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# Viral biogeography of the mammalian gut and parenchymal organs

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The mammalian virome has been linked to health and disease but our understanding of how it is structured along the longitudinal axis of the mammalian gastrointestinal tract (GIT) and other organs is limited. Here, we report a metagenomic analysis of the prokaryotic and eukaryotic virome occupying luminal and mucosa-associated habitats along the GIT, as well as parenchymal organs (liver, lung and spleen), in two representative mammalian species, the domestic pig and rhesus macaque (six animals per species). Luminal samples from the large intestine of both mammals harboured the highest loads and diversity of bacteriophages (class *Caudoviricetes*, family *Microviridae* and others). Mucosal samples contained much lower viral loads but a higher proportion of eukaryotic viruses (families *Astroviridae*, *Caliciviridae*, *Parvoviridae*). Parenchymal organs contained bacteriophages of gut origin, in addition to some eukaryotic viruses. Overall, GIT virome composition was specific to anatomical region and host species. Upper GIT and mucosa-specific viruses were greatly under-represented in distal colon samples (a proxy for faeces). Nonetheless, certain viral and phage species were ubiquitous in all samples from the oral cavity to the distal colon. The dataset and its accompanying methodology may provide an important resource for future work investigating the biogeography of the mammalian gut virome.

he gastrointestinal tracts (GIT) of humans and other mammals contain highly individualized microbiomes<sup>1-4</sup>, composed of bacteria<sup>5</sup>, archaea<sup>6</sup>, eukarvotic microorganisms<sup>7</sup> and viruses<sup>8-10</sup>. The close association of microbes and their mammalian host as an ecological unit is increasingly recognized as important for health<sup>11,12</sup>. The gut virome, which is largely composed of bacterial viruses (bacteriophages or phages) remained a relatively unexplored area until recently, when a potential role for the virome in shaping bacterial communities was postulated<sup>13-16</sup>. A number of potential mechanisms by which such shaping could occur have been suggested, and include 'kill-the-winner' dynamics in bacterial communities caused by phage predation (at least at strain and substrain level<sup>17</sup>); diversifying selection acting upon both adaptive mutations<sup>18,19</sup> and phase variations<sup>20,21</sup>; as well as phage-mediated horizontal gene transfer that could involve diverse mechanisms such as generalized, specialized and lateral transduction<sup>22-24</sup>.

Our current understanding of the virome, and the phageome in particular, is limited and based mostly on sequencing-based studies of faecal samples, which represent static snapshots of the distal gut virome. Neither the temporal dynamics  $^{17}$ , nor the variation and flux of viral populations along the longitudinal and transverse axes of GIT (the 'viral biogeography' of the gut  $^{25,26}$ ), have received proper attention. Recent human cohort studies highlighted a tight association between the gut virome and gut bacteriome in terms of both  $\alpha$ - and  $\beta$ -diversity  $^{17,27}$ . In addition, multiple lines of evidence suggest that many successful gut bacteriophages, such as the crAss-like phages, engage in long-term persistent relationships with their hosts  $^{28,29}$ , in line with the 'piggyback-the-winner' dynamics of temperate bacteriophages  $^{30}$ . It is important to obtain a more detailed

view of both the temporal and spatial dynamics of the virome to understand its interplay with the bacterial microbiome, its importance for human health and its potential role in disease<sup>31,32</sup>.

Complex macro- and micro-anatomy of the digestive tract, together with the exocrine functions of GIT mucosa and accessory organs create a series of longitudinal and radial biochemical gradients, affecting the composition of local resident microbiota, including viruses<sup>25,33,34</sup>. Adaptation to such microhabitats is clearly evident among bacteria, such as body site-specific lactobacilli35 or various mucin-foraging bacteria<sup>36</sup>. Host-associated mutualistic and commensal bacteria have evolved persistence mechanisms such as adsorption and embedding into mucus layers, and potentially have access to anatomical sites protected from the luminal stream and the action of bacteriophages<sup>25,34</sup>. Similarly, the ability to bind to and accumulate in the mucous layer and potentially restrict bacterial invasion has also been reported for certain bacteriophages, which prompted a discussion on the role of bacteriophages as a quasi-immune system of the digestive tract<sup>37–39</sup>. Pronounced physiological and anatomical differences between homologous GIT segments in different species of mammals, associated with digestive adaptations, adds another layer of complexity to this system<sup>40</sup>.

In this study, we present a comprehensive biogeographical analysis of viruses in the GIT of two mammalian species, the domestic pig (Sus scrofa domesticus) and rhesus macaque (Macaca mulatta), chosen for their phylogenetic, physiological and anatomical relevance for humans. We focus our attention on bacteriophage populations and attempt to answer two key questions. First, what are the differences in virome composition between different digestive tract regions and how representative are distal gut samples of the virome

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in the upper GIT? Second, to what extent can the virome be shared between the digestive tract regions and with extra-GIT organs?

#### Results

Virome sequencing approach. We applied shotgun metagenomic sequencing of virus-like particle (VLP) enriched samples<sup>41-43</sup>, to characterize luminal/mucosal viral DNA and RNA content in different locations along the digestive tract. To adopt a broader taxonomic outlook and gain insights into spatial virome organization that go beyond the physiological and anatomical specifics of a particular mammalian species, we included six healthy domestic pigs (Sus scrofa domesticus) and six rhesus macaques (Macaca mulatta). Thirteen anatomical locations were sampled for each species: skin, tongue, stomach, small intestine (SI; proximal (duodenum), medial (jejunum) and distal (ileum)), caecum, large intestine (LI; proximal, medial and distal colon), as well as parenchymal organs (liver, lung and spleen). At relevant sites, both the mucosal and the surrounding luminal content were sampled (Fig. 1). Given the overwhelming prevalence of bacteriophages in mammalian faecal viromes<sup>17,41</sup>, their possible role in shaping the gut bacteriome<sup>16,19,44</sup> and a lack of knowledge on their spatial distribution and populations dynamics in the GIT<sup>9,34</sup>, bacterial viruses were the primary focus of this study.

Genomic DNA and complementary DNA of mixed viral populations were sequenced using the Illumina NovaSeq platform to a depth of  $6.2 \pm 5.8 \,\mathrm{M}$  per sample (median  $\pm$  interquartile range (IQR); Supplementary Information). Unlike many previous studies, our viral metagenomics approach was designed to be relatively unbiased. A simple nucleic acid extraction procedure was adopted that deliberately avoided the use of microfiltration, VLP precipitation using polyethylene glycol/NaCl, chloroform extraction or density gradient ultracentrifugation, all of which are known to introduce different biases in virome profiling studies<sup>42,43,45</sup>. By avoiding whole-genome amplification we also avoided artificial virome composition skewness, loss of viral diversity and overamplification of small circular single-stranded DNA genomes<sup>17,46</sup>. Lastly, including lactococcal phage Q33 as an artificial internal viral standard in our extraction procedure allowed us to estimate the abundance of viral genomes in the sample by comparing their mean sequence coverage with that of the internal standard<sup>17,43</sup>.

Assembly of reads into contigs47, removal of redundancy across individual samples and animals<sup>17</sup>, and selection of viral sequences from a bacterial and mammalian host DNA background yielded of catalogue of 107,680 contigs corresponding to putative complete and fragmented viral genomes (Extended Data Fig. 1). At least 24 families of prokaryotic and eukaryotic viruses<sup>48,49</sup> were recognized across the viromes of the two mammalian species using an automated taxonomic assignment algorithm (Extended Data Fig. 2). Approximately half (58,573) of the contigs were broadly similar (≥50% sequence identity over 85% of contig length) to previously reported genomes of either cultured or uncultured viruses<sup>50-55</sup>, but the remaining half were only identifiable as viral using a de novo multiclassifier approach<sup>56</sup>. However, even within sequences homologous to previously reported viral genomes and genome fragments, the majority (31,032) constitute unclassified viral species by the recently proposed standard of metagenomic viral species delineation ( $\geq$ 95% sequence identity over  $\geq$ 85% of its length)<sup>57</sup>.

**Absolute viral counts along the GIT proximal—distal axis.** Absolute quantitation of viral genomic contigs with  $\geq$ 50% calculated completeness level (n=2,442), grouped at viral family level, revealed pronounced differences in the virome between GIT locations, as well as the differences between the two animal species. The pig LI lumen is dominated by tailed bacteriophages (class *Caudoviricetes*, including crAss-like phages<sup>9,53,58</sup>) with total viral loads exceeding 10° genome copies per g of contents. Similar total counts are evident in macaques, although small ssDNA *Microviridae* phages<sup>59</sup> are the

most numerous group of taxonomically classified viruses (Fig. 1). Total viral loads in LI mucosa samples were two orders of magnitude lower than matched luminal samples, and eukaryotic viruses (families *Circoviridae*, *Astroviridae*, *Calicivirdae* and *Parvoviridae*) had higher relative weights in those locations. Stomach and SI lumen and mucosa were colonized by relatively even mixes of bacteriophages and eukaryotic viruses, with a characteristic prevalence of *Parvoviridae* in the pig SI mucosa. Similar combinations of viral families were detectable in tongue mucosa and skin samples in both animal species.

