non-quantitative MDA-based approaches and the availability now of low-input quantitative approaches pioneered studying ocean viruses <sup>61,75</sup> suggest that gut virome studies should move away from the former towards the latter. GVD, combined with the means to classify uncultivated virus genomes<sup>47</sup>, are prime starting requirements for enabling ecosystem-wide examinations<sup>76</sup> of the dynamics and impacts of the virome within the human gut. Other environmental advances also invite such studies to include assessing the role of micro- and macro-diversity on virus

- 286 persistance<sup>41</sup>, and metabolic reprogramming via virus-encoded auxiliary metabolic genes<sup>73,76</sup>. These
- 287 combined efforts are critical to enable studies of the human gut virome to advance from 'stamp
- 288 collecting' diversity studies towards the kinds of comprehensive efforts needed to incorporate viruses into
- 289 mechanistic, predictive models. Such efforts, with future viral mapping outside the gut to parallel efforts
- for the 'non-gut' human microbiome <sup>77</sup>, should help transform personalized medicine and lead to a better
- 291 understanding of human ecosystems.

# **293 FIGURE LEGENDS:**

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295 Figure 1. Overview of studies and meta-analyses comprising the Gut Viral Database (GVD). (a) 296 Global heatmap of the world showing the number of individual's gut viromes coming from different 297 countries within the GVD. Importantly, individual's viromes coming from the Cameroon were pooled 298 based on their location, age, and contact with bats. The pools were counted as a single individual's virome 299 for our analyses. (b) Pipeline for the selection and processing of human gut virome datasets (see 300 **Methods**). Datasets were processed individually and, within each dataset, viromes were pooled by 301 individual, except for fecal microbiota transfer (FMT) studies and data that was given to us prior to 302 publication (Yinda et al., 2019; Neto et al. (unpublished)). Reads were filtered for quality and trimmed 303 and reads that mapped to  $\Phi x 174$  and the human genome were removed. The remaining reads were 304 assembled into scaffolds, filtered for lengths  $\geq 1.5$ kb, and run through tools that collectively utilize 305 homology to viral reference databases, probabilistic models on viral genomic features, and viral k-mer 306 signatures to identify viral contigs. Viral contigs were then deduplicated to get a total of 13,203 viral 307 populations.

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**Figure 2. The Gut Viral Database (GVD)**. (a) Pie charts showing the number of bacteriophages,

310 eukaryotic viruses, and archaeal viruses in the GVD (center) and their familial taxonomic composition by

311 the bacteriophages (left) and the eukaryotic viruses (right). (b) Gene-sharing taxonomic network of the

312 GVD, including viral RefSeq viruses v88. RefSeq viruses are highlighted in red. Every node represent a

313 virus genome, while connecting edges identify significant gene-sharing between genomes, which form the

basis for their clustering in genus-level taxonomy. (c) Bar chart showing the number of bacterial host

315 phyla of the GVD bacteriophages, with an inset providing resolution for the low frequency bacteria host

316 phyla. Putative host phyla per each bacteriophage population are in **Supplementary Table 3** 

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Figure 3. GVD as a reference database increases viral population detection. (a) Boxplots showing median and quartiles of the number of viral populations detected per study using the IV, Viral Refseq v88, JGI IMG/VR, or GVD databases. Studies where the reads were given to us prior to publication are excluded from this analysis (Yinda et al., 2019; Neto et al. (unpublished)). (b) Venn diagram showing the number of viral populations unique and shared between the different databases. Importantly, we only compared dereplicated viral populations from IMG/VR that came directly from human gut samples or had reads mapping to them from GVD gut samples.

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Figure 4. Individual Viromes (IV) Study Databases and Cross-Study Comparisons. (a) Barplot
 showing the proportion of those viruses that are bacteriophages, archaeal viruses, or eukaryotic viruses.

snowing the proportion of those viruses that are bacteriophages, archaeal viruses, or eukaryotic vi

328 The total number of assembled viral contigs and viral populations per study are available in 329 Supplementary Fig. 1a. (b) Barplot showing the proportion of those viruses that are dsDNA, ssDNA, or 330 RNA viruses. Studies where multiple displacement amplification (MDA) was used show a higher 331 prevalence of ssDNA viruses. No viral contigs  $\geq$ 1.5kb were assembled from the Reves *et al.* 2010 study. 332 (c) Hierarchically clustered heatmap showing the number of viral populations shared within and between 333 studies. The barplot on top of the heatmap shows the total number of sequenced base pairs following 334 quality control within each study. (d) Viral population co-occurrence network per individual within each 335 study shows that individuals within a study cluster together regardless of health status. The squares 336 represent the healthy individuals within each study.

