



Revisiting fecal metatranscriptomics analyses of macaques with idiopathic chronic diarrhoea with a focus on trichomonad parasites

Research Article

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Abstract

Trichomonads, anaerobic microbial eukaryotes members of the phylum Parabasalia, are common obligate extracellular symbionts that can lead to pathological or asymptomatic colonization of various mucosal surfaces in a wide range of animal hosts. Results from previous *in vitro* studies have suggested a number of intriguing mucosal colonization strategies by Trichomonads, notably highlighting the importance of interactions with bacteria. However, *in vivo* validation is currently lacking. A previous metatranscriptomics study into the cause of idiopathic chronic diarrhoea in macaques reported the presence of an unidentified protozoan parasite related to *Trichomonas vaginalis*. In this work, we performed a reanalysis of the published data in order to identify the parasite species present in the macaque gut. We also leveraged the information-rich metatranscriptomics data to investigate the parasite behaviour *in vivo*. Our results indicated the presence of at least 3 genera of Trichomonad parasite; *Tetratrachomonas*, *Pentatrachomonas* and *Trichomitus*, 2 of which had not been previously reported in the macaque gut. In addition, we identified common *in vivo* expression profiles shared amongst the Trichomonads. In agreement with previous findings for other Trichomonads, our results highlighted a relationship between Trichomonads and mucosal bacterial diversity which could be influential in health and disease.

Introduction

Trichomonads are a group of microbial eukaryotes within the phylum Parabasalia, almost all of which are known as obligate mucosal symbionts that colonize a wide range of mammals, birds and reptiles (Malik *et al.*, 2011). Parasitic Trichomonads have no known free-living stages and are assumed to be transmitted almost exclusively by direct contact. The molecular basis of virulence, mucosal colonization (Sommer *et al.*, 2005; de Miguel *et al.*, 2010; Handrich *et al.*, 2019; Martínez-Herrero *et al.*, 2019) and metabolism (Matthews, 1986; Westrop *et al.*, 2017) of Trichomonads has been the subject of extensive *in vitro* investigation. The vast majority of this work has focused on the human sexually transmitted parasite *Trichomonas vaginalis*. However, the importance of the proposed mechanisms during colonization of the complex mucosal environment *in vivo* is unclear. Validation of hypotheses in the natural setting is essential to avoid misinterpretation of results (Bello-Ortí *et al.*, 2015; Marzano *et al.*, 2017). The conservation of genes encoding virulence and mucosal colonization mechanisms across a wider range of Trichomonad species is also largely unknown, necessitating comparative studies.

There is extensive evidence for an interaction between Trichomonads and mucosal bacteria which is influential in health and disease. For example, *T. vaginalis* infection can induce dysbiotic changes in the urogenital tract (UGT) microbiota (Fichorova *et al.*, 2013). Such results have been validated *in vivo* in several Trichomonad species and hosts (Wei *et al.*, 2020; Bierlein *et al.*, 2021). Notably, *Trichomonas gallinae* infection was correlated with changes in the microbiota at local and distant mucosal sites in pigeon squabs (Ji *et al.*, 2020). However, previously, methods have been exclusively limited to 16S profiling, which provides no information on the potential mechanisms underlying parasite–bacteria interactions. *T. vaginalis* is a phagocytic predator of bacteria (Juliano *et al.*, 1991; Rendon-Maldonado *et al.*, 1998) and fungi (Pereira-Neves and Benchimol, 2007), with some evidence for selective preference of prey species (Juliano *et al.*, 1991). In addition, *T. vaginalis* can form symbiotic associations with *Mycoplasma* spp. (Dessi *et al.*, 2019). Functional work is required to determine the contribution of predation, symbiosis or other mechanisms to Trichomonad-induced *in vivo* microbiota changes.