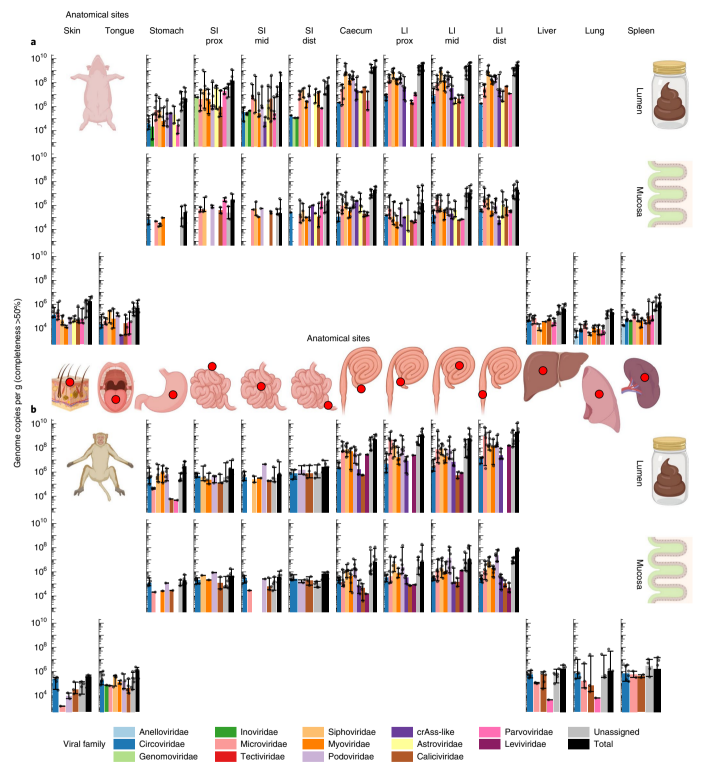
Interestingly, samples taken from lung, spleen and liver parenchyma in both species contained unexpectedly high viral loads, approaching and exceeding 10<sup>6</sup> genome copies per g of tissue. In macaques, these viral populations that are apparently associated with the interior body milieu of healthy animals, were mainly represented by eukaryotic viruses of the *Circoviridae* and *Caliciviridae* families. In both species, and especially in pigs, the viral consortia of interior milieu included bacteriophages, primarily from the *Microviridae* family (Fig. 1).

We then used all 107,680 viral contigs, both high quality and highly fragmented<sup>57</sup>, to identify compositional virome differences between different body sites in both animal species (Fig. 2 and Extended Data Fig. 1). Although highly fragmented viral contigs are less useful for taxonomic classification and host identification purposes<sup>17,32</sup>, omitting them from diversity analyses would leave the majority of viral diversity untapped (>50% of all Illumina reads from most body sites) (Supplementary Information). To compensate for interindividual virome differences and make the virome more comparable across animal cohorts we used gene sharing networks<sup>60</sup> to group all non-singleton viral genomic contigs (n=12,262) into 3,770 viral clusters (VCs).

Virome composition along the GIT proximal-distal axis. Multivariate virome comparison, based on the fractional abundance of VCs at different sites, revealed a strong separation of LI viromes from the SI and gastric viromes in both animal species (Fig. 2a and Extended Data Fig. 3). When viewed across the two species, differences between organs were responsible for 11.1% of the variance (Adonis with 1000 permutations;  $P \le 0.001$ ). Surprisingly, interindividual virome differences accounted for 9.6% of the variance, higher than the per cent variance explained by animal species (4.9%;  $P \le 0.001$ ). This is despite the fact that within each cohort animals were relatively inbred, lived in the same facility and were fed with a standardized diet. Moreover, between-organ variance in interaction with the individual animal factor accounted for 30.0% of the virome data variance ( $P \le 0.001$ ), much higher than the per cent variance explained by a similar interaction between organ and animal species factors (9.4%;  $P \le 0.001$ ). Differences between mucosal and luminal virome explained only a relatively minor fraction of the variance (1.0% for the main effect, 1.9% in the interaction with organ factor;  $P \le 0.001$ ). The major compositional separation between viromes of LI, SI and other organs seems to be closely aligned with overall diversity and total viral load ( $P \le 0.001$  in permutational multivariate analysis of variance), with caecal and LI viromes being simultaneously the most taxonomically diverse and the most populous (Figs. 1 and 2a-c).

In a single macaque (M6) and pig (E6), all mucosal sites were sampled twice, with 1 cm of separation between each pair of samples, to assess whether close proximity of mucosal sites in the gut correlates with increased similarity of the virome composition. In both SI and LI there was a tendency for these paired samples to resemble each other more closely than more distant sites within these and other animals, but this did not reach the level of statistical significance (Supplementary Information).

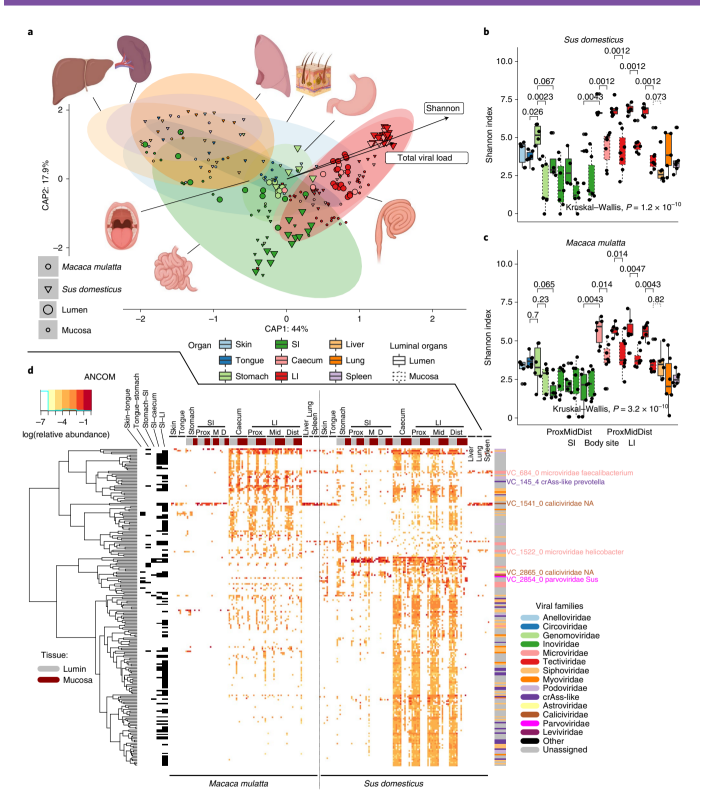
We attempted to identify specific VCs driving the separation between organ-specific viromes (Fig. 2d), as well as VCs responsible



**Fig. 1** Abundance of different viral families along the GIT and in parenchymal organs of domestic pigs and rhesus macaques.  $\mathbf{a}$ ,  $\mathbf{b}$ , Absolute abundance of viral genomes for ( $\mathbf{a}$ ) domestic pigs (n = 6) and ( $\mathbf{b}$ ) rhesus macaques (n = 6) was calculated by comparing coverage with that of the spike-in standard (phage Q33). Only genomes with >50% of estimated completeness were taken into account when calculating viral loads. Bar heights correspond to median values across six animals of each species, error bars denote IQR. Rows are tissue types (top to bottom: lumen, mucosa and skin/parenchyma); columns are anatomical sites. The middle portion of the figure shows schematic locations of the sampled anatomical sites. Dist, distal; Mid, medial; Prox, proximal.

for separation between luminal and mucosal viromes and the two animal species. Across the two species, a total of 217 VCs were differentially abundant between organ pairs in the following sequence: skin-tongue-stomach-SI-caecum-LI (P<0.05 in ANCOM test

with Benjamini–Hochberg correction; see Supplementary results for more detail), with the largest fraction of these VCs (n=119) being discriminatory between the SI and caecum/LI. Twenty VCs were found to be differentially abundant between luminal



**Fig. 2** |  $\alpha$ - and  $\beta$ -diversity of viromes in various anatomical sites in domestic pigs (n = 6) and rhesus macaques (n = 6). **a**, Canonical analysis of principal coordinates of Bray-Curtis dissimilarities between virome samples, based on fractional VC counts, Anatomical locations, Shannon diversity index and total viral load were used as constraining explanatory variables (vectors are only shown for the latter two). Ellipses represent 95% confidence regions. **b**,**c**, Shannon diversity index calculated with read counts for individual viral genomic contigs in pigs (**b**) and macaques (**c**). Organ colours match those in **a**, dashed boxplots represent mucosal sites. Boxplots are standard Tukey type with IQR (box), median (bar) and Q1 - 1.5 × IQR/Q3 + 1.5 × IQR (whiskers). **d**, VCs differentially abundant between organs selected using ANCOM-II test (P < 0.05 after Benjamini-Hochberg correction). Rows represent VCs and columns represent sites in individual animals. A series of post hoc tests identified VCs (black blocks) discriminatory between the following anatomic locations: skin-tongue, tongue-stomach, stomach-SI, SI-caecum and SI-LI. The upper and the right-hand side annotation bars represent tissue types (lumen versus mucosa) and viral families of VCs, respectively. The tree represents hierarchical clustering of VCs based on relative abundance patterns. An expanded version of panel **d** is provided as Supplementary Fig. 5.

and mucosal sites in both species of animals, 11 of them being over-represented in mucosal sites compared with the luminal sites in the same organs. As described in the Supplementary results, many of the organ-discriminatory VCs were positively correlated with bacterial genera characteristic of a particular segment in the GIT (Supplementary Information).

Sharing of virome components between different regions in the GIT. Having observed only this partial separation of GIT sites by virome composition, we reasoned that there should be extensive sharing of individual viral species/strains between multiple GIT sites in each of the animals. To investigate this, we returned to the level of individual viral contigs and visualized their sharing between organs in a particular sequence. Agreeing with the individualized nature of gut viromes demonstrated above, patterns of viral contig sharing between different organs were also unique, not only between pigs and macaques, but also between individual animals within each cohort (Extended Data Figs. 4 and 5). Despite that, common trends in viral sharing between organs could also be easily observed.

As shown in aggregate maps of viral contig sharing, summarizing the data from all pigs (Fig. 3) and all macaques (Fig. 4), high-diversity populations of LI bacteriophages (Fig. 2b,c) are also efficiently shared between all locations in caecum and colon (Figs. 3 and 4). Summing up data from all pigs, for instance, >2,000 crAss-like phage and 65–130 *Microviridae* genomic contigs are shared between sites from the caecum to the distal colon (in luminal and mucosal samples together). Similarly, in macaques, 65–109 crAss-like contigs and 33–98 *Microviridae* were found shared between same anatomical sites. The same pattern was also true for tailed bacteriophage genomic contigs in the families *Siphoviridae*, *Myoviridae* and *Podoviridae* (Extended Data Fig. 6).

As a rule of thumb, >50% of viral contig diversity in pigs and >30% in macagues was shared between all locations in the LI. Extensive sharing of viral contigs was observed within the SI of both animal species. By contrast, only 1%-2% of pig distal SI viral diversity and <1% of the same in macaques was detectable in caecum samples (Figs. 3 and 4). This picture is, however, complicated by the fact that some of the gastric and SI viral contigs that we failed to detect in the distal segment were nevertheless found in caecum and/or lower segments of the LI. This suggests that limitations in sequencing depth and/or strict criteria of contig detection might introduce some artificial gaps of contig detection across multiple anatomical sites in our data. Distal colonic samples (a proxy for faecal samples in our study), appeared to be good representatives of total viral diversity in the lower GIT (>50% represented) and poorer representatives of the upper GIT (~10% represented of gastric virome). Only a small fraction of tongue virome could be detected in the distal colon (Figs. 3 and 4).

Nevertheless, our data contain numerous examples of prokaryotic and eukaryotic viruses (genomic contigs with ≥50% estimated completeness) shared across six or more different anatomical sites. In pigs, such examples include *Astroviridae* and *Caliciviridae* species in luminal samples, *Parvoviridae* in mucosal samples and parenchymal organs, as well as numerous bacteriophage types across anatomical locations. In macaques *Circoviridae* and *Caliciviridae* species were ubiquitously found (Extended Data Fig. 7).

Livers, lungs and spleens of both animal species shared with the GIT sites not only the genomes of eukaryotic viruses (*Circoviridae*, *Caliciviridae*, *Parvoviridae*), but also small genomes of *Microviridae* phages and other phage genomic contigs (Figs. 3 and 4 and Extended Data Figs. 4–6). In the light of some recent publications, this can be interpreted as evidence for possible translocation of some digestive tract bacteriophages across healthy gut epithelia<sup>61</sup>, ending up in the internal organs (liver, lung, spleen), presumably via macropinocytosis, the portal vein (liver), lymphatic system or perhaps via regurgitation of stomach contents (lung).

#### Discussion

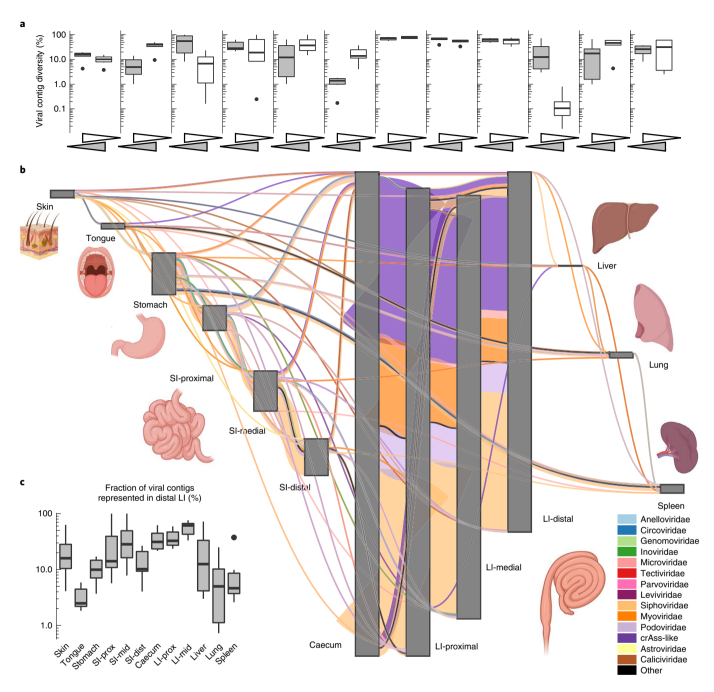
Recent studies have observed correlations in gut bacteriome and phageome composition and claimed associations between altered virome composition and GIT diseases in humans<sup>31,32</sup>. It has been speculated that phages could play a decisive role in controlling bacterial population density and structure via 'kill-the-winner' or similar types of ecological dynamics<sup>62-64</sup>. Indeed, in simplified microbiota models, exponential growth of phage under optimal conditions can lead to the rapid collapse of sensitive bacterial populations<sup>65</sup>, resulting in cascades of knock-on effects in non-susceptible bacterial populations via inter-bacterial interactions<sup>16</sup>.

On the other hand, there is also convincing evidence that points towards a much less disruptive role of phages in microbiome composition, in that most numerically prevalent phage types are either temperate (existing in the form of prophages as well as free viral particles) or have evolved to support a long-term, stable persistence in the microbiome with only limited effects on the density of bacterial host populations<sup>66</sup>. A number of potential persistence mechanisms have been proposed that include phase variation of phage receptors in bacteria<sup>20,28</sup> leading to herd immunity<sup>21</sup>, or physical segregation of mucus-embedded sensitive bacteria from luminal phages ('source-sink' model)<sup>34</sup>. It would be impossible to fully understand the dynamics of phage-host interaction and therefore the role of phages as either 'drivers' or 'passengers' in real-world complex microbiomes without having a detailed map of the virome in both temporal and spatial (biogeographic) dimensions. In this study we provide such a spatial map for two mammalian species, pigs and macaques.

From a technical perspective, the study was designed to minimize the biases typically associated with virome analysis 42,47,67. We used unamplified nucleic acids and assembly-based cataloguing of unclassified viruses, coupled with quantitation by comparison against a spike-in viral standard. We also clustered individual sequences into VCs to allow us to robustly detect and quantify both known and unclassified viruses with DNA or RNA genomes. Unlike in many previous studies<sup>67</sup>, we revealed an abundance of RNA viruses, including unclassified phages belonging to Leviviricetes class, and mammalian viruses belonging to the Astroviridae and Caliciviridae (Supplementary Information). Small ssDNA Microviridae phages were found to be a dominant group in the macaque colon, a finding that previously would have been dismissed as a DNA amplification bias 17,46. A limitation of this assembly-based approach was, however, that we almost certainly missed some of the low-abundance viruses seen in a previous study of the macaque virome<sup>26</sup>.

In the mammalian GIT, a number of factors may influence differences in the bacterial microbiota and virome between SI and LI. Lower pH, higher oxygen tension, faster transit time and bile acid activity may limit bacterial growth in the SI, whereas a thicker mucus gel layer, slower transit time and shift to fermentation contribute to a large increase in microbial density in the LI<sup>68</sup>. As expected, the vast majority of phage biomass and diversity were concentrated in the colonic lumen, reflecting the dense community of bacterial hosts in that site. Upper GIT viromes were distinctly different and reflective of differences in bacteriome composition between different GIT regions (Supplementary Information). Direct correlations in density and composition between virome and bacteriome in the gut have been reported previously<sup>17</sup> and are consistent with the 'piggyback-the-winner' ecological model<sup>30</sup>.