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Figure 5. There are no core viral populations across GVD samples. (a) Histogram showing the number of viral populations present in different percentages of GVD samples. The vast majority of viral populations are found in <10% of the individuals. (b) Hive plot showing the percentage of GVD sample each viral population is detected within. The dots on the x-axis represent each GVD viral population in ascending order of the percentage of GVD samples that they are found within. The y-axis is the percentage of GVD samples that each viral population is detected within. CrAssphage viral populations are highlighted in red. (c) Heatmap showing the presence or absence of each crAssphage viral populationacross the different GVD samples.

346 347 Figure 6. Diet and geography widely influence gut virome. (a) World map showing the geographical 348 distribution of the Rampelli et al., 2017 dataset. (b) Boxplots showing median and quartiles of the number 349 of viral populations detected using the GVD database and the Rampelli et al., 2017 viral database alone 350 (IV) within each geographic group. (c) Boxplots showing median and quartiles of the  $\alpha$ -diversity metrics 351 - richness, Shannon's H and Peilou's J – across the different geographic groups using the GVD database 352 (see Fig. S3 for  $\alpha$ -diversity metrics using the IV database). (d) Principal coordinate analysis (PCoA) of a 353 Bray-Curtis dissimilarity matrix calculated from mapping the Rampelli et al., 2017 dataset against the IV 354 (left) and GVD (right) databases. Analyses show that the virones significantly (Permanova p < 0.05) 355 structure into based on the geographic groups, with mapping to the GVD showing revealing much 356 stronger clustering based on geography. Ellipses in the PCoA plot are drawn around the centroids of each 357 group at a 95% confidence interval. The dashed lines connecting the different points reveal the 358 connections determined by hierarchically clustering between the different samples. (e) Heatmap of the 359 abundances of the viral populations found across >50% of individuals within the study. Individuals on a 360 Western diet (from the USA and Italy) lack phages that infect Bacteroidetes, specifically those that infect 361 *Prevotella* sp. All pairwise comparisons were performed using a two-tailed Mann-Whitney U-tests. 362

Supplementary Figure 1. Number of assembled viral contigs and populations. (a) Barplot showing
 the number of assembled viral contigs versus the number of deduplicated viral populations per study. (b)
 Scatterplots with linear regressions showing the impact of increased sequencing on the number of
 assembled contigs per study divided by studies that did not have multiple displacement amplification
 (MDA; top) and those that did have MDA (bottom).

Supplementary Figure 2. There are no core viral populations across healthy samples and across healthy western samples. Hive plots showing the percentage of GVD samples each viral population is detected within across (A) all healthy individuals and (B) across only healthy western adults. The dots on the x-axis represent each GVD viral population in ascending order of the percentage of GVD samples that they are found within. The y-axis is the percentage of GVD samples that each viral population is detected within.

Supplementary Figure 3. Boxplots showing median and quartiles of the α-diversity metrics – richness,
Shannon's H and Peilou's J – across the different geographic groups in the Rampelli *et al.* 2017 study
using the IV and GVD databases.

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Supplementary Figure 4. The number of gut viral populations will still increase with more samples
 added to GVD. Collector's curve for gut viral populations in the GVD. (inset) Collector's curve for just
 the viromes from samples from the USA.

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384 Supplementary Table 1. Origin of datasets and associated metadata used to create the gut virome
 385 database.
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**Supplementary Table 2.** Gut Viral Database contigs family-level taxonomy and putative hosts.