A recent fecal metatranscriptomics investigation by Westreich *et al.* (2019) into the cause of idiopathic chronic diarrhoea (ICD) in laboratory macaques revealed the presence of GIT-localized protozoa. The authors stated ‘Protozoans with the most abundant transcripts in the faecal samples from the macaques were *Blastocystis* sp. and *Trichomonas vaginalis*’; however, also qualified ‘*T. vaginalis* was the only species in the reference data set representative of the *Trichomonas* genus, so it is possible that the particular species with increased gene expressed in macaques with ICD was not *T. vaginalis*’. Trichomonads do exhibit mucosal and host plasticity (Maritz *et al.*, 2014). For example, *T. vaginalis* has been detected in the oral cavity (Costello *et al.*, 2017) and respiratory tract (Duboucher *et al.*, 2003). In addition,

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T. vaginalis is thought to originate from zoonosis of an avian oral parasite (Maritz *et al.*, 2014). However, *T. vaginalis* is essentially a human UGT parasite. We suggest that misidentification as '*T. vaginalis*' due to sequence database incompleteness (Watts *et al.*, 2019) is very likely, as there are no reports of *T. vaginalis* in non-human animals or the GIT. Thus, further investigation into the parasite identity is warranted. Studies conducted on wild macaques did not report the presence of intestinal Trichomonads (Adhikari and Dhakal, 2018), although infection with *Pentatrichomonas hominis* has been reported in immunocompromised laboratory macaques (Zaragoza *et al.*, 2011) and those suffering from ICD (Laing *et al.*, 2018). Thus, *P. hominis* is a likely candidate for the parasite infecting the macaques with ICD.

In this work, we utilized the existing fecal metatranscriptomics data from macaques (Westreich *et al.*, 2019) to identify the Trichomonads present, and to investigate their *in vivo* gene expression. Our results also suggested relationships between Trichomonads and the mucosal microbiota *in vivo*, with potential implications for the aetiology of ICD.

Materials and methods

Macaque fecal metatranscriptomics data analysis

Full details for experimental methodology used to generate the previously published macaque fecal metatranscriptomics data are available from Westreich *et al.* (2019). Briefly fecal samples were collected from 12 macaques suffering from ICD and 12 healthy control animals. For 30 days prior to sample collection animals were housed indoors in pairs, separating macaques with ICD and healthy controls. For stool collection, cage pans were placed under the cages of individually housed animals overnight and collected in the morning. Stool samples were used as a proxy for the intestinal mucosal environment. Total RNA was extracted from stool samples and used for cDNA library preparation.

The workflow used to assess macaque fecal metatranscriptomics data is shown in Fig. S1. The metatranscriptomics dataset for macaques was obtained from the NCBI SRA database (Leinonen *et al.*, 2011) under accessions SRX3517701-SRX3517724 (Westreich *et al.*, 2019). There were approximately 95 million paired end reads per animal. The average Phred quality score for all reads did not fall below 30, as assessed using FastQC version 0.11.9 (Andrews, 2018). For quantitative analysis of taxonomic abundance and functional expression, reads derived from rRNA were filtered by alignment to a prokaryotic and eukaryotic rRNA database using SortMeRNA version 4.2.0 (Kopylova *et al.*, 2012). If both reads in a pair aligned, the pair was excluded. Kraken2 version 2.0.8-beta (Wood *et al.*, 2019) with default parameters was used to taxonomically classify reads. The Kraken2 reference database was enriched by including *de novo* assembled contigs derived from *in vitro* RNA-Seq data for *P. hominis* strain PhGII, *Tetratrichomonas gallinarum* strain M3, and *Trichomitus batrachorum* strain BUB, kindly provided by Sriram Garg and Sven Gould (Handrich *et al.*, 2019) (Heinrich Heine University, Düsseldorf) to improve the Trichomonad sequence diversity. The NCBI Taxonomy Toolkit version 0.5.0 (Shen and Xiong, 2019) was used to manipulate taxonomy IDs generated by Kraken2.

A *de novo* assembly was generated from the data using metaSPAdes version 3.13.0 (Nurk *et al.*, 2017). To assess the accuracy of the assembly, reads were aligned to assembled contigs using STAR version 2.7.3a (Dobin *et al.*, 2013). Samtools version 0.1.20 (Li *et al.*, 2009) was used to manipulate alignment files. Overview for the dataset and assembly are presented in Table S1. *De novo* assembled contigs derived from combined Parabasalia reads from all samples to maximize coverage (available in Data files S1, selected genes, and S2, all genes) were

used to examine Parabasalia gene expression. For transcript annotation, assembled parasite contigs were aligned to the *T. vaginalis* G3 annotated proteins (Carlton *et al.*, 2007) using BLASTx version 2.9.0+ (Altschul *et al.*, 1990). A single top hit for each contig was selected after sorting by, respectively and in priority order, E value and percentage identity, and excluding hits with an E value greater than 1×10^{-10} or a query coverage of less than 70%. Annotated contigs are presented in Table S2 (specific genes used as phylogenetic markers) and Table S3 (all genes).

Phylogenetic analysis

To sequence type parasites, BLASTn version 2.9.0+ (Altschul *et al.*, 1990) was used to identify *de novo* assembled contigs homologous to Parabasalia genes of interest, with E value, percentage identity and query coverage cut-off values of 1×10^{-10} , 88% and 90%, respectively. To broaden the taxonomic sampling for genes of interest, additional homologues were identified by consulting the literature and through the use of online BLAST (Altschul *et al.*, 1990) searches against the NCBI non-redundant protein or nucleotide databases (O'Leary *et al.*, 2016). Alignments were generated using Clustal omega version 1.7 (Sievers *et al.*, 2011) and visually inspected in Seaview version 5.0.4 (Gouy *et al.*, 2010). To improve phylogenetic resolution of parasite sequencing typing, DNA alignments for protein-coding genes were generated. Protein sequences were aligned using Clustal omega version 1.7 (Sievers *et al.*, 2011), and corresponding codon alignments were derived using pal2nal version 14 (Suyama *et al.*, 2006). Poorly aligned sequences were removed, and alignments were trimmed to remove excessive gaps (sites containing a gap for greater than 90–95% of sequences) using TrimAl version 1.2 (Capella-Gutiérrez *et al.*, 2009). Alignments are available in supplementary data files S3–S8. IQ-tree version 1.6.1 (Nguyen *et al.*, 2015) was used to generate maximum likelihood phylogenies, using automatic model selection. Support for tree topology was assessed by computing 1000 bootstrap replicates. iTol (Letunic and Bork, 2019) was used to generate annotated figures from the phylogenies.

Microbial diversity and expression analysis

Microbial diversity analysis was performed using the R packages PhyloSeq version 1.34.0 (McMurdie and Holmes, 2013) and Microbiome version 1.12.0 (Lahti and Shetty, 2017), excluding reads assigned within equivalent or child taxa to animals, viruses or Parabasalia. ANCOM-BC version 1.0.5 (Lin and Peddada, 2020) was used for differential abundance analysis, excluding reads assigned within equivalent or child taxa to animals or plants. SparCC (Friedman and Alm, 2012) was used for microbial correlation analysis, including only bacterial and parabasalid genera representing at least 0.005% of the sequencing library in at least 1 sample. Bootstrapped samples (100 replicates) of microbial abundance were used to calculate 2-sided pseudo-*P* values. Microbial correlation networks were derived from the sparCC results using the R package igraph version 1.2.11 (Csardi and Nepusz, 2006), with edges linking genera sharing a correlation coefficient greater than 0.8 and pseudo *P* value lower than 0.05. Networks were split into modular components by the Louvain method (Blondel *et al.*, 2008) and Cytoscape version 3.6.1 (Shannon *et al.*, 2003) was used to generate figures and calculate network summary statistics. For functional analysis of microbial transcription, reads assigned within equivalent or child taxa to animals, plants, Parabasalia or viruses were excluded. HUMAnN version 2.8.2 (Franzosa *et al.*, 2018) was used to functionally classify reads at the gene family level by translated alignment to UniRef90 protein families (Suzek *et al.*, 2015). Gene

family abundance values were normalized to library size (in counts per million; CPM) prior to assignment to MetaCyc pathways (Caspi *et al.*, 2016) to calculate pathway abundance using HUMAnN version 3.0.0 (Beghini *et al.*, 2021). Pathway abundance values were \log_2 transformed, with an added pseudocount of 0.01, before differential abundance test by the limma-trend method (Law *et al.*, 2014).

Results

Identity of trichomonads colonizing the macaque gut

We performed an analysis on the unidentified Trichomonads which were reported in published fecal metatranscriptomics data from rhesus macaques with ICD (Westreich *et al.*, 2019). Metatranscriptomes were available for 12 macaques with ICD (Macaques 1–12) and 12 healthy control animals (Macaques 13–24).

We aimed to investigate the identity of Trichomonad parasites reportedly present in this dataset (Westreich *et al.*, 2019). To generate sequences for molecular typing, we generated a *de novo* assembly of the metatranscriptome. Contig length statistics suggested an overall low degree of assembly, with a mode contig length of 161 bp across all samples, and N50 values ranging from 544 to 1021 bp. Alignment of reads to the assembly indicated no major compositional biases (Fig. S2). Summary statistics for the dataset and assembly are presented in Table S1. The *de novo* assembly is available in Data file S2.

Due to the low sequence coverage and high complexity of parasite sequences, we utilized the 18S rRNA, actin and elongation factor 1 alpha (EF-1 α) loci to identify the Parabasalia colonizing the macaque gut. Amongst all the samples, we assembled 58, 10 and 11 18S rRNA, actin and EF-1 α sequences, respectively, which shared greater than 88% sequence identity with reference Parabasalia sequences for at least 90% of their length (Table S2). We assessed the diversity of sequences present by maximum likelihood analysis and identified 10 well-supported clades for the 18S rRNA locus (Fig. S3), 4 for the actin locus (Fig. S4), and 1 for the EF-1 α locus (Fig. S5). We generated phylogenies using representative sequences from each clade alongside a range of Parabasalia reference sequences in order to refine the identity of the parasite sequences. Analysis of a single representative sequence from each of the 18S rRNA sequence groups revealed at least 3 major lineages, related to *Tetratrichomonas*, *Pentatrichomonas* and *Trichomitus* spp., with strong bootstrap support only present for the latter (99%; Fig. 1). In contrast, there was strong bootstrap support (99%) for grouping of all identified actin sequences with *Tetratrichomonas gallinarum* (Fig. 2), and all identified EF-1 α sequences with *P. hominis* (100%; Fig. 3). Integrating these analyses, we inferred that there are likely to be 3 Parabasalia lineages, related to *Trichomitus*, *Tetratrichomonas* and *Pentatrichomonas*, present amongst the macaque fecal samples.

To taxonomically assign the metatranscriptome reads, we included *de novo* assembled contigs derived from *in vitro* RNA-Seq analysis of *P. hominis*, *Tetratrichomonas gallinarum* and *Trichomitus batrachorum* (Handrich *et al.*, 2019) in the reference database to improve assignment for sequences derived from the putative parasite genera of interest. In agreement with the phylogenetic results, *Trichomitus*, *Pentatrichomonas* and *Tetratrichomonas* were the 3 most abundant (mean across all samples) parabasalid genera which were identified (Fig. 4A). According to read assignment, *Trichomitus* was the most abundant individual genus of interest (mean abundance 0.096%), followed by *Pentatrichomonas* (mean abundance 0.025%) and *Tetratrichomonas* (mean abundance 0.020%). A substantial number of sequences (mean abundance 0.093%) were identified as

parabasaliid in origin but could not be assigned to a particular genus. Unidentified parabasalid reads appeared more abundant among animals in which *Tetratrichomonas*, *Pentatrichomonas* or *Trichomitus* classified reads were abundant, likely suggesting that they originated from 1 or more of these genera.

In addition, a notable fraction of reads were classified as *Trichomonas* (mean abundance 0.019%). This most likely reflects the greater representation of *Trichomonas* whole genome sequences available in the reference database, including *T. vaginalis* and *T. gallinae*, whereas the genera of interest were only represented by *in vitro* RNA-Seq data, which is likely to have an incomplete gene content. However, while it cannot be ruled out that *Trichomonas* spp. were present amongst the samples, we have focused our analysis on the most likely genera based on the phylogenetic results. Only 2 control macaques showed a total abundance of Parabasalia greater than 0.125%, limiting the statistical power for tests correlating variables with Parabasalia abundance amongst the control animals.

Trichomonad gene expression

We focused on the most abundant putative Parabasalia-derived contigs to explore the most biologically important functions, which are summarized in Table 1. Potential energy-generation pathways included glycolysis, hydrogenosome metabolism, catabolism of GlcNAc, GalNAc, galactose and glucosamine and amino acid catabolism, including the arginine dihydrolase (ADH) pathway. The presence of a putative xanthine dehydrogenase could also indicate catabolism of nucleotides as an additional nutrient source (Wang *et al.*, 2016). Synthesis of glucose and storage as glycogen was suggested by gluconeogenesis and glycogen processing enzymes.

A number of Parabasalia contigs were annotated with putative lysozyme activity, thus potentially targeting the bacterial microbiota. A maximum likelihood phylogeny was generated to investigate the possibility that the contigs were bacterial in origin (Fig. S6). Results suggested that the lysozyme-like contig NODE_2008 originated from *Trichomitus* with strong support, and an additional contig NODE_1833 may have originated from *Pentatrichomonas* or *Tetratrichomonas*, although this was poorly supported. In addition, we identified a contig with high similarity to *T. vaginalis* coronin, an actin-binding protein implicated in phagocytosis (Bricheux *et al.*, 2000), thus consistent with parasite phagocytosis targeting microbial or host cells.

Numerous contigs showed strong similarity to *T. vaginalis* genes previously implicated in pathobiology. Of particular interest for parasite adhesion to host or microbial cells (Handrich *et al.*, 2019), we detected expression of 762 contigs with substantial sequence similarity to *T. vaginalis* BspA proteins. As the BspA family represent strong candidate LGTs of prokaryotic origin (Handrich *et al.*, 2019), it is likely that this list could include bacterial contigs, due to high similarity between the Trichomonad and bacterial sequences. The list included 70 contigs with greater than 60% sequence identity with the nearest *T. vaginalis* homologue (by BLASTx). Cysteine peptidases are also implicated in *Trichomonas* pathobiology (Sommer *et al.*, 2005) and 93 contigs were detected which shared high similarity with *T. vaginalis* Clan CA, family C1, cathepsin L-like cysteine peptidases. Of particular interest, 17 contigs were close homologues of TvCP39 (locus tag TVAG_298080; mean percentage identity 69%), a secreted cysteine peptidase demonstrated to induce host cell apoptosis (Arroyo *et al.*, 2015).

Trichomonad interactions with the microbiota

To further investigate potential interactions between parabasalid parasites and the microbiota, we examined the taxonomic

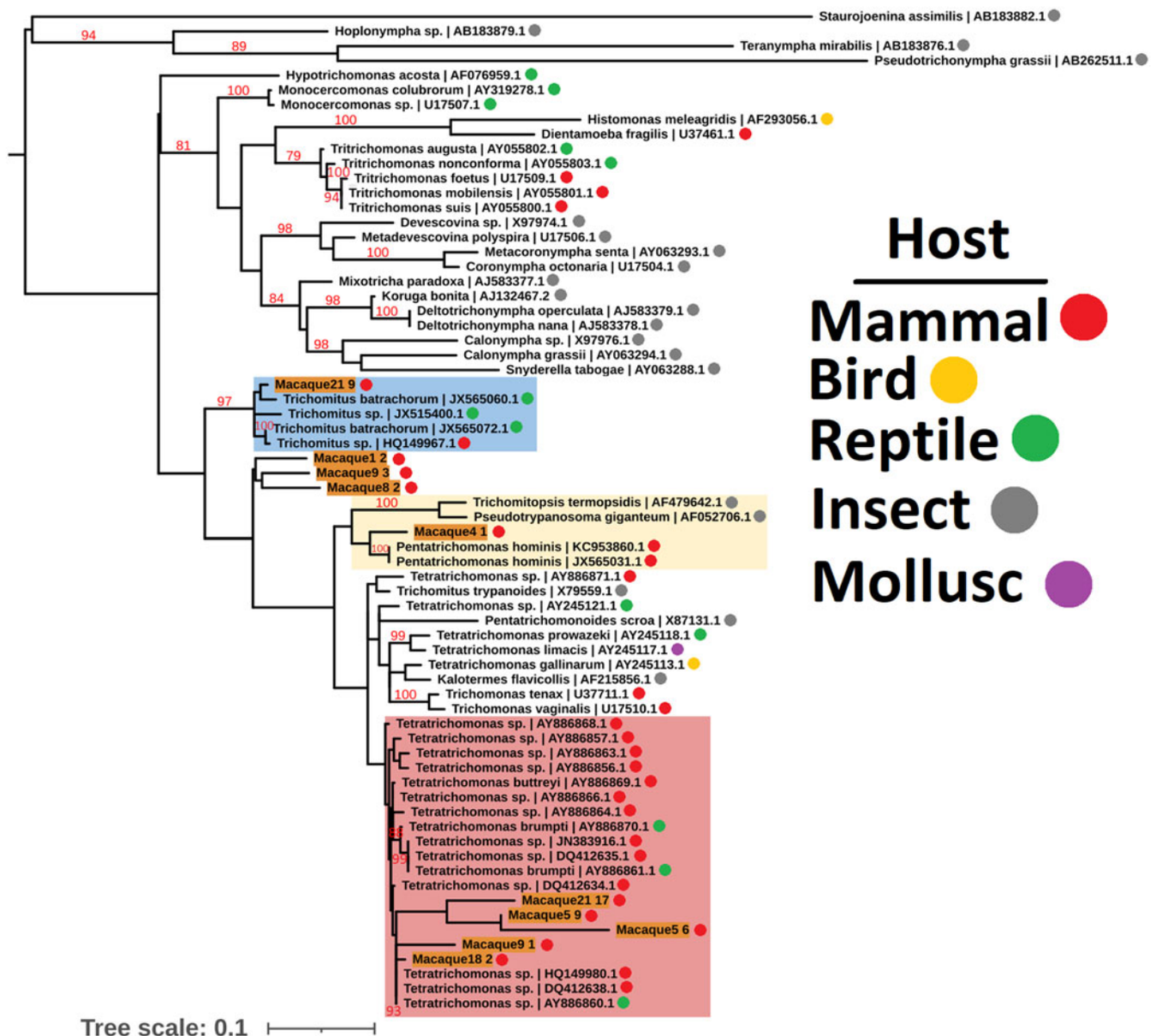


Fig. 1. Unrooted maximum likelihood phylogeny (GTR model with empirical base frequencies, invariable sites and the discrete gamma model) of Parasbasalia-like 18S rRNA sequences from the macaque fecal metatranscriptome, alongside a range of Parasbasalia species. Bootstrap values (1000 replicates) greater than 75% are shown on branches in red. Units for tree scale are inferred substitutions per base pair. Macaque-derived sequences (highlighted in orange) are named sequentially according to the animal from which they originated, e.g. Macaque1:1, Macaque1:2. Major lineages of macaque-derived *Tetratrichomonas*-like, *Pentatrichomonas*-like and *Trichomitus*-like sequences are highlighted in red, yellow and blue, respectively. Where available, Genbank accessions (Benson *et al.*, 2015) are shown at the ends of tip labels.

composition of the samples (Fig. 4). We reproduced a microbiota profile which was in agreement with the previous report (Westreich *et al.*, 2019); Bacteroidetes and Firmicutes were the most abundant phyla (Fig. 4B), the former largely dominated by the genus *Prevotella* (Fig. 4C) and the latter composed of a diverse range of genera (Fig. 4D). A large proportion of sequences could not be taxonomically classified at the phylum level (mean 66.7% of reads across all samples).

Principal component analysis (PCA) based on the microbial profile showed clear separation between the healthy and ICD groups. The macaques with ICD appeared to resolve into 3 sub-groups, potentially indicating distinct microbial communities (Fig. 5). An obvious association between PCA-based clustering and abundance of the parasitid genera of interest was not clear. However, we tentatively suggest a loose clustering of diseased animals somewhat consistent with *Tetratrichomonas* abundance (Fig. 5B–D). Of particular interest, a single healthy control

animal, macaque 17, clustered amongst the diseased animals. Macaque 17 showed the greatest abundance of total Parasbasalia, *Trichomitus* and *Pentatrichomonas* of all macaques, and the greatest *Tetratrichomonas* abundance amongst the macaques with ICD.

Our results suggested a possible relationship between parasite abundance and microbiota diversity. Amongst the macaques with ICD, there was a significant positive relationship between Parasbasalia abundance and microbial alpha diversity measures (number of observed taxa, Chao1 and Fisher diversity). There was also a significant negative relationship between Parasbasalia abundance and Simpson evenness, indicating a more non-uniform distribution of abundance amongst microbial taxa in animals with greater abundance of Parasbasalia (Fig. 6). However, this may be restricted to the ICD condition, as a significant relationship between Parasbasalia abundance and alpha diversity could not be demonstrated amongst the control macaques (*P* values derived from linear regression ranged from 0.21 to 0.27), although

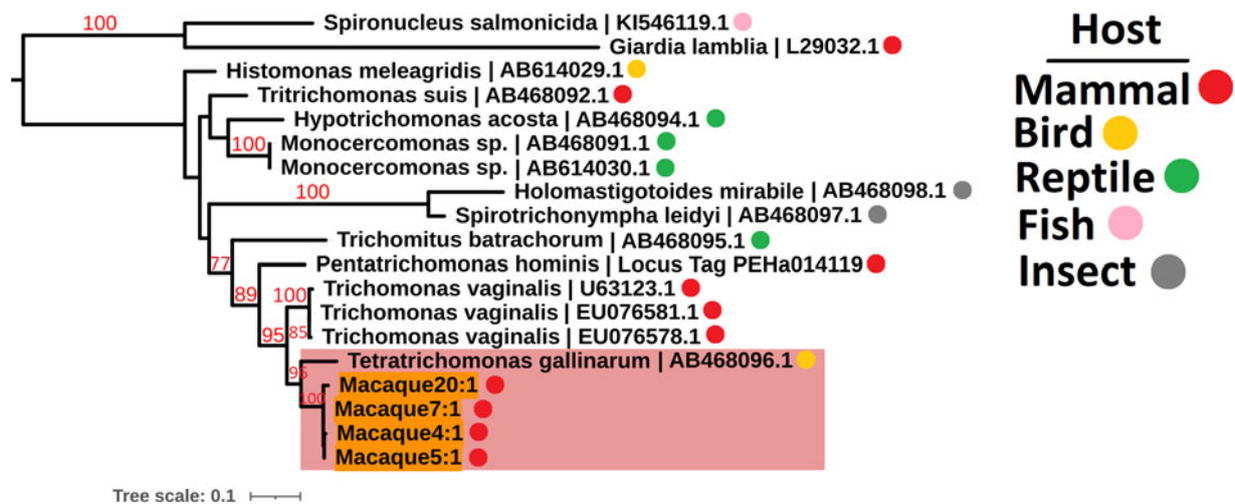


Fig. 2. Maximum likelihood phylogeny (TIM2e model with equal base frequencies and the discrete gamma model) of Parabasalia-like actin sequences from the macaque fecal metatranscriptome alongside a range of Parabasalia species. Phylogeny is rooted using sequences from *Giardia lamblia* (accession L29032.1) and *Spiroucleus salmonicida* (accession K1546119.1) as an outgroup (not shown). Bootstrap values (1000 replicates) greater than 75% are shown on branches in red. Units for tree scale are inferred substitutions per base pair. Macaque-derived sequences (highlighted in orange) are named sequentially according to the animal from which they originated, e.g. Macaque1:1, Macaque1:2. The major *Tetratrichomonas*-like lineage of macaque-derived sequences is highlighted in red. Coloured dots indicate animal host taxa. Where available, Genbank accessions (Benson *et al.*, 2015) are shown at the ends of tip labels.

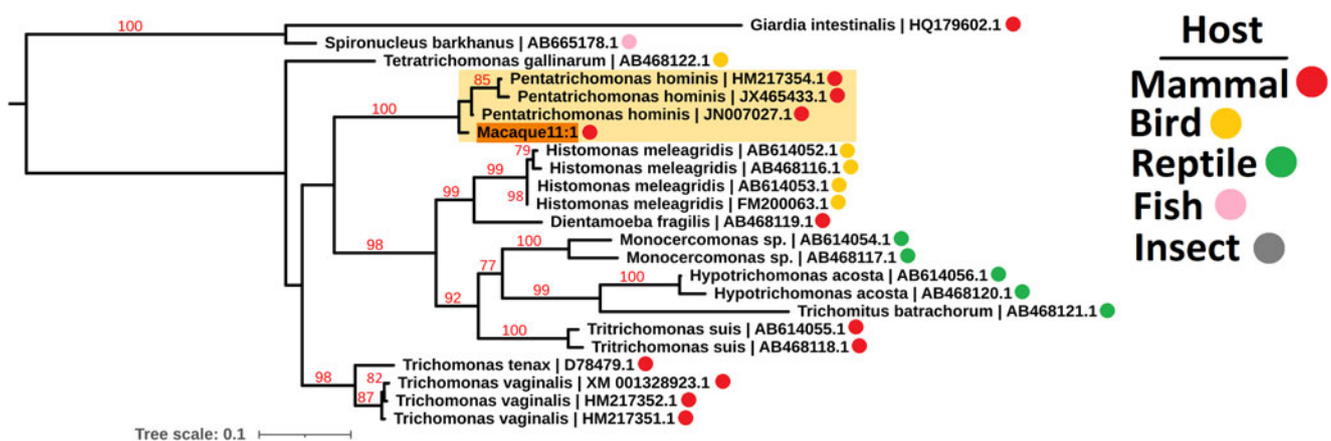


Fig. 3. Maximum likelihood phylogeny (TIM2 model allowing unequal base frequencies, with empirical base frequencies, invariable sites and the discrete gamma model) of Parabasalia-like EF-1 α sequences from the macaque fecal metatranscriptome, alongside a range of Parabasalia species. Phylogeny is rooted using sequences from *Giardia intestinalis* (accession HQ179602.1) and *Spiroucleus barkhanus* (accession AB665178.1) as an outgroup (not shown). Bootstrap values (1000 replicates) greater than 75% are shown on branches in red. Units for tree scale are inferred substitutions per base pair. Macaque-derived sequences (highlighted in orange) are named sequentially according to the animal from which they originated, e.g. Macaque1:1, Macaque1:2. The major *Pentatrichomonas*-like lineage of macaque-derived sequences is highlighted in yellow. Coloured dots indicate animal host taxa. Where available, Genbank accessions (Benson *et al.*, 2015) are shown at the ends of tip labels.

this is likely to have been influenced by the Parabasalia scarcity amongst the control animals (Fig. 5A). In addition, amongst the full cohort of macaques, using a combined linear model with disease state and Parabasalia abundance as predictors, only disease state showed a significant relationship with the same alpha diversity measures (P value derived from linear regression <0.0001), whereas Parabasalia abundance did not (P value ranged from 0.074 to 0.18). Intriguingly, ICD macaques with greater Parabasalia abundance appeared to more closely resemble control macaques in terms of alpha diversity.

To investigate specific interactions between Parabasalia and bacterial members of the microbiota, we performed an all vs all correlation analysis at the genus level. We focused on the macaques with ICD and included only the most abundant taxa (greater than 0.005% in at least 1 sample). Amongst 358 taxa, with a total of 64 261 possible interactions, our results indicated 11 606 significant abundance correlations between genera. Of the 3 parabasalid genera of interest, *Tetratrichomonas* showed the greatest

number of significant correlations with bacteria (110) followed by *Pentatrichomonas* (53), and *Trichomitius*, which showed far fewer significant correlations (17). *Tetratrichomonas* and *Pentatrichomonas* substantially overlapped in terms of bacterial genera showing significant positive and negative correlations, possibly indicating shared relationships with bacteria. In contrast, *Trichomitius* did not share common negative or positive relationships with any bacterial genera with either of the other parabasalid genera (Fig. 7A). Amongst the full complement of significant correlations, *Tetratrichomonas* stood out as participating in a large number of positive correlations. To investigate this further, we performed a network analysis by linking genera which shared a strong positive correlation. The genera were resolved into 29 connected components (Fig. 7). Of the 12 larger connected components (greater than 3 genera), 4 had a network clustering coefficient of greater than 0.5, indicating the majority of genera correlated with a given genus were also correlated with one another, suggesting well-supported and interdependent networks.