Interestingly, distal gut luminal viromes appeared to be very homogenous, from caecum to distal colon, and compositionally much more reflective of an individual animal, than of a particular location in the colon. This confirms that interindividual variability remains a hallmark of the intestinal virome<sup>17</sup>, even within these highly controlled environments. Recently, stochastic assembly effects have been shown to drive interindividual variability in the



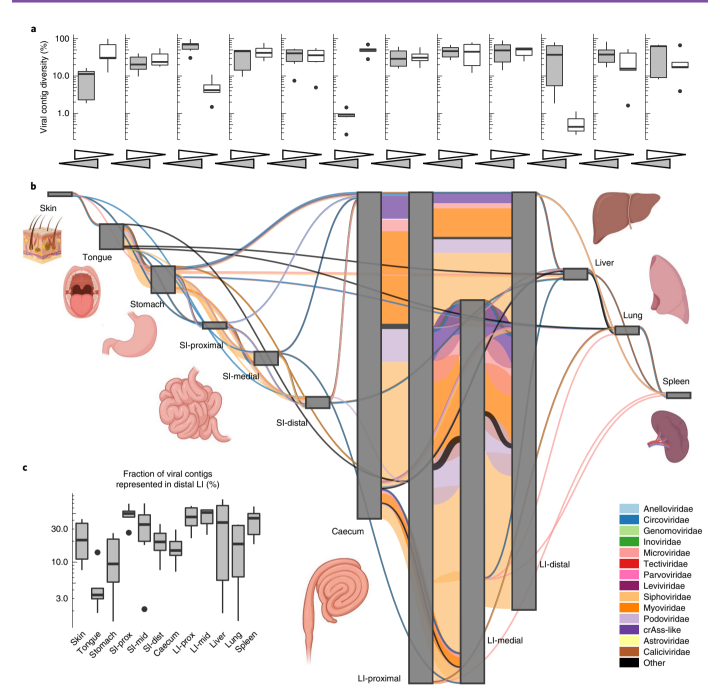
**Fig. 3 | Sharing of viral genomic contigs between different anatomical sites in pigs (n = 6). a**, Fraction of viral contig diversity shared between pairs of sites in both directions (white arrows/boxplots are the forward direction, and grey arrows/boxplots are the reverse), in the order indicated in panel **b**. Boxplots are standard Tukey type with IQR (box), median (bar) and Q1 – 1.5 × IQR/Q3 + 1.5 × IQR (whiskers). **b**, Aggregated map of viral contig sharing across six animals; the height of the vertical grey rectangles is proportional to viral richness (individual genomic contig counts) at each location, aggregated across luminal and mucosal samples. The thickness of coloured connectors is proportional to the number of genomic contigs of each viral family shared between pairs of locations. Unclassified genomic contigs were excluded. **c**, Fraction of viral contig diversity from each organ represented in the distal LI; boxplots are standard Tukey type as above.

bacterial microbiota in mice<sup>69</sup> and this phenomenon should apply similarly to the virome in pigs and macaques.

Our results are in close agreement with research conducted on bacterial biogeography in the macaque gut, in which Yasuda et al. observed predominantly interindividual, and less location-specific, variation of luminal microbiota in jejunal, ileal and colonic sites<sup>70</sup>. The same authors noted differences between luminal and mucosal microbiota in the same locations, with the latter being more influenced by biogeography than by an individual animal. In line with

this, subsets of VCs appear to be specifically associated with both types of habitat (Supplementary Information). At the same time, mucosal samples show drastically reduced viral load and increased prevalence of viruses infecting mammalian cells.

These results may support a recently proposed 'source-sink' model<sup>34</sup> arguing that exclusion of bacteriophages from the mucous layer creates a refuge for bacterial cells, allowing the co-existence of virulent phage and sensitive bacterial cells in close proximity. This apparently disagrees with an earlier 'bacteriophage adherence



**Fig. 4 | Sharing of viral genomic contigs between different anatomical sites in macaques (***n* = **6). a**, Fraction of viral contig diversity shared between pairs of sites in both directions (white arrows/boxplots are the forward direction, grey arrows/boxplots are the reverse), in the order indicated in panel **b**. Boxplots are standard Tukey type with IQR (box), median (bar) and Q1 – 1.5 × IQR/Q3 + 1.5 × IQR (whiskers). **b**, Aggregated map of viral contig sharing across six animals; the height of the vertical grey rectangles is proportional to viral richness (individual genomic contig counts) at each location, aggregated across luminal and mucosal samples. The thickness of coloured connectors is proportional with the number of genomic contigs of each viral family shared between pairs of locations. Unclassified genomic contigs were excluded. **c**, Fraction of viral contig diversity from each organ represented in the distal LI; boxplots are standard Tukey type as above.

to mucus' (BAM) model<sup>37</sup>, which argued that an accumulation of bacteriophages and an increased virus-to-microbe ratio (~39:1) in the mucus creates a barrier limiting bacterial invasion and segregating bacterial population to the luminal space. In the absence of quantitative data on bacteria, our study cannot testify to the virus-to-microbe ratios in the lumen and mucosa. The BAM model, therefore, can still accommodate our results, with a caveat that certain bacteriophages possessing immunoglobulin-like

protein domains required for binding to mucus<sup>37</sup> are equally abundant in the mucus and in the lumen, whereas phages lacking this ability are excluded into the luminal space. One can envisage complex scenarios of phage–host interaction in the GIT, with some phage–host pairs following 'source–sink' dynamics, whereas others show behaviours more conforming with the BAM model.

We observed extensive sharing of individual viral strains throughout the entire GIT. The most prominent examples were

phages found continuously across multiple sites. For the majority of strains, however, the continuous flow of phages from SI to LI seems to be interrupted at the ileocaecal valve. This can be explained in part by drastic differences in the composition (and presumably total biomass) of bacteriomes between the SI and LI, which in turn support the growth of completely different phage populations. However, a complete extinction of SI phages during passage from SI to LI seems unlikely, and therefore, the dilution effect, caused by a vastly larger viral biomass supported by greater numbers of bacteria, combined with limitations imposed by sequencing depth, is a probable cause of the apparent disappearance of gastric and SI phages in the caeca and LI.

Despite our original expectations, we could not definitively confirm a tendency for mucosal samples taken at 1 cm distance to be closer in virome composition to each other than to other sites in the same anatomical region (Supplementary Information), which again suggests relative homogeneity of the virome along the proximal-distal axis within each region of the digestive tract. This observation calls for future longitudinal studies to examine viral flow and local temporal differences in virome composition in the gut.

Luminal samples from the distal colon, which can roughly be equated with faecal samples for the purpose of this study, are only representative of a fraction of the viral diversity present in different segments of the digestive tract. This is especially evident in the case of eukaryotic viruses, many of which are readily detectable in colonic mucosa (Astroviridae in pigs) or SI lumen (Caliciviridae in both pigs and macaques), and in parenchymal organs such as liver, lung and spleen, but not in the distal LI lumen. Interestingly, and in agreement with our earlier notion of virome individuality, each animal harboured a unique pattern of eukaryotic viruses, with regards to their taxonomic composition, strain variation and biogeographic distribution (Supplementary Information). The epidemiological and pathological importance of the biogeographic distribution of these common viruses in porcine and murine GIT (in particular porcine Astroviridae<sup>71</sup>) is difficult to establish without collection of further extensive population and longitudinal data.

Our findings in this study were largely consistent between pigs and macaques, despite differences in species, environment, diet and age. Notably, the pigs at 3 months old were weaned and in early adolescence, whereas macaques were adults (5–12 years old). We note that all animals were female, thus preventing any determination of possible sex-effects on intestinal biogeography.

One of the interesting findings in this study was possible evidence of bacteriophage translocation from the gut to the systemic circulation and eventually parenchymal organs such as the liver, spleen and lungs. Although animal dissection and sample collection for this study were conducted within a sanitary research environment, we could not achieve fully aseptic conditions in our pig facility. Therefore, it is possible some of the viral biomass in pig parenchymal organs that was orders of magnitude lower than was found in the gut could represent cross-contamination of solid organ samples. Nevertheless, we believe that this cannot fully explain our findings. Parenchymal organ viromes were dominated by eukaryotic viruses, and although phages present in these organs were specific strains shared with digestive tract viromes, they were not the most dominant strains. It has previously been demonstrated that at least specific phage types are able to adhere and translocate through the intestinal epithelial lining<sup>37,61</sup>. In our study, a tendency towards enrichment for smaller phages (family Microviridae) was observed in parenchymal organ viromes, which might indicate increased transepithelial diffusion of small viral particles. The exact fate of translocating phage and their systemic effects has so far remained unclear<sup>72,73</sup>, and our observations might be insightful for studying anti-phage immune responses<sup>74</sup>.

This work highlights that focusing on distal LI sampling (or faecal sampling) dramatically under-represents gastrointestinal (GI) viral communities (particularly eukaryotic viruses), and points to a consistent drop-out of upper GI viral communities in colonic samples. In addition to these findings, we detected some overlap between viral communities in parenchymal organs and the GIT that was not related to their overall abundance, suggesting that there may be some degree of specificity to viral translocation. Finally, we propose that this dataset and its accompanying methodology may provide an important catalogue of gut viruses and resource for future investigators in the field.

#### Methods

Ethical approval and study design. The study design was developed with consideration to the three Rs for the ethical use of animals in science: replacement, reduction and refinement. The proposed euthanasia-only study was reviewed by the Animal Welfare Body of University College Cork (Euthanasia Only Authorization 17-005). With authorization and under the remit of authorized and experienced personnel, the study was performed succinctly and with minimal distress to the animals involved. No statistical methods were used to predetermine sample sizes but our sample sizes are similar to those reported in previous publications<sup>26,70</sup>. Data collection and analysis were not performed blind to the conditions of the experiments. No randomization procedures were used and no data points were excluded from any of the analyses.

Animal sampling procedures. Six healthy female Landrace pigs (Sus scrofa domesticus, body mass ~30 kg, approximately 3 months of age) were sourced from a local farm in Cork, Ireland. All pigs were raised in a shared environment and on the same diet, although the relatedness of their parentage is unknown. Pigs were transported to the research facility on the morning they were to be killed, with two animals sampled back-to-back per day. Before euthanasia, work surfaces and necessary tools were disinfected using Virkon S disinfectant. Following anaesthetic overdose with pentobarbital (150 mg kg<sup>-1</sup>) death was confirmed by an authorized person, and tissue samples were collected. All biopsies (minimum 3 cm × 3 cm) were minimally handled on site. Therefore, samples were not washed or stored in a buffer but placed directly into 50 ml Falcon tubes and stored on dry ice and then at -80 °C. Initially, external biopsies of the tongue and skin were collected. Skin biopsies were taken from around the shoulder. Once external biopsies were obtained, pigs were rolled onto their backs and a midline incision was performed from below the neckline of the animals to immediately preceding the genitalia. The complete GIT was removed from the abdominal cavity, with the connective tissue severed where required. Surgical thread was used to seal sections of the GIT. Two knots, ~2 cm apart, were tied tightly without severing the GIT. Subsequently, sections of the GIT were separated by cutting between the tied knots, which prevented the intestinal contents from leaking. Both the SI and LI of animals were sealed in three approximately equal length sections to represent the proximal, medial and distal regions. All sections of the GIT were treated similarly. Briefly, an opening into the sealed GIT tube was created and the contents removed before large representative sections of the bowel were cut and stored. Finally, stomach mucosa was from fundic region, and parenchymal organs were removed from the abdominal cavity of animals with large biopsies sections stored for later analysis. The processing time per animal was ~3 h.

Six healthy Indian-origin, female adult rhesus macaques (*Macaca mulatta*) aged 5–12 years and weighing 5.3 to 10.6 kg were used. All animals were born and raised in naturalistic multigenerational breeding groups at the Biomedical Primate Research Centre (BPRC), Rijswijk, The Netherlands, in comparable environments. All enclosures contained environmental enrichment and bedding to stimulate their natural behaviour. They were daily fed monkey chow pellets (Ssniff) in the morning, complemented with fruit and vegetables. Over a period of 5 months animals were killed using pentobarbital (70 mg kg<sup>-1</sup>) following sedation with ketamine (10 mg kg<sup>-1</sup>). The necropsy and collection of samples were done immediately after euthanasia.

For isolation and collection of macaque samples, strict sterility protocol and safety procedures were used. The sterility of the necropsy table and the surgical instruments was assured using Virkon S, sterilization procedures and the use of disposable scalpels. Macaque tissue samples were retrieved and stored similarly to the procedures outlined for pigs. For the collection of parenchymal and intestinal samples, disposable scalpels and autoclaved scissors and forceps were used. To avoid contamination, after opening the thoracic and abdominal cavity, the first samples collected were from the parenchymal organs (liver, spleen and lung) following by the intestinal samples. After each animal, the table was thoroughly cleaned with hot water and detergent followed by disinfection with Virkon S, to prepare for the next animal. All samples were placed immediately on dry ice and stored at  $-80\,^{\circ}\text{C}$ . Tissue samples were transported on dry ice to APC Microbiome Ireland for further processing.

**Biopsy preparation procedure, VLP enrichment and nucleic acid sequencing.** GI and parenchymal organ sections of pigs and macaques were processed identically, in the same research facility, by the same team members, but on

different days. Tissue samples were thawed on ice until completely defrosted. Excess faecal material on caecal and colon tissue sections was washed off with sterile SM buffer (50 mM Tris–HCl, 100 mM NaCl, 8.5 mM MgSO $_4$ , pH 7.5). Tissue sections were stretched and pinned to a Styrofoam board using sterile syringes. Defined volume pinch biopsies of mucosal surfaces were collected with endoscopic biopsy forceps. A 'double-bite' of tissue samples at the same site ensured accurate and complete loading of the forceps' jaws. Mucosal pinches were removed from the forceps directly into pre-labelled Eppendorf tubes, filled with 400  $\mu$ l of sterile SM buffer for processing.

To enable comparisons of viral load across biopsy samples, 10 µl of 107 plaque-forming units per millilitre of lactococcal phage Q33 were added to all samples. Additionally, Q33 in SM buffer or SM buffer alone were processed as negative controls. Fresh 0.5 M dithiothreitol was prepared in 1 ml of SM buffer. A volume of 16 µl of the dithiothreitol stock was added to samples to achieve a final concentration of 20 mM, and samples were incubated at 37 °C for 30 min. Dithiothreitol was used to gently solubilize mucin with minimal disruption of phage virions because this disulphide bond reducing agent was previously demonstrated to release large quantities of non-mucin proteins from SI porcine preparations75. Host cellular debris and bacterial cells were pelleted by gentle centrifugation at 4,000g for 30 min at room temperature. Subsequently, 400 µl of liquid was aspirated and treated with 40 µl of DNase/RNase buffer (50 mM CaCl<sub>2</sub>, 10 mM MgCl<sub>2</sub>), 12 µl (24 U) of Turbo DNase (Thermo Fisher) and 4 µl (40 U) of Fermentas RNase I (Fisher Scientific), and incubated at 37 °C for 1 h with intermittent inversion approximately every 15 min. Enzymes were inactivated by incubating at 65 °C for 10 min.

Viral-enriched samples void of free nucleic acids were lysed using the QIAgen Blood and Tissue Kit following the manufacturer's guidelines. However, samples were eluted in only  $20\,\mu$ l of Qiagen AE elution buffer (10 mM Tris-HCl, 0.5 mM EDTA, pH 9.0) to increase the final concentration of nucleic acid obtained.

Virome shotgun library preparation and sequencing. A reverse transcription reaction was performed using SuperScript IV First Strand Synthesis System (Invitrogen/Thermo Fisher Scientific) with 11  $\mu$ l of purified VLP nucleic acids sample and random hexamer oligonucleotides according to the manufacturer's protocol. The concentration of DNA purified using a DNeasy Blood & Tissue kit (Qiagen) was determined using the Qubit dsDNA HS kit and the Qubit 3 fluorometer (Invitrogen/Thermo Fisher Scientific). DNA/cDNA yields varied between 0.05 and 29 ng  $\mu$ l<sup>-1</sup>, with some samples being below the detection limit.

Library preparation was carried out using Accel-NGS 1S Plus kit (Swift Biosciences) according to the manufacturer's instructions. Briefly, 20 µl of reverse transcription product (regardless of DNA concentration because the kit is flexible with regards to the amount of input DNA) were taken for sonication after adjusting the volume to 52.5 µl with low-EDTA TE buffer. Shearing of unamplified DNA/cDNA mixture (variable amounts of DNA) was performed on M220 Focused-Ultrasonicator (Covaris) with the following settings: peak power of 50 W, duty factor of 20%, 200 cycles per burst, total duration of 35 s. All subsequent steps were performed in accordance with the manufacturer's protocol. A 0.8 DNA/AMPure beads v/v ratio was used across all purification steps in the Accel-NGS 1S Plus protocol. Post-preparation library QC (fragment length distribution and quantitation) was performed using Agilent Bioanalyzer 2100 with High Sensitivity DNA kit and Invitrogen Qubit. Dual-indexed pooled library was sequenced using 2×150 nucleotide paired-end sequencing run on an Illumina NovaSeq platform at GENEWIZ (Leipzig, Germany).

To control for contamination of samples with exogenous viruses and viral nucleic acids, including laboratory reagent-derived and environmental, we also performed extraction from 400  $\mu$ l of sterile SM buffer alone. Two samples were processed simultaneously with pig and macaque samples using the same protocol. Both yielded DNA/cDNA below the detection limit after extraction, and only a trace (insufficient for sequencing) amount of DNA was visible. To compensate for the low yield, a third sample was subjected to whole-genome multiple displacement DNA amplification (Illustra GenomiPhi V2 DNA Amplification Kit) as described previously  $^{17}$ .

Analysis of virome shotgun sequencing data. Raw reads were processed using Cutadapt v.2.4 to remove adaptor sequences. Trimmomatic v.0.36 (ref. \*\*) was used for quality-based trimming and filtration of reads with the following parameters: 'SLIDINGWINDOW:4:20 MINLEN:60 HEADCROP:10'. Reads aligning to mammalian genomes were identified using Kraken v.1.1.1 in combination with *Macaca mulatta* (GCF\_000772875.2\_Mmul\_8.0.1) and *Sus scrofa* (GCF\_00003025.6\_Sscrofa11.1) reference genome files.

Following removal of mammalian reads, levels of contamination with bacterial genomic reads were assessed using ViromeQC tool?". Reads were then assembled into contigs on a per sample basis using SPAdes assembler v.3.13.0 in metagenomic mode with standard parameters?8. Additionally, in an attempt to assemble low-abundance genomes, reads were pooled by animal and assembled using MEGAHIT v.1.2.1-beta?9. All contigs >1 kb were then pooled and an all-vs-all BLASTn search was performed with an e-value cut-off of  $\leq 1\times 10^{-20}$ . Contig redundancy was removed by identifying pairs sharing 90% identity over 90% of the length (of the shorter contig in each pair) retaining the longest contig in each case.

To extract viral contigs from a background of bacterial contamination several selection criteria were used. First, contigs aligning using BLASTn v.2.10.0+ $^{80}$  against viral sequences in the National Center for Biotechnology Information (NCBI) RefSeq database (release 208), Gut Virome Database $^{51}$ , JGI IMG/VR database (v.3, release 12-10-2020) $^{50}$ , Gut Phageome Database $^{54}$  and the recent human gut phage MGV database  $^{55}$ , as well as our in-house database of crAss-like phage genomes ( $n\!=\!1,\!576$ ), with at least 50% identity over 85% of contig length (e-value cut-off of individual hits  $\leq\!1\!\times\!10^{-10}$ ) were deemed as viral. Second, contigs that identified as viral using VirSorter2 pipeline  $^{56}$  with strict criteria (score  $\geq\!0.9$  or score  $\geq\!0.7$  with at least one viral hallmark protein-coding gene present) were added. Completeness level of viral genomic contigs was determined using CheckV $^{81}$  with default parameters. VirSorter2-identified viral contigs marked as prophages by CheckV (Provirus = Yes), were eliminated. Viral genomic contigs identified by these approaches constituted the final non-redundant viral sequence catalogue ( $n\!=\!107,680$ ).

Protein-coding genes on viral contigs were predicted using Prodigal  $^{\mathbb{N}^2}$  (-meta mode). Translated protein sequences were searched against PHROGs database of virus-specific protein family profile HMMs, using hmmscan (HMMER v.3.1b2; e-value cut-off of  $\leq 1\times 10^{-5}$ ); viral protein sequences from NCBI non-redundant protein sequence database (as of 2 November 2021), viral RefSeq (release 208) database and crAss-like phage proteins from an in-house database (n = 7,356) using BLASTp v.2.10.0+ (e-value cut-off of  $\leq 1\times 10^{-10}$ ). Circular genomic contigs were identified using LASTZ. G+C content was calculated using EMBOSS geecee.

Assignment of contigs to viral families was accomplished using Demovir script (https://github.com/feargalr/Demovir), as described previously<sup>17</sup>. Clustering of viral genomic contigs (only for contigs with >3 kb in length) into VCs (approximately genus-level operational taxonomic groups) was done using vConTACT2 software<sup>60</sup> with the following optional parameters: --rel-mode Diamond --db ProkaryoticViralRefSeq85-Merged --pcs-mode MCL --vcs-mode ClusterONE. The viral genomic contig catalogue was further manually curated to remove coliphage phiX174 genome (commonly used as a spike-in by sequencing facilities). Phage lifestyle (temperate versus virulent) was predicted using BACPHLIP<sup>84</sup> using a 0.95 confidence threshold.

Remaining (non-viral) non-redundant contigs were assigned to bacterial taxa by performing a BLASTn search against bacterial RefSeq (release 99) and HMP Reference Genomes databases. Taxonomic assignments were made at the genus level for contigs having 90% identity over  $\geq$ 85% of combined alignment(s) length against a reference bacterial genome. CRISPR arrays were predicted on bacterial contigs and spacer sequences were extracted using PILER-CR v.1.06 (ref.  $^{85}$ ).

To predict the hosts of phage, data were aggregated from several sources. First, previously predicted hosts for viral species included in the IMG/VR database were assigned to viral contigs in our catalogue belonging to the same species (≥95% identity over ≥85% of viral genomic contig length, in accordance with MIUViG criteria for viral species demarcation in metagenomic sequence data<sup>57</sup>). Second, a search against an in-house CRISPR spacer database (derived from bacterial RefSeq (release 89) and HMP Reference Genomes) was performed as described previously<sup>17</sup> to assign hosts to viral contigs, missing close homologues in the IMG/ VR database. In a similar fashion, matches were found with CRISPR spacers encoded by bacterial contigs (with taxonomy assigned as described above) in the current study's dataset. Third, BLASTn similarity of viral contigs to closely related (≥90% identity over ≥85% of viral contig length) prophages in bacterial genomes (RefSeq database of bacterial genomes, release 99; HMP Reference genomes database86) was used to assign hosts where neither IMG/VR nor CRISPR approaches were successful. Lastly, transfer RNA gene hits against the NCBI nucleotide database (release 28 November 2020) and bacterial RefSeq database (release 99) were used to predict hosts for cases in which all other methods failed. At the VC level, host was assigned using the majority vote rule, after aggregating host predictions from individual viral contigs-members of a particular VC.

Quantitative analysis of viral metagenomic data was performed essentially as described previously<sup>17</sup>. Quality filtered reads were aligned to the curated viral contig database on a per sample basis using Bowtie2 v.2.3.4.1 in the 'end-to-end' mode. A count table of contigs versus samples was subsequently generated using SAMTools v1.7. Sequence coverage was calculated per nucleotide position per contig per sample using SAMTools 'mpileup' command. Read counts for contigs in samples showing less than a minimum of 1× coverage of 75% of a contig length, were set to zero<sup>17</sup>.

Absolute viral counts were calculated for viral genomic contigs based on comparison of their relative abundance with that of the externally added standard (lactococcal phage Q33). Only viral contigs with estimated completeness of >50% were taken into account based on an assumption that additional genomic fragments, which together constitute the remaining <50% portion of the complete genome, will not be counted and therefore will not artificially inflate the calculated total viral loads.

**Bacterial 16S rRNA amplicon sequencing.** During the biopsy preparation procedure, porcine and macaque biopsy samples were reduced with dithiothreitol followed by centrifugation to reduce host tissue and bacterial cells and enrich the viral-like particles. However, the bacterial-containing pellet was used as the starting material for complementary 16S rRNA analysis of bacterial communities

associated with the same biopsy samples analysed with respect viromes. The preparation and sequencing of 16S rRNA gene V3–V4 segment libraries followed the procedure outlined previously<sup>43</sup>.

Analysis of bacterial 16S rRNA amplicon sequencing data. Bacterial 16S rRNA amplicon sequencing data were processed using a pipeline based on USEARCH v.8.1 (64 bit). Forward and reverse reads of the 16S rRNA V3–V4 segment were merged allowing for an expected error rate of <0.5 per nucleotide position at overlap. Merged sequences were truncated to remove forward (first 17 nucleotides) and reverse (last 21 nucleotides) 16S rRNA primers. Reads were then de-replicated and singletons were removed, followed by clustering into operational taxonomic units (OTUs) at a 97% sequence identity level. Chimeras were removed using the *-uchime\_ref* function with the  $rdp\_gold$  reference database. Individual reads were then assigned to OTUs generated above at 97% sequence identity cut-off and a read count matrix was generated. Finally, taxonomic assignment of OTUs was performed using RDP Classifier v.2.12.

Statistical methods. All statistical analysis of sequencing data was carried out in R environment v.4.1.0. Descriptive statistical visualizations were created using ggplot2 v.3.3.3. Network visualizations were created using igraph v1.2.6. Heat maps were produced using gplots v.3.1.1. Sankey diagrams were made using networkD3 v.0.4. Permutational multivariate analysis of variance was performed using the adonis() function in Vegan with Bray-Curtis distances. Virome β-diversity was visualized through canonical analysis of principal coordinates with Bray-Curtis distances (capscale() function in Vegan v.2.5-7 with default parameters). Comparison of Bray-Curtis distances between viromes within organs was done using Wilcoxon test with Benjamini-Hochberg corrections. VCs differentially abundant between organs, tissues and animal species were identified using ANCOM-II<sup>87,88</sup> with Benjamini-Hochberg correction,  $\alpha = 0.05$  and  $w_0$  threshold set at 0.7. For between-organ tests, individual animal was used as a random effect variable and models were adjusted for tissue type (lumen versus mucosa) as a covariate. This was followed by post hoc ANCOM-II tests for specific pairs of organs. For between-tissue tests (lumen versus mucosa) and between-species tests (macaques versus pigs), models were adjusted for individual animal or organ type as a covariate, respectively. Correlations between fractional abundances of individual viral genomic contigs (or VCs) and bacterial 16S rRNA OTUs (or genera) were calculated using the Spearman rank correlation method with Bonferroni correction for multiple tests.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

#### Data availability

All data needed to evaluate the conclusions in the paper are present in the paper, Supplementary Information file and the additional dataset available at https://doi.org/10.6084/m9.figshare.15149247.v2. Raw sequencing data are available from NCBI databases under BioProject PRJNA753514. Source data are provided with this paper.

#### **Code availability**

Source data and custom R code used in this study are available at https://doi.org/ 10.6084/m9.figshare.15149247.v2. Further information and requests for data, code and resources should be directed to and will be fulfilled by A. Shkoporov and C. Hill.

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#### **Author contributions**

A.N.S., S.R.S., R.P.R. and C. Hill conceived the study. A.N.S., S.R.S., A.L., I.K., L.A.D., C. Heuston and J.A.M.L. designed the study. A.N.S., S.R.S., A.L., I.K., C. Heuston, A.U., E.V.K., I.v.d.K. and B.O. undertook data acquisition. A.N.S., S.R.S., A.L., I.K., C. Heuston, J.A.M.L., L.A.D., R.P.R. and C. Hill analysed and interpreted the results. A.N.S. was responsible for the software. All authors contributed to drafting and revising of the manuscript.

#### Competing interests

The authors declare no competing interests.

#### Additional information

Extended data is available for this paper at https://doi.org/10.1038/s41564-022-01178-w.

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s41564-022-01178-w.

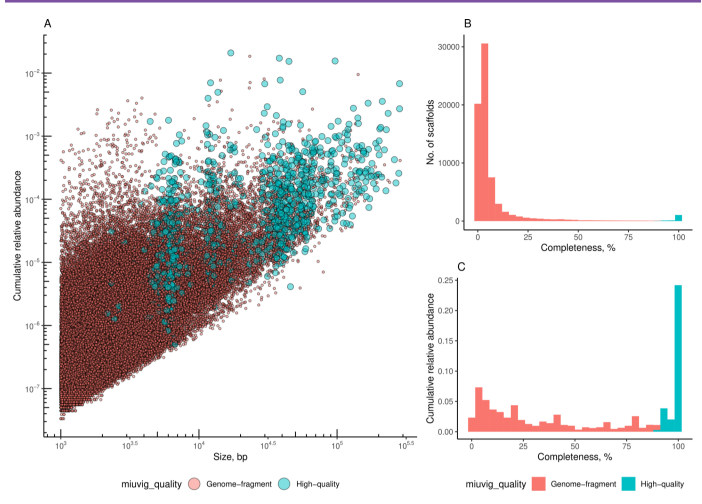
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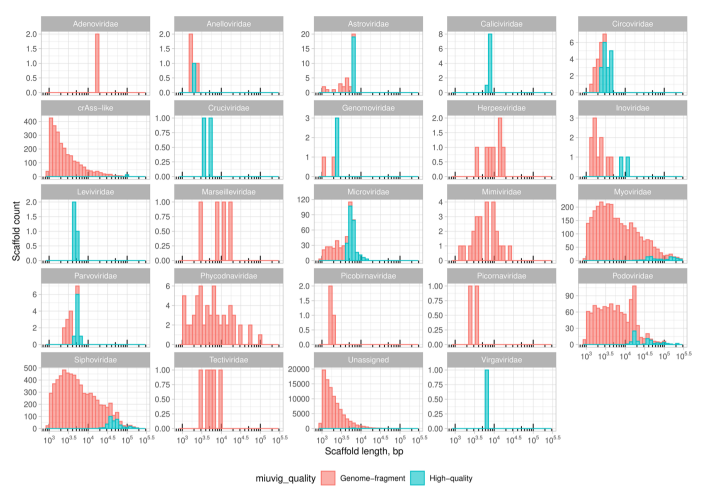
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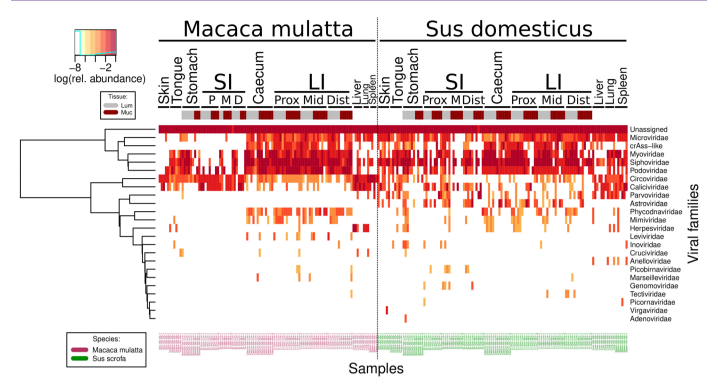
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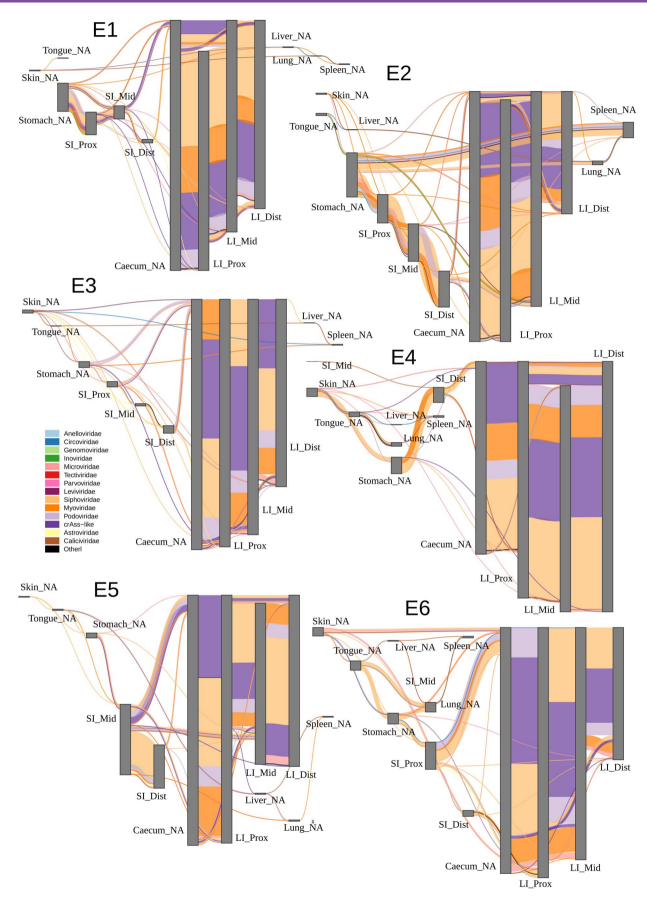
**Extended Data Fig. 1** | Catalogue of viral genomic contigs assembled from trimmed and filtered Illumina reads (n = 107,680). a, Average read coverage vs. contig length, categories of viral genomic contigs identified by CheckV (high-quality genomes vs genome fragments according to definitions given by the MIUViG standard); **b**, distribution of viral genomic contigs by completeness level as predicted by CheckV with high quality draft and complete genomes by MIUViG standard highlighted in blue; **c**, cumulative fractional abundance of genomic contigs with different levels of completeness.



**Extended Data Fig. 2 | Taxonomic distribution, size, and completeness of viral genomic contigs.** Different viral families are shown in separate panels. Assignments are based on Demovir script. Contig size is plotted on log10-scaled x-axis. Contig completeness is predicted using CheckV.

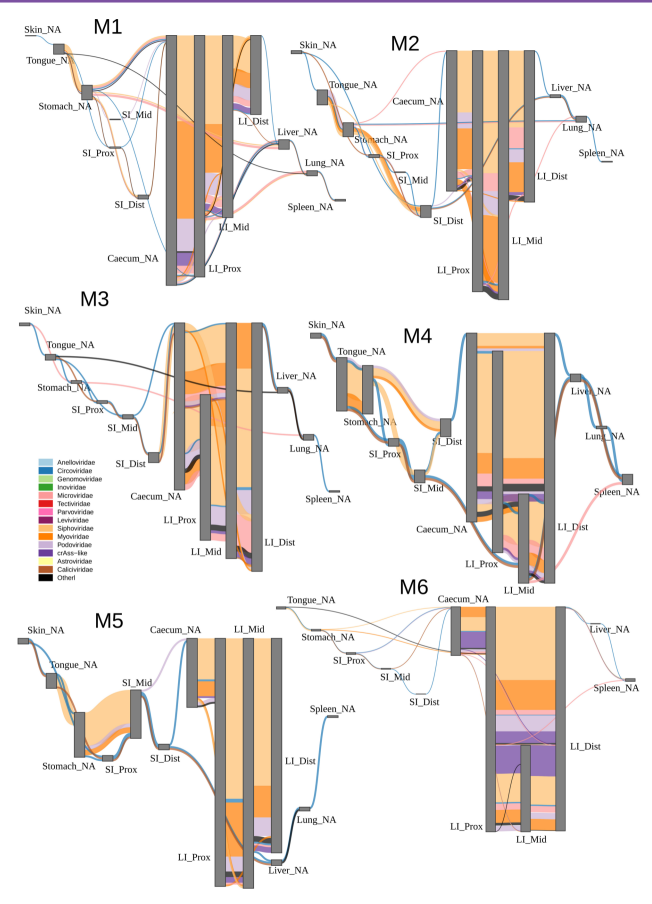


**Extended Data Fig. 3** | Aggregated fractional abundance of viral families across all anatomical sites in pigs (n = 6) and macaques (n = 6) in the **study.** Rows represent viral families, columns – sites in individual animals; the top annotation bar represent tissue types (lumen vs mucosa). Data is log10-transformed and presented with hierarchical clustering based on relative abundance patterns.



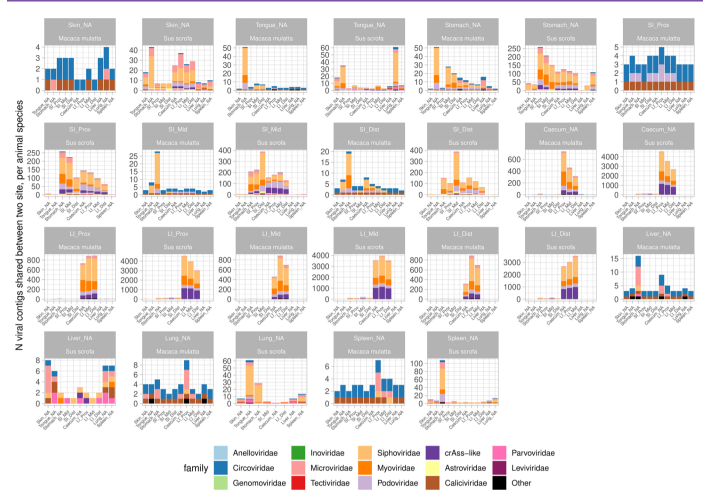
Extended Data Fig. 4 | See next page for caption.

**Extended Data Fig. 4 | Sharing of viral genomic contigs between different anatomical sites in individual pigs (n = 6).** Vertical grey rectangles height is proportional to viral richness (individual genomic contig counts) at each location, aggregated across luminal and mucosal samples; thickness of coloured connectors is proportional with the number of genomic contigs of each viral family shared between pairs of locations; SI, small intestine; LI, large intestine; Prox/Mid/Dist, proximal, medial and distal portions, respectively; unclassified genomic contigs were excluded; C, fraction of viral contig diversity from each organ represented in the distal LI.

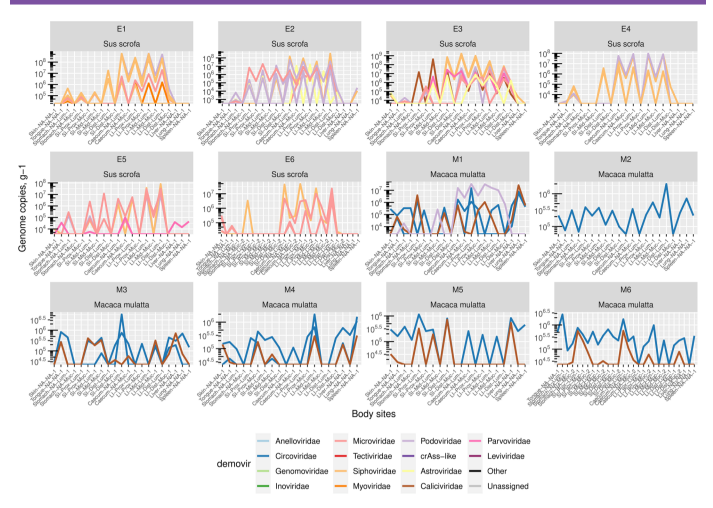


Extended Data Fig. 5 | See next page for caption.

**Extended Data Fig. 5 | Sharing of viral genomic contigs between different anatomical sites in individual macaques (n = 6).** Vertical grey rectangles height is proportional to viral richness (individual genomic contig counts) at each location, aggregated across luminal and mucosal samples; thickness of coloured connectors is proportional with the number of genomic contigs of each viral family shared between pairs of locations; SI, small intestine; LI, large intestine; Prox/Mid/Dist, proximal, medial and distal portions, respectively; unclassified genomic contigs were excluded; C, fraction of viral contig diversity from each organ represented in the distal LI.



**Extended Data Fig. 6 | Numbers of viral genomic contigs shared between pairs of organs in pigs and macaques.** Numbers of shared contigs are expressed as aggregate counts of unique contigs shared between sites across all animals for each of the two species; SI, small intestine; LI, large intestine; Prox/Mid/Dist, proximal, medial and distal portions, respectively.



**Extended Data Fig. 7 | Absolute counts of some of the most ubiquitous viral genomic contigs present in pigs and macaques.** Only contigs with >50% estimated completeness and shared between 6 or more sites in any of the animals are displayed. Each line corresponds to an individual genomic contig (potentially collapsing multiple viral strains). Colours are according to viral families. Each panel represent an individual animal.

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#### Software and code

Policy information about availability of computer code

Data collection

Nucleotide sequene data was generated by an external sequencing provider (GENEWIZ) using Illumina NovaSeq 6000 platform.

Data analysis

The following open source programs were used in this study: Cutadapt v2.4, Trimmomatic v0.36, Kraken v1.1.1, ViromeQC pipeline, SPAdes v3.13.0, MEGAHIT v1.2.1-beta, BLASTn v2.10.0+, VirSorter2 pipeline, CheckV pipeline, HMMER v3.1b2, Demovir script for bash, vConTACT2 pipeline, BACPHLIP pipeline, PILER-CR v1.06, Bowtie2 v2.3.4.1, SAMTools v1.7, USEARCH v8.1, RDP Classifier v2.12, R environment v4.1.0, ggplot2 v3.3.3, gplots v3.1.1, igraph v1.2.6, networkD3 v0.4, Vegan v2.5-7, ANCOM-II script for R, custom R scripts available from https://doi.org/10.6084/m9.figshare.15149247.v1

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All data and code needed to evaluate the conclusions in the paper are present in the paper, supplementary materials, or the additional dataset available at https://doi.org/10.6084/m9.figshare.15149247.v1. Raw sequencing data are available from NCBI databases under BioProject PRJNA753514.

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All studies must dis	close on these points even when	the disclosure is negative.			
Sample size	No statistical methods were used to (Zhao et al., 2019; Yasuda et al., 201	p pre-determine sample sizes but our sample sizes are similar to those reported in previous publications [15]			
Data exclusions No data were excluded from the analysis and reporting.		alysis and reporting.			
Replication	Replication This study was performed with successful replication in six individual animals of each of the two species (pigs and macaques).				
Randomization Not applicable, as this is an exploratory (observational)					
Blinding Data collection and analysis were not performed blind to the conditions of the experiments.					
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Six healthy female Landrace pigs (body mass approximately 30 kg, approximately 3 months of age) were sourced from a local farm in Laboratory animals

Cork, Ireland. Six healthy Indian-origin, female adult rhesus macaques aged 5-12 years with bodyweight 5.3 to 10.6 kg were used. All animals were born and raised in naturalistic multi-generational breeding groups at the Biomedical Primate Research Centre (BPRC),

Rijswijk, The Netherlands, in comparable environments.

Wild animals No wild animals were used in this study.

Ethics oversight

Field-collected samples No field collected samples were used in this study.

The proposed euthanasia only study was reviewed by the Animal Welfare Body (AWB) of University College Cork (Euthanasia Only Authorisation 17-005). With authorisation and under the remit of authorised and experienced personnel, the study was performed succinctly and with minimal distress to the animals involved.

Note that full information on the approval of the study protocol must also be provided in the manuscript.