388389 Supplementary Table 3. Distributions of viral populations across GVD samples.

- **391** Supplementary Table 4. Core, common, low-overlap, and unique GVD viral populations
- **393** Supplementary Table 5. Human gut virome studies since the end of 2017

#### 394 **METHODS**

395 Experimental Model and Subject Details. Gut virome database (GVD) studies were selected by doing a 396 thorough and manually curated search of the Web of Science Core Collection of Thomson Reuters for 397 studies looking at viruses in the gut published prior to 2018. All studies that used next-generation 398 sequencing and looked for viruses within the gut microbiome were selected to be part of GVD (see full 399 list of studies in **Supplementary Table 1**). Additionally, we were given access to the reads of two studies 400 that were unpublished at the time. One of the studies, however, is now published (Yinda et al., 2019). 401

402 Viral contig assembly, identification, and dereplication. Previously published GVD reads were 403 downloaded from their respective hosting databases (e.g. SRA, iVirus, or MG-RAST). Prior work 404 revealed that an individual's gut virome is stable across time (Minot et al., 2013), so reads were pooled 405 per individual regardless of the number of time points, with a few exceptions (Fig. 1). These exceptions 406 included studies with fecal microbiota transfers and studies whose reads were given to us prior to 407 publication. For fecal microbiota transfers, all time points per individual were kept separate and processed 408 independently. Read sets from two studies were given to us prior to publication (Yinda et al., 2019; Neto 409 et al., unpublished). For the Yinda et al., 2019 study, individual's reads were pooled based on their 410 location, age, and contact with bats. The pools were counted as a single individual's virome for our 411 analyses. For the Nadia et al., (unpublished), all reads from all individuals were pooled together. A global 412 map showing the number of individuals (or pooled read sets) originating from each country was created 413 using the R packages 'rworldmap.' In total, there were 648 GVD samples from 572 individuals.

Pooled reads were then assembled using metaSPAdes 3.11.1<sup>78</sup>. Following assembly, contigs 414 >1.5kb were piped through VirSorter<sup>79</sup> and VirFinder<sup>80</sup> and those that mapped to the human, cat or dog 415 416 genomes were removed. For viral-enriched metagenomes (i.e. viromes), contigs >5kb or >1.5kb and 417 circular that were sorted as VirSorter categories 1-6 and/or VirFinder score  $\geq 0.7$  and p < 0.05 were pulled 418 for further investigation. Of these contigs, those sorted as VirSorter categories 1 and 2, VirFinder score 419  $\geq$ 0.9 and p <0.05 or were identified as viral by both VirSorter (categories 1-6) and VirFinder (score  $\geq$ 0.7 and p < 0.05) were classified as viral. The remaining contigs were run through CAT <sup>81</sup> and those with 420 421 <40% (based on an average gene size of 1000) of the genome classified as bacterial, archaeal, or 422 eukaryotic were considered viral. For the microbial metagenomes, we took a more conservative approach 423 with only contigs >5kb or >1.5kb and circular that were sorted as VirSorter categories 1-2 and VirFinder 424 score >0.6 and p < 0.05 were considered viral. Across the both the viral-enriched and microbial 425 metagenomes, contigs  $\geq$ 5kb or  $\geq$ 1.5kb and circular that were classified as eukaryotic viral contigs by 426 CAT were also considered viral. In total, 29,345 viral contigs were identified.

427 Viral contigs that were from known ssDNA or RNA viral families using CAT were grouped into 428 populations if they shared  $\geq$ 95% nucleotide identity across  $\geq$ 100% of the genome. Because there are no 429 benchmarked metagenomic population boundaries for ssDNA and RNA viral families, we chose to not 430 use stringent dereplication. All other contigs were considered double-stranded DNA and were grouped into populations if they shared  $\geq$ 95% nucleotide identity across  $\geq$ 70% of the genome (*sensu*<sup>82</sup>) using 431 nucmer<sup>83</sup>. All the viral contigs that were assembled were dereplicated per study to create the individual 432 433 virome (IV) databases and across all of GVD (see Supplementary Fig. 1). For GVD, this resulted in 434 13,203 total viral populations found in GVD (see Supplementary Table 3 for VirSorter, VirFinder, and CAT results), of which 6,373 were  $\geq 10$ kb in length. 435

437 Core Viral Population Analyses. To explore if there were any core viral populations, the abundance 438 table was turned into a binary presence-absence matrix. The number of GVD samples that each viral 439 population was detected within was then calculated using R and divided by the total number (648) to get 440 the percentage of samples. Each viral population's percentage was plotted in hive plot using 441 'geom\_curve' in ggplot2<sup>84</sup>. This process was repeated on subsets of the matrix including all healthy 442 individuals and only the healthy western adults. The number of viral populations that were present across 443 different percentages were calculated using R and their distributions plotted using 'geom\_histogram' in ggplot2<sup>84</sup>. CrAssphage viral populations in GVD were identified using CAT results and by dereplicating 444 445 GVD viral populations with the crAssphage genomes identified in Guerin et al.<sup>64</sup> and seeing which GVD 446 genomes cluster. In total, there were 95 unique crAssphage populations. The binary presence-absence 447 data for the crAssphage populations were plotted using pheatmap in R.

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Viral taxonomy. For each viral population, ORFs were called using Prodigal<sup>85</sup> and the resulting protein 449 sequences were used as input for vConTACT2<sup>47</sup> and for BLASTp. Double-stranded DNA viral 450 451 populations represented by contigs >10kb were clustered with Viral RefSeq release 88 viral genomes using vConTACT2. Those that clustered with a virus from RefSeq based on amino acid homology based 452 on DIAMOND<sup>86</sup> alignments were able to be assigned to a known viral taxonomic genera. For viral 453 454 dsDNA populations that could not be assigned taxonomy or were <10kb, family level taxonomy was 455 assigned using a majority-rules approach, where if >50% of a genome's proteins were assigned to the 456 same viral family using a blast bitscore >50 with a Viral RefSeq virus, it was considered part of that viral family (see Supplementary Table 3 for family-level taxonomy). For ssDNA and RNA viruses, 457 458 CAT was used to assign the viral family (see **Supplementary Table 3** for family-level taxonomy). 459

- 460 Viral Host Prediction. Bacteriophage hosts were predicted using a variety of bioinformatic methods 461 including: (i) CRISPR-spacer matches, (ii) prophage blasts, (iii) tRNA genes matches, and (iv) WiSH matches<sup>87</sup> against Bacterial Refseq v88. CRISPR spacers were predicted using MinCED 462 463 (https://github.com/ctSkennerton/minced) and the CRISPR Recognition Tool (CRT<sup>88</sup>) and a BLASTn (-464 task blastn-short -word\_size 5) was used to assess matches between the CRISPR spacers and viral 465 populations in GVD. Those with 1 mismatch were considered a match. For prophage blasts, a blastn of 466 the viral population against Bacterial RefSeq was performed. A bacterial genome with  $\geq$ 2500bp regions 467 of their genome matching at 95% ID with a viral population genome were considered putative hosts of that viral population (see <sup>76</sup>). Viral tRNA genes and Bacterial RefSeq tRNA genes were predicted using 468 tRNA-scan<sup>89</sup> and then a blastn was performed between the viral and bacterial tRNA genes. Bacterial 469 470 tRNA genes that matched viral tRNA genes at 95% ID across 100% of the length were considered putative bacterial hosts. Lastly, WIsH was used to predict hosts according to default settings <sup>87</sup>. Priority 471 host assignment was given to CRISPR, then prophage, WIsH and tRNA results. Viruses with putative 472 473 archaeal hosts were predicted using MarVD<sup>90</sup>. Viruses with predicted eukaryotic hosts were assigned 474 based on their assigned taxonomic viral family. 475
- 476 **Detecting viral populations and calculating their raw abundances.** To calculate the raw abundances of
- the different viral populations in each sample, reads from each GVD pooled read set were first non-
- 478 deterministically mapped to all GVD viral population genomes using bowtie2. Further, reads from each
- 479 GVD pooled read set per study were mapped to their respective IV databases. BamM
- 480 (https://github.com/ecogenomics/BamM) was used to remove reads that mapped at <95% nucleotide

identity to the contigs, bedtools genomecov<sup>91</sup> was used to determine how many positions across each 481 482 genome were covered by reads, and custom Perl scripts were used to further filter out contigs without 483 enough coverage across the length of the contig. All contigs  $\leq 5kb$  in length with >70% of the contig 484 covered were considered detected in the sample. Contigs >5kb in length with  $\ge 5kb$  in length covered were also considered detected in the sample<sup>92</sup>. BamM was used to calculate the average read depth ('tpmean' -485 minus the top and bottom 10% depths) across each detected contig. For the alpha-diversity calculations, 486 487 the average read depth was used as a proxy for abundance and normalized by total read number per 488 metagenome to allow for sample-to-sample comparison. However, because most of the studies in GVD 489 involved MDA, which can skew abundances, we chose to use only a presence-absence statistic (richness) 490 for most of our  $\alpha$ -diversity calculations. Collector's curves and the whole GVD and across only American 491 samples were calculated using the function 'specaccum' in the R 'vegan' package <sup>93</sup>.

492

493 Comparisons to IMG/VR, Viral RefSeq v88, and IV databases. The IMG/VR (1.1.2018 release) 494 included all viral contigs assembled from different datasets. All of the viral contigs in GVD, Viral Refseq 495 v88, and IV databases are dereplicated at the population level. In order to make IMG/VR comparable to 496 GVD, Viral Refseq and IV databases, we needed to dereplicate the IMG/VR database. IMG/VR (1.1.2018 497 release) is composed of 715,672 contigs. Because dereplication is extremely computationally intensive, 498 we decided to only focus on dereplicating viral contigs that originated from the human gut and had at 499 least 1 read from a GVD metagenome map. These IMG/VR viral contigs were then dereplicated using the 500 same methodology as previously described in the methods section. In total, 29,378 IMG/VR viral contigs 501 were dereplicated into 6,652 viral populations. GVD pooled read sets were mapped to this IMG/VR 502 human gut viral population database, Viral RefSeq v88, and the IV databases for each individual study in 503 GVD. The raw abundances of the different IMG/VR and Viral RefSeq viral populations in each sample 504 were calculated the same way as described in the previous section. The total number of viral populations 505 detected per sample per study using the different databases were then plotted and comparative statistics 506 using the 'ggboxplot' function from the 'ggpubr' package in R.

507 All of the viral populations from GVD, the dereplicated IMG/VR gut-specific dataset, and Viral 508 Refseq were then dereplicated to see how many viral populations overlapped between databases. The 509 results were then plotted using the 'VennDiagram' package in R. Importantly, in the dereplication 510 process, some of the original viral populations in each database may be dereplicated down due to the 511 presence of a longer viral contig from the same population that links the two together into the same population. Across the databases, 329, 177, and 459 viral populations were dereplicated in GVD. 512 513 IMG/VR, and Viral Refseq, respectively. This is why the total number of populations displayed in the 514 Venn diagram does not add up to the total number of viral populations in each database.

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516 Clustering studies based on shared viral populations. To test how studies clustered together, the viral 517 population presence-absence data from individuals (or pooled read sets) within a study were merged. In 518 Study 1, individual A had viral population 1, 2, 4, 5 and individual B had viral population 3, then Study 1 519 had viral populations 1, 2, 3, 4, and 5. The different studies were then assessed for the number of shared 520 viral populations that were present in both studies. These values were then displayed and hierarchically 521 clustered using the R 'pheatmap' package and the stability of the hierarchical clusters were assess using 522 the R 'pvclust' package. The number of shared viral populations between individuals (or pooled read sets 523 within a sample) were clustered using the R 'SPIEC-EASI' package <sup>94</sup> using the Meinshausen and

524 Bühlmann (MB) method to infer associations between samples based on the shared number of viral525 populations. The network was plotted using the R 'igraph' package.

526

527 Alpha- and Beta-Diversity calculations. The  $\alpha$ - (Richness, Shannon's *H*, and Peilous' *J*) and  $\beta$ - (Bray-Curtis dissimilarity) diversity statistics were performed using VEGAN <sup>93</sup> in R. For all studies, except for 528 Rampelli et al.<sup>26</sup>, only richness was calculated for both abundances based on read mapping to IMG/VR, 529 Viral Refseq, the IV databases and GVD. Comparisons were plotted using 'ggboxplot' function in the R 530 'ggpubr' package. The Rampelli et al.<sup>26</sup> did not use MDA, so we went ahead with scaling the raw 531 532 abundances based on the number of quality controlled base pairs sequenced to normalize the data. All  $\alpha$ -533 diversity statistics were calculated and  $\beta$ -diversity was used to look at community structure using both the 534 IV and GVD databases. Principal Coordinate analysis (function capscale of VEGAN package with no 535 constraints applied) was used as the ordination method to plot the Bray-Curtis dissimilarity matrices 536 (function vegdist; method "bray") after a cube root transformation (function nthroot; n = 3). To 537 determine if the Rampelli et al. samples clustered by geographic region, a permanova test (function 538 "adonis") and the 95% confidence interval were plotted using function "ordiellipse." Further, the samples 539 were hierarchically clustered and plotted within the PCoA. To specifically look at abundance differences in the most abundant viral populations in the Rampelli et al.<sup>26</sup> study, viral populations that were present 540 541 in 50% study individuals and their hosts information were plotted using the R 'pheatmap' package. 542 543 Code availability. Scripts used in this manuscript are available on the Sullivan laboratory bitbucket under 544 Gut Virome Database.

545

546 Data availability. All raw reads are available through SRA, iVirus, or MG-RAST using the identifiers

547 listed in **Supplementary Table 1**. GVD viral populations can be downloaded directly from iVirus

- 548 through the following link: <u>https://de.cyverse.org/dl/d/E83EFBFF-2A23-4794-8819-</u>
- 549 ADD34160D018/FINAL\_Gut\_Viral\_Database\_GVD\_1.7.2018.fna
- 550

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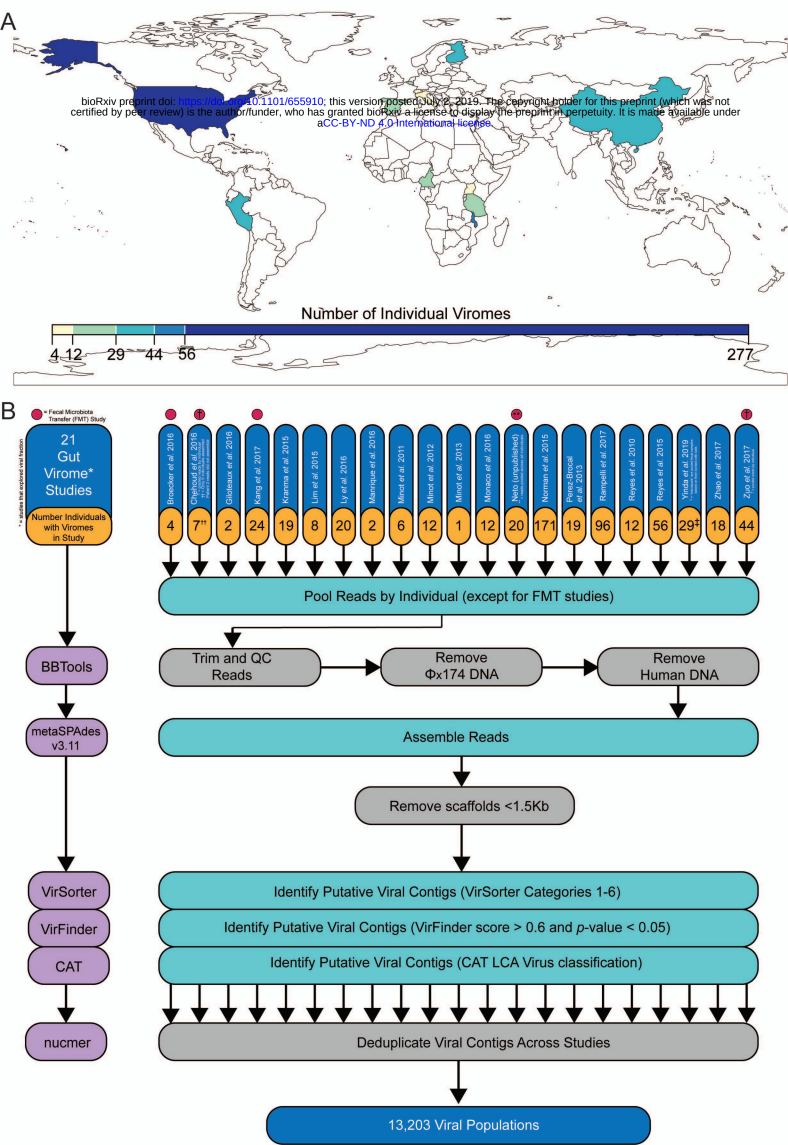
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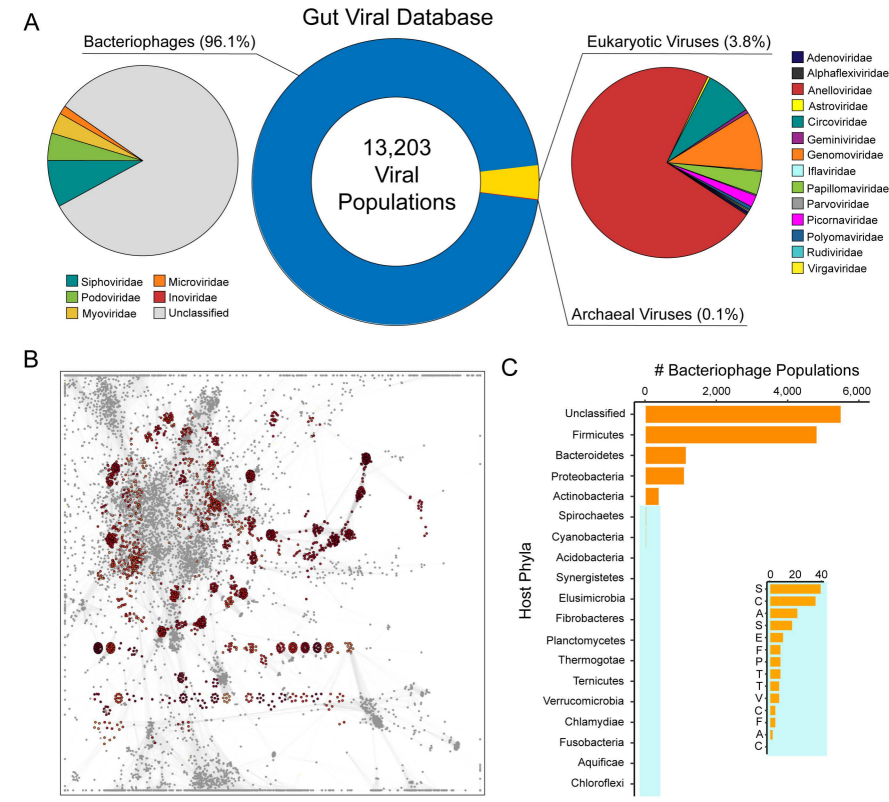
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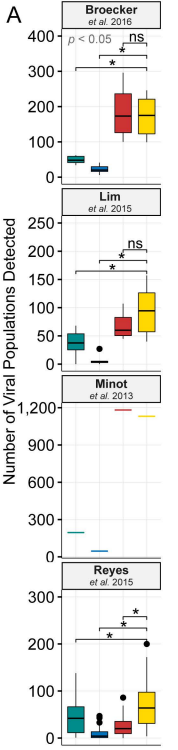
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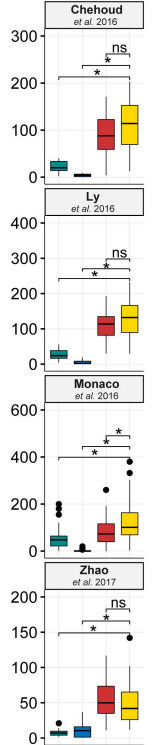
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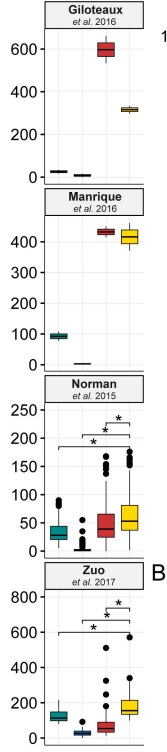
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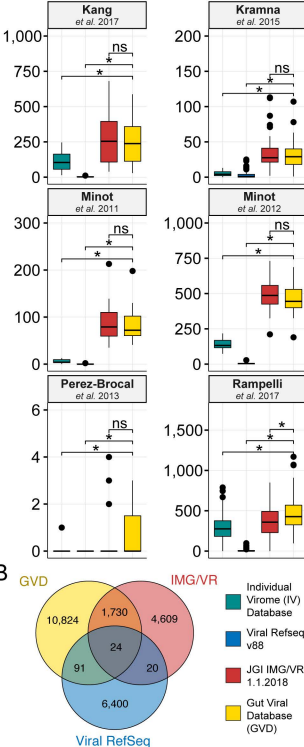












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