Hologenomic adaptations underlying the evolution of sanguivory in the common vampire bat

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Adaptation to specialized diets often requires modifications at both genomic and microbiome levels. We applied a hologenomic approach to the common vampire bat (Desmodus rotundus), one of the only three obligate blood-feeding (sanguivorous) mammals, to study the evolution of its complex dietary adaptation. Specifically, we assembled its high-quality reference genome (scaffold N50 = 26.9 Mb, contig N50 = 36.6 kb) and gut metagenome, and compared them against those of insectivorous, frugivorous and carnivorous bats. Our analyses showed a particular common vampire bat genomic landscape regarding integrated viral elements, a dietary and phylogenetic influence on gut microbiome taxonomic and functional profiles, and that both genetic elements harbour key traits related to the nutritional (for example, vitamin and lipid shortage) and non-nutritional (for example, nitrogen waste and osmotic homeostasis) challenges of sanguivory. These findings highlight the value of a holistic study of both the host and its microbiota when attempting to decipher adaptations underlying radical dietary lifestyles.
inflammatory bowel disease\textsuperscript{15–17}. In response to the growing awareness of the key roles that host–microbiome relationships can play across the spectra of life, various studies have advocated for the ‘hologenome’ concept\textsuperscript{18–20}. In brief, this argues that natural selection acts on both the host and its microbiome (together forming the holobiont); thus, evolutionary studies should incorporate both. The extreme dietary adaptation of vampire bats provides a suitable model to investigate the effects of selection across the genome and microbiome, and thus allows exploration of the role of host-associated microbiome in the evolution of specialized diets.

**Results and discussion**

Here, we explore the contributions of both the common vampire bat’s nuclear genome and gut microbiome to its adaptation to obligate sanguivory. To this end, we generated both the common vampire bat genome and fecal metagenomic data sets as a proxy to study its gut microbiome, as well as faecal metagenomic data sets from other non-sanguivorous bat species. We used these data sets for comparative genomic and metagenomic analyses. Specifically, we analysed the common vampire bat genomic landscape, the ratio of substitution rates at non-synonymous and synonymous sites (dN/dS), putative gene loss and gene family expansion/contraction, and computational predictions of the functional impact of amino acid substitutions. We also performed microbial taxonomic and functional profiling, identified the microbial taxonomic and functional core of the common vampire bat, and identified enriched microbial taxa and functions. Following a hologenomic approach, we identified elements in both the host genome and microbiome that could have played relevant roles in adaptation to sanguivory.

**Genomic landscape.** We sequenced and de novo assembled the ~2 gigabase (Gb) common vampire bat genome using Illumina sequencing technology (Supplementary Information 1). The genome is smaller than that of other mammals, but similar to previously reported bat genomes\textsuperscript{11}. The initial assembly (~100× mean coverage, scaffold N50 = 5.5 Mb and N90 = 933 kb; Supplementary Figs. 1 and 2) was subsequently improved using the in vitro proximity ligation-based technology for assembly contiguity refinement developed by Dovetail Genomics\textsuperscript{22}. We obtained a final high-quality assembly with scaffold N50 = 26.9 Mb and N90 = 9.46 Mb, contig N50 = 36.6 kb and N90 = 8.8 kb (Supplementary Table 1, Supplementary Fig. 3 and Supplementary Information 1). We used our annotated common vampire bat genome (see Methods—Protein-coding gene and functional annotation) for comparative genomic analyses with publicly available bat genomes and other mammalian genomes (Supplementary Table 2 and Fig. 1).

Repetitive elements can significantly contribute to genome evolution. Thus, for a genomic landscape characterization, we first compared transposable elements in the common vampire bat genome to those within the genomes of non-phyllostomid bats with other diets: the carnivorous *Megaderma lyra* (greater false vampire bat, Megadermatidae), the insectivorous *Pteronotus parnellii* (Parnell’s moustached bat, Mormoopidae) and the frugivorous *Pteropus vampyrus* (large flying fox, Pteropodidae) (Fig. 1, Supplementary Table 3 and Supplementary Note 1). We identified a 1.6- to 2.26-fold higher copy number of the MULE-MuDR transposon in the common vampire bat genome relative to the other bat genomes. The high mutagenic capacity of MULE-MuDR has been demonstrated to have played a critical role in the evolution of some plants\textsuperscript{23}. Furthermore, transposable elements in general may cause structural or functional changes within the genome and alter epigenetic regulation of the genes into which they are inserted\textsuperscript{24}. Therefore, we explored whether these elements might have also played a role in the evolution of sanguivory by analysing their genomic location in the nuclear genome of the common vampire bat. We found that the identified common vampire bat transposable elements, MULE-MuDR elements in particular, were located within genomic regions enriched for gene ontology (GO) functions related to the challenges of sanguivory, such as antigen processing and presentation, defence response to viruses, lipid metabolism, and vitamin metabolism (Supplementary Information 2).

A sanguivorous diet facilitates exposure to blood-borne viruses that could lead to an increase in genomic invasion by retroviral and non-retroviral endogenous viral elements (EVEs). Thus, we next characterized their presence in the common vampire bat genome. Compared to previously published EVE studies on non-Chiropteran mammals, the common vampire bat exhibits a greater diversity of non-retroviral EVEs in terms of the number of integrations, including endogenized viral genes from avian Bornaviridae

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\caption{**Comparative genomic analyses.** Species with a light blue circle were used for comparison of transposable elements. In golden colour are the silhouettes of the species used for the identification of putative lost genes and their corresponding number of putatively lost genes compared to the human genome. The number of gene families under expansion (green) and contraction (red) in Chiroptera using *H. europaeus* as an outgroup are indicated. Estimated dates of nodes (in million years) are indicated in black. The gut microbiomes of bats with different diets derive from *R. aegyptiacus* (blue, frugivorous), *M. gigas* (green, carnivorous), *R. ferrumequinum* (grey, insectivorous) and the common vampire bat (red, sanguivorous). *H. sapiens*, *Homo sapiens*; *E. helvum*, *Eidolon helvum*.}
\end{figure}
and Parvoviridae/Dependovirus. However, these findings are not restricted to vampire bats and are similar to those in other bat species25 (Supplementary Note 2, Supplementary Fig. 4 and Supplementary Information 3). Surprisingly, and in contrast to the prior expectations given its sanguivorous diet, the diversity of endogenous retroviral elements (ERVs) in the common vampire bat is very low compared to other bat species26. The only proviral elements detected were DrERV27 and DrgERV2, both present in low copy numbers (Supplementary Note 3, Supplementary Fig. 5 and Supplementary Information 4). We hypothesize that genome colonization by ERVs could have been restricted by the genomic adaptations in the common vampire bat genome against ERV insertion and proliferation. In support of our hypothesis, we identified expansion of the anti-retroviral gene TRIM5 family (Viterbi $P=0.00088$, Supplementary Information 5 and Supplementary Note 4).

**Genomic adaptations to sanguivory.** Feeding specialization often requires morphological and physiological adaptations in traits such as the sensory apparatus (for example, infrared sensing), locomotion, digestion, kidney function and immunity (Fig. 2a and Supplementary Information 6). For example, it has been shown that vampire bats have a loss of sweet taste genes and reduction of bitter taste genes28. In agreement, such genes were also identified in our putative gene loss analysis (Supplementary Information 7 and Supplementary Note 5). It is likely that the function of those genes is related to sanguivory, because sweet and bitter taste receptor genes influence glucose homeostasis in humans29. Interestingly, we found that the common vampire bat bitter taste receptor gene TAS2R3 has experienced episodic positive selection and shows two species-specific positively selected sites (PSSs) on topological domains, one of them having a potential impact on protein function (PROVEAN score = −4.4) (Supplementary Table 4 and Supplementary Fig. 6). Among the enriched GOs of the differentially evolving genes, we identified functions related to sanguivory, such as the sensory apparatus (for example, infrared sensing), locomotion, digestion, kidney function and immunity (Fig. 2b). However, it is clear that despite the number of detected genomic features related to sanguivory adaptation, they alone cannot address all of the challenges posed by this diet (Fig. 2b).

**Gut microbiome diet and phylogenetic influence.** We generated Illumina shotgun metagenome data in order to compare the gut microbiomes of 13 faecal samples from common vampire bats with those of non-sanguivorous non-phyllostomid bats; specifically, eight frugivorous Rousettus aegyptiacus (Egyptian fruit bat, Pteropodidae), five insectivorous Rhinolophus ferrumequinum (greater horseshoe bat, Rhinolophidae) and five carnivorous Macroderma gigas (ghost bat, Megadermatidae) bats (Supplementary Information 9). We obtained a median of 15.8 Gb of sequencing data (~37.6 million 100-bp paired-end reads) for each dietary category. After filtering low-quality bases, adaptor sequences and bat-genome-derived reads, we obtained a median of 2.77 Gb of high-quality data for each species, totalling 86.73 Gb of data. We identified taxa and functions present only in the common vampire bat microbiome (gut microbiome core), as well as taxa and functions that exhibit statistically significant differences in abundance or contribution to variation between the different microbiomes (Supplementary Information 6 and 8, Supplementary Tables 5 and 6 and Supplementary Note 8).

It has been observed previously that similarity in the taxonomic composition of vertebrate gut microbiomes (including bats) can be influenced by the diet and the phylogenetic relationships of the respective host species33. Overall, the common vampire bat microbiome taxonomic composition is more similar to that of the insectivorous and carnivorous bats than to that of the frugivorous bat. This may reflect a phylogenetic influence on the microbiome taxonomic profile (Fig. 3a, Supplementary Fig. 7 and Supplementary Notes 9 and 10). In contrast, the vampire bat microbiome is strikingly different to that of the compared bats at the functional level, which was characterized by the KEGG annotations of the microbial non-redundant gene catalogues assembled from the metagenomic data sets. While there is little differentiation between the functional gut microbiomes of carnivorous, insectivores and frugivorous bats, the common vampire bat functional gut microbiome is almost completely different to that of the compared bats at the functional level, which was characterized by the KEGG annotations of the microbial non-redundant gene catalogues assembled from the metagenomic data sets. While there is little differentiation between the functional gut microbiomes of carnivorous, insectivores and frugivorous bats, the common vampire bat functional gut microbiome is almost completely different, and exhibits the least intra-species variation between the samples (Fig. 3b, Supplementary Fig. 8 and Supplementary Table 7).

**Fig. 2 | Dietary challenges overcome by the common vampire bat.** a. Adapational contributions to sanguivory accountable to genomic changes alone (blue labels). b. Adapational contributions to sanguivory within a hologenomic context (blue labels for host genes; red labels for gut microbial traits).
This suggests that the functional profile is less influenced by phylogeny than the taxonomic profile, and that the common vampire bat gut microbiome harbours a set of functions specialized to its extreme diet (Supplementary Note 11). Subsequently, we analysed the comparative genomic and metagenomic results in a hologenomic framework to demonstrate how both components contribute to adaptation to sanguivory (Fig. 4, Supplementary Figs. 9–12, Supplementary Tables 7–9 and Supplementary Information 8 and 10).

The hologenomic framework of sanguivory. Viscosity and subsequent coagulation represent a challenge for ingestion and digestion of blood. Besides developing potent anticoagulants in its saliva34, the common vampire bat hologenome addresses this challenge in various ways. For instance, REG4, involved in metaplastic responses of the gastrointestinal epithelium, was found to be under ongoing positive selection (M8a/M8 test \( P = 0.047 \)) with possible functional impact on its carbohydrate-binding capacity, including binding of the anticoagulant heparin (Supplementary Table 4 and Supplementary Information 8). Furthermore, we identified genes in the common vampire bat microbial functional core related to pathways for degradation of heparan sulfate and dermatan sulfate, both being polysaccharides involved in blood coagulation (Supplementary Information 10). We also identified an enrichment in the common vampire bat microbial gene L-asparaginase (Fisher’s \( P = 0.00027 \)), which decreases protein synthesis of coagulation factors35 (Supplementary Information 11).

Besides specialized digestion, sanguivory poses other challenges related to the poor nutritional value of blood itself, as well as to the side effects that may arise due to the blood components being the sole dietary source (Fig. 2b). We identified elements in both the genome and gut microbiome that might be involved in solving each of these challenges discussed next.

Hologenomic solutions to nutritional challenges. Low nutrient availability. Obligate sanguivory requires adaptation to very low levels of some nutrients, such as essential amino acids and the vitamin B complex47, and very high levels of others, such as salt46. Our data clearly demonstrate how both the host and its associated gut microbiome have dealt with these challenges. We found the gene LAMTOR5 to be positively selected in the common vampire bat genome (false discovery rate (FDR)=2.02×10\(^{-6}\), Supplementary Information 8). This gene is involved in the response to nutrient starvation39, suggesting that the common vampire bat metabolism has adapted to the low nutrient content available in blood. Similarly, we identified KOs in the common vampire bat microbial core related to energy and carbohydrate metabolism (Fig. 4b and Supplementary Information 11). For example, when compared to the other bats, we identified an enrichment in the common vampire bat microbial genes involved in response to low nutrient availability (RelA/SpoT family protein, Fisher’s \( P = 0.0064 \), and guanosine pentaphosphate, Fisher’s \( P=0.0018 \)), and enzymes in the common vampire bat core involved in the reverse Krebs cycle (Supplementary Information 10), which is used by some bacteria to produce carbon compounds from CO\(_2\) and water (abundant blood components). We speculate that the presence of such metabolic pathways indicates the growth of specific microbes in the gut’s environmental conditions resulting from blood consumption.

The holobiont has also provided solutions to the lack of important nutrients in blood (Fig. 4c and Supplementary Information 8). For example, the PDZD11 gene in the common vampire bat genome, involved in vitamin B\(_6\) metabolism, evolved faster in the common vampire bat genome relative to the other examined bats (branch test \( P=1.97 \times 10^{-15} \)). We further postulate that the microbiome also contributes in tackling the low nutritional challenge by providing necessary nutrients. For example, compared to the other bat microbiomes, the common vampire bat gut microbiome had the highest number of enriched enzymes related to the biosynthesis of cofactors and vitamins, such as carotenoid (Supplementary Fig. 8B and Supplementary Information 10 and 11). Furthermore, we identified enzymes involved in the metabolism of butyrate, an
important nutrient for cells lining the mammalian colon derived from bacterial fermentation\(^6\), enriched in the common vampire bat gut microbiome as well as in the vampire bat gut microbiome core (Supplementary Information 10 and 11).

**Lipid and glucose assimilation.** Besides vitamins and other nutrients, lipids and glucose may not be readily available in blood. Furthermore, vampire bats have a reduced capacity to store energy reserves. In agreement with this, we identified GO enrichment of lipid metabolism on genes with a dN/dS ratio that is statistically higher or lower compared to other bats (Fig. 4a,c and Supplementary Information 8). For example, we identified the gene FFAR1, which plays an important role in glucose homeostasis, as evolving faster in the common vampire bat compared to the other bats (branch test \(P = 3.68 \times 10^{-5}\)) and containing amino-acid substitutions with a possible functional impact on its binding ability (Supplementary Table 4). This may enable the common vampire bat to better utilize the available glucose. The common vampire gut microbiome also exhibits unique solutions to the challenge (Fig. 4b,d and Supplementary Information 10 and 11). Differences in the carbohydrate and glycan metabolism functional profile were identified in the principal component analysis (PCA) comparing the different microorganisms (Supplementary Fig. 11), which place the vampire bat gut microbiome profile within a cluster separate from those of the non-sanguivorous bats. Importantly, we identified enrichment in the microbial gene glycerol kinase in the common vampire bat (Fisher’s \(P = 0.0027\)), which plays a key role in the formation of triacylglycerol and in fat storage, and its deficiency causes symptoms such as hypoglycaemia and lethargy in a mouse model\(^41\).

**Hologenomic solutions to non-nutritional challenges. Immunity.** Due to its sanguivorous lifestyle, the common vampire bat risks direct contact with blood-borne pathogens from prey. Consequently, we observed >280 bacterial species known to be pathogenic to some mammalian species present exclusively in the common vampire bat gut microbiome (Supplementary Information 12). For example, we identified enrichment of genes from *Borrelia* (Supplementary Information 11) and *Bartonella* as one of the most abundant genera in the common vampire bat compared to the other bats (Supplementary Information 13). These bacteria are known to be transmitted by sanguivorous invertebrates (ticks, fleas, mosquitoes and lice). This suggests that the abundance of this genus could be a shared pattern of sanguivorous species. While several studies have elucidated part of the expected genomic immunity-related adaptations\(^5\), analysis of the full genome enabled us to identify more elements related to immunity, such as defence response to virus and antigen processing and presentation (Fig. 4a,c and Supplementary Information 8). For example, we identified the antimicrobial gene *RNASE7* to be positively selected (branch-site test \(P = 0.004\)) and containing amino-acid substitutions that may increase its bactericidal capacity (Supplementary Table 4). In addition, when compared to the gut microbiomes of non-sanguivorous bats, that of the common vampire bat contains a large abundance of potentially protective bacteria, such as *Amycolatopsis mediterranei* (\(P < 0.05\)).

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**Fig. 4** | Traits in both the genome and gut microbiome with direct roles in the adaptation for sanguivory. a, GOs of the single-copy orthologous genes with a dN/dS ratio that is statistically higher or lower in the common vampire bat in comparison to the other bats. b, KOs from the common vampire bat gut microbiome functional core. c, Genes with a dN/dS ratio that is statistically higher or lower in the common vampire bat in comparison to the other bats and that are directly associated with sanguivory challenges. d, Taxa and genes from the common vampire bat gut microbiome core directly associated with the adaptation to sanguivory.
which has been shown to produce antiviral compounds against bacteria phages and poxviruses\(^2\) (Supplementary Information 13).

**Iron assimilation.** Iron concentration represents a significant challenge to sanguivory. Although the concentration of free iron is not high in the blood, severe haemolysis (for example, during digestion of blood) could result in high levels of iron that, if absorbed in excess, could accumulate and disrupt the normal function in organs such as the liver, heart and pancreas. Interestingly, we identified the light and heavy chains of the iron-storing protein ferritin (encoded by *FTL*, *FTH1*) under gene family expansion in the common vampire bat genome (*Viterbi* \(P = 0\) and *Viterbi* \(P = 0.0012\), respectively, Supplementary Information 5). In addition, we identified an enrichment of the iron-storing protein ferritin (Fisher’s \(P = 0.0014\)), suggesting that the gut microbiome also contributes to solving this challenge (Fig. 4d and Supplementary Information 11).

**Nitrogen waste and blood/osmotic pressure.** The high abundance of protein in the blood and its rapid ingestion could lead to accumulation of nitrogenous waste products, primarily urea, which could lead to renal disease-like symptoms (for example, high blood pressure and fluid retention). This challenge is exacerbated by the abundance of salts in blood, which pose additional osmotic and blood pressure challenges. We see at the genome level that this is addressed by a higher rate of evolution in the common vampire bat genes compared to the other bats involved in disposal of excess nitrogen (Fig. 4a,c and Supplementary Information 8), such as *PSMA3* (branch test \(P = 2.08 \times 10^{-10}\)). This challenge seems to also be addressed by the gut microbiome. The PCA of the copy number of genes involved in amino-acid metabolism distinguishes the common vampire bat in a single cluster separated from the other bat species analysed (Supplementary Fig. 11), suggesting a specialized microbial amino-acid metabolism capacity. We also identified enrichment in the common vampire bat microbial gene urease subunit alpha (*ureA*, Fisher’s \(P = 0.016\)) involved in urea degradation (Supplementary Information 11).

**Conclusions**

It is clear from our results that the common vampire bat has adapted to sanguivory through a close relationship between its genome and gut microbiome. We identified a phylogenetic and dietary impact on the common vampire bat gut microbiome and uncovered an unexpected genomic viral and repetitive element genomic makeup. We showed that extreme dietary specializations, such as that of the common vampire bat, provide a comparative framework with which to tease apart the relative roles of genomes and microorganisms in adaptation. In conclusion, our study illustrates the benefits of studying the evolution of complex adaptations under a holobiome framework, and suggests that vertebrate adaptation studies that do not account for the action of the hologenome may fail to recover the full complexity of adaptation.

**Methods**

**Genome sequencing and raw-read processing.** We shotgun-sequenced the *D. rotundus* genome using a wing biopsy from a sample collected by the NIH through the Catoctin Wildlife and Zoo in Thurmont, Maryland, USA. The capturing method and dead preservation procedure of the specimen are unknown. The age and sex of the dead individual are unknown. Sampling permits were given to BGI (Fig. 7a and Supplementary Information 5). For shotgun-sequenced paired-end sequences, reads with overlapping length \(\geq 100\) bp between the Read1 and Read2 were removed.

**Genome assembly.** We estimated the genome size of *D. rotundus* using KmerFreq\(^1\) by dividing the total number of seven decamers by the peak of the seven decamer Poisson distribution. High-quality reads were assembled using SOAPdenovo\(^2\) as follows. (1) Short-insert library reads were assembled as initial contigs ignoring the sequence pair information. (2) Reads were aligned to the previously generated contig sequences. Scaffolds were constructed from short-insert-size libraries to large-insert-size libraries by weighting the paired-end relationships between pairs of reads or at least three reads pairs required for forming the gap between any two contigs. (3) Gaps in the scaffolds were closed using GapCloser\(^3\). Genome quality assessment was performed by downloading a publicly available *D. rotundus* transcriptome\(^4\) and aligning the transcripts to the genome using BLAT\(^5\).

**Genome contiguity improvement.** We prepared two Chicago libraries\(^6\) using 5 \(\mu\)g of high-molecular-weight DNA obtained from *D. rotundus* cultured cells from the San Diego Zoo collection, which were originally derived from a skin sample taken from between the shoulder blades of a *D. rotundus* individual. Permits for this were obtained from the San Diego Zoo Global. The capturing method and dead preservation procedure of the specimen are unknown. The age and sex of the dead individual are unknown. DNA was extracted with Qiagen Blood and Cell Midi kits according to the manufacturer’s instructions. The steps required for building the Chicago libraries were performed as described in ref. \(^7\). The libraries were sequenced using Illumina HiSeq 2500 \(\times 100\) bp rapid run. Our initial *D. rotundus* assembly, shotgun sequence data and Chicago library sequences were used by Dovetail Genomics as input data for HiRise, as described in ref. \(^8\). Genome assembly contiguity statistics were obtained using a minimum N track length of 1 to delimit the contig blocks within the scaffolds.

**Protein-coding gene and functional annotation.** Homology-based gene prediction was performed using a reference the Ensembl genome set of *Myotis lucifugus*, *Pteropus alecto*, *Myotis davidii*, horse and human. We aligned the protein sequences of the reference gene sets to the *D. rotundus* assembly using tblastn\(^9\) and linked the blast hits into candidate gene loci with genBlastA\(^10\). We filtered out those candidate loci with homologous block length <90% of the query length. We extracted genomic sequences of candidate gene loci, including the intronic regions and 3 kb upstream/downstream sequences. The sequences were passed to GeneWise\(^11\) to search for accurately aligned alignments. We filtered out pseudogenes containing more than one frame error for single-exon genes. Potentially pseudogenized single exons were removed if they were part of a multi-exon gene. We then aligned protein sequences of these genes against UniProt using blastp and filtered out genes without matches. We also filtered out genes that had >80% repeat regions. De novo gene prediction was performed with AUGUSTUS\(^12\) using a published common vampire bat transcriptome\(^13\) as a training data set and with masked transposable-element-related repeats. We filtered out partial and <150 bp predicted genes. Genes that aligned over 50% of their length to annotated protein database were filtered out. Finally, we built a non-redundant protein set with the homology-based evidence prioritized over the de novo evidence. If de novo genes were chosen in the reference gene set, we retained only those with >30% of their length aligning against UniProt\(^14\) and that contained at least 3 exons. The integrated gene set was translated into amino-acid sequences, which were used to search the InterPro database with iprscan_4.8\(^15\). We used the BLAST to search the metabolic pathway database in KEGG\(^16\) and homologues in the SwissProt and TrEMBL databases in UniProt. The quality and annotation of the *D. rotundus* genome was assessed and compared to that of *E. europaeus*, *R. ferrumequinum*, *M. brandtii*, *M. davidii*, *M. lucifugus* and *P. parnellii* with BUSCO\(^17\).

**Repeat annotation.** Repeat annotation was performed on the genomes of *D. rotundus*, *P. parnelli*, *M. lyra* and *V. vampyrus*. Transposable elements were identified using RepeatMasker\(^18\) and RepeatProteinMask against the Repbase transposable element library\(^19\). We used RepeatScount, PILER-DF and RepeatModeler-1.0.5\(^20\) to construct a de novo transposable element library, which was then used by RepeatMasker to predict repeats. We predicted tandem repeats using TRF\(^21\), LTR_Finder\(^22\) was used to detect long terminal repeats (LTRs). The Repbase-based annotations and the de novo annotations were merged.

**Non-retroviral EVs.** We constructed a comprehensive library of all non-retroviral virus protein sequences available in GenBank and EMBL. We used DIAMOND\(^23\) to search these sequences against the *D. rotundus* genome. We extracted the matching amino-acid sequences and performed reciprocal blast-like searches using DIAMOND with the selected subset of *D. rotundus* amino-acid sequences and the set of non-redundant protein sequences. *D. rotundus* genome sequences were considered of viral origin if they unambiguously matched viral
proteins in the reciprocal best hits. Putative viral open reading frames were inferred from automated alignments, using exonerate. For each putative D. rotundus viral peptide, we retrieved the function and predicted the taxonomic assignment by comparison to the best reciprocal blastp-like hit viral proteins. Phylogenetic analyses were performed by aligning the sequences using MAFFT and curated using AliView. Maximum-likelihood inferences were performed on each multiple amino-acid alignment using RAxML. Support for nodes was obtained from 100 non-parametric bootstrap iterations, and the root of maximum-likelihood trees was determined by midpoint rooting. We confirmed the presence of selected viruses by constructing double-indexed Illumina libraries on DNA derived from the spleen tissue of four different D. rotundus. Libraries were pooled and sequenced using 150-bp paired-end Illumina platform NextSeq 500. The raw paired-end reads were quality assessed, merged and filtered by mapping against the bacterial, human and chiropteran reference genome database with SMALT (tiptop). PSSs were filtered with SMALT software. ERV sequences were retrieved through a viral assignment pipeline using blastx to search against the viral non-redundant protein database. Viral-matching reads were assigned a taxonomy with MEGAN. Reads matching viral sequences were manually verified by reciprocal blastx analysis. Positive viral hits were defined as those with at least two different reads matching two different proteins or two different regions within the same viral protein.

ERVs. We identified flanking LTR regions using RepeatMasker and RMBLAST (http://www.repeatmasker.org/RMBlast.html), Tandem Repeats Finder and RepeatMasker to the D. rotundus genome. We retrieved retro-viral reference genome sequences in the RefSeq dataset of NCBI against the D. rotundus genome. Reads matching each of the RefSeq sequences were extracted and sorted by GI, collapsed by ID and manually verified by reciprocal blast. Tblasts were re-run and the kept reads were mapped to the D. rotundus genome with SMALT and manually verified by reciprocal blast. ERV sequences validation was carried out using tblasts against the Retroviridae. The retrieved ERV sequences were mapped using SMALT v0.7.6 against the genomes of D. rotundus, Mastomys natalensis, Eptesicus fuscus, M. lucifugus, M. brandtii, D. alecto, R. aegyptiacus and P. paradiisi. Tblasts of the full genomic sequences of D.ERV and DgrERV was also performed against Chirottera. Selected genomic contigs were further analysed by reciprocal blast against the non-redundant GenBank coding DNA sequence (CDS) translations + PDB + SwissProt + PIR + PRF database restricting the search to the Retroviridae, or against the DgrERV sequences. Retrieved sequences were aligned using MUSCLE in SeaView. The best-fit amino-acid substitution model for each alignment was identified using ModelTest and trees were inferred under maximum-likelihood criteria using RAxML.

Orthologous gene families. Using the D. rotundus genome against a range of other mammalian species, we performed clustering of orthologous genes using jModelTest2 and trees were inferred under maximum-likelihood criteria. Genes with significant P values. To adjust for multiple testing, we used the FDR method. To identify genes with positively selected sites in D. rotundus, we also used a branch site with PAML codeml. LRT and FDR were computed as carried out for the branch model tests.

Gene family expansion/contraction. We used CAFE on the results from the single-copy orthologous gene families. To obtain the dated tree required as input for CAFE, we obtained divergence times from the fossil dating records from TimeTree. We concatenated the CDSs aligned with MAFFT and used PAML mcmtree to determine split times with the approximate likelihood calculation method. Genes with >200 copies in 1 of the species were filtered out.

Putative gene loss. We identified genes putatively lost in D. rotundus as previously described in ref. We filtered genes putatively lost in D. rotundus as previously described in ref. We filtered genes putatively lost in D. rotundus as previously described in ref. The human gene set was downloaded from Ensembl and we obtained a non-redundant gene catalogue to be used as a reference by collapsing redundant genes and keeping the longest open reading frame. We used Bos taurus, Equis caballus and E. europaeus as outgroups. Genes identified as missing in D. rotundus were retrieved against the GenBank databases of the other available bath species. Further validation on selected genes was performed by: evaluating the conservation of their syntenic regions compared to other species, and the GC content of the syntenic regions compared to the D. rotundus genome average GC content; searching them against the published D. rotundus transcriptome; and through PCR assays.

Functional characterization. We performed GO analysis using GOrilla as well as manual characterization using the GO annotations of the human genes downloaded from Ensembl and literature mining. We used PROVEAN to characterize the functional impact of the non-synonymous substitutions present only on the proteins in D. rotundus.

Species-specific PSS identification and protein modelling. We tested for positive selection and positive selected sites (PSSs) using the complete CDS alignments for the proteins FFAR1, PLXNA4, REG4, RNAS7 and T2A23 from various species downloaded from the OrthoMCL database. Sequences were realigned using MUSCLE in SeaView and phylogenetic analysis was performed using PhyML. We used PAML codeml to test: the M1/M2 branch model constraining the D. rotundus node; the M8A/M8 site model; and the branch-site model A constraining the D. rotundus node. The branch-site model A was evaluated under a LRT against the null model without branch-site constraints. We scored were scored under naive empirical Bayes (NEB) and Bayes empirical Bayes (EBE). We performed modelling of the mentioned proteins from D. rotundus using the PyMOL Molecular Graphics System (Schrodinger, LLC). The three-dimensional protein models of the D. rotundus sequences were constructed using Phyre2 based on profile hidden Markov model analysis using the structures reported in refs.

Metagenomic data. We used fecal and anal swab samples from D. rotundus, R. aegyptiacus, R. ferrumequinum and M. gigas. The D. rotundus samples were obtained from wild D. rotundus individuals captured using mist nets and cloth traps and from the non-sanguivorous bat species. The R. aegyptiacus samples were collected from the Copenhagen Zoo. We have complied with all relevant ethical regulations for the collection of these samples. DNA from fecal samples were PCR-amplified targeting the 16S rRNA gene and the PCR products were cloned into a pCR2.1 vector (Invitrogen). The R. ferrumequinum samples were sequenced and cDNA isolation Kit (MoBio) following the manufacturer’s protocol. All samples were kept at −20°C. Libraries were built using the NEBNext DNA Library Prep Master Mix Set for 454 (New England BioLabs) following the manufacturer’s instructions. Samples were 100-bp pair-end sequenced on the Illumina 2500 HiSeq platform.

Taxonomic and functional identification. The reads were cleaned with trimmomatic and prinseq-lite. The data sets were mapped against the closest available bat genome and only the non-mapping reads were kept. Mgmapper was used for the taxonomic identification in invertebrates, protozoa, fungus, virus and bacteria. We kept the species identified with a coverage value higher than the first-quartile from the coverage distribution of the corresponding database and filtered out those found on the corresponding extraction blanks. Rarefaction curves from each data set were obtained using an in-house script with the MGmapper results. We then performed de novo assembly using IDBA_UD, predicted genes using Prodigal and generated a non-redundant gene catalogue with usearch. The non-redundant catalogue was searched against the UniProt database for functional and taxonomic annotation. We used DIAMOND to search the unmapped reads against the UniProt database keeping only the best hit for functional and taxonomic annotation. We annotated the UniProt protein identifications using KEGG orthology (KO) and eggNOG IDs.

Taxonomic and functional metagenomics comparison. We filtered on the basis of the breadth of coverage and the identifications of the extraction blanks. We removed any non-microbial hit and any taxa in which the paired reads matched different genera or only one of the reads had a hit. The counts were normalized by percentage. We identified a microbial taxonomic and functional sanguivorous core by comparing the filtered sets of the bats and keeping as core those taxa and gene functions common to all samples. We removed any non-microbial hit and any taxa in which the paired reads matched the filtered taxonomic identifications, and the KO and COG annotations from the filtered non-redundant gene set catalogue. We compared the normalized abundance of taxa and functions between D. rotundus and the non-sanguivorous bats as follows. (1) Using the distributions of the different functional categories from D. rotundus and each non-sanguivorous bat species with a Wilcoxon rank-sum test. (2) Using the entire taxonomy and functional data sets, as well as down-sampling the normalized count values. Sampling values were the minimum, median and third-quartile values of the count distributions. With the resulting data sets, we calculated the Euclidean, Bray–Curtis and Jaccard distance metrics with the corresponding extraction blanks. Rarefaction curves from each data set were obtained using an in-house script with the MGmapper results. We then performed de novo assembly using IDBA_UD, predicted genes using Prodigal and generated a non-redundant gene catalogue with usearch. The results from each data set were obtained using an in-house script with the MGmapper results. We determined the abundance of taxa and functions between the filtered taxonomic identifications, and the KO and COG annotations from the filtered non-redundant gene set catalogue. We compared the normalized abundance of taxa and functions between D. rotundus and the non-sanguivorous bats as follows. (1) Using the distributions of the different functional categories from D. rotundus and each non-sanguivorous bat species with a Wilcoxon rank-sum test. (2) Using the entire taxonomy and functional data sets, as well as down-sampling the normalized count values. Sampling values were the minimum, median and third-quartile values of the count distributions. With the resulting data sets, we calculated the Euclidean, Bray–Curtis and Jaccard distance metrics with the R package vegan, and used the Ward hierarchical clustering method using UPGMA and a p-value with p<0.01, and performed PCAs with precomp and the GPA method. (3) We identified taxa and functions significantly contributing to the variation between the D. rotundus and the non-sanguivorous bat species. We examined the rotation matrix from the PCA of the normalized counts, excluding the four deepest sequenced samples, of the species and genus microbial taxonomic levels and the KEGG functional pathways. We identified the most significantly abundant
D. rotundus microbial taxa as those with a significantly higher median normalized count value (P < 0.05) in D. rotundus and a median and mean normalized count value of 0 in the other 3 bat species for the first 3 principal components. We also identified significantly more abundant genes in D. rotundus by generating and annotating with KEGG a non-redundant gene set with all of the predicted genes from all of the bat samples. The reads of the samples were mapped against this bat non-redundant gene set, and a normalized count matrix was generated and used for Fisher tests on each of the functional pathways.

Life Sciences Reporting Summary. Further information on experimental design is available in the Life Sciences Reporting Summary.

Code availability. In-house scripts used for the processing of the data are available from the corresponding authors upon request.

Data availability. The NCBI BioProject accession code for the genome assembly is PRJNA444273, and the sequence reads are available at the NCBI sequence read archive (SRA) under the accession SRA619672. The BioProject code for the metagenomic sequencing data is PRJNA415003, and the reads can be accessed at SRA with the accession SRA620977.

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Competing interests
The authors declare no competing financial interests.

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Experimental design

1. Sample size
   Describe how sample size was determined.
   - We obtained as many samples as possible for the chosen species to perform our metagenomics comparisons.

2. Data exclusions
   Describe any data exclusions.
   - We used all the generated data. Different sets for the different analyses, each one described in the extended methods. For example, the identification of the microbiome cores was done using all the datasets, and the microbiome profiles distances were obtained both using all the sets downsampled, and excluding the 3 most heavily sequenced samples to avoid bias.

3. Replication
   Describe whether the experimental findings were reliably reproduced.
    - All attempts at replication were successful

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups.
   - Allocation was done by species. This is not relevant to the comparative genomics. And for the metagenomics, we controlled the variability by normalizing the sequencing depths and performed the appropriate tests for finding enrichments. For identification of the microbial cores that is also not relevant.

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   - Blinding is not relevant to our analyses, which compare the genomes and metagenomes of the species, our analyses do not make predictions of their microbes/genes, but characterize their identifications.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
   For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

   n/a  Confirmed

   - The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
   - A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
   - A statement indicating how many times each experiment was replicated
   - The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
   - A description of any assumptions or corrections, such as an adjustment for multiple comparisons
   - The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
   - A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)

   See the web collection on [statistics for biologists](https://statisticsforbiologists.com) for further resources and guidance.
### Software
Policy information about availability of computer code

7. Software
Describe the software used to analyze the data in this study.

We used many different softwares. All of them are specified in the Methods sections in the main and supplementary text, including the versions. In house scripts upon request.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

### Materials and reagents
Policy information about availability of materials

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

Laboratory reagents are all commercially available. The unique materials are the bat fecal samples and *Desmodus rotundus* skin cells, which were all used for the generation of the sequencing data.

9. Antibodies
Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

We did not use any antibodies.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

b. Describe the method of cell line authentication used.

No eukaryotic cell lines were used.

c. Report whether the cell lines were tested for mycoplasma contamination.

No eukaryotic cell lines were used.

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

N/A

### Animals and human research participants
Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals

Provide details on animals and/or animal-derived materials used in the study.

We used samples from *Deamodus rotundus*, *Roussettus aegyptiacus*, *Rhinolophus ferrumequinum*, *Macrotus gigas*, *Rhinolophus euryale*, *Miniopterus schreibersi*, *Nyctalus leisleri*, and *Nyctalus lasiopterus*. Physiological information of the vampire individuals sampled by net capturing is presented in the Supplementary Information.

Policy information about studies involving human research participants

12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

The study did not involve human research participants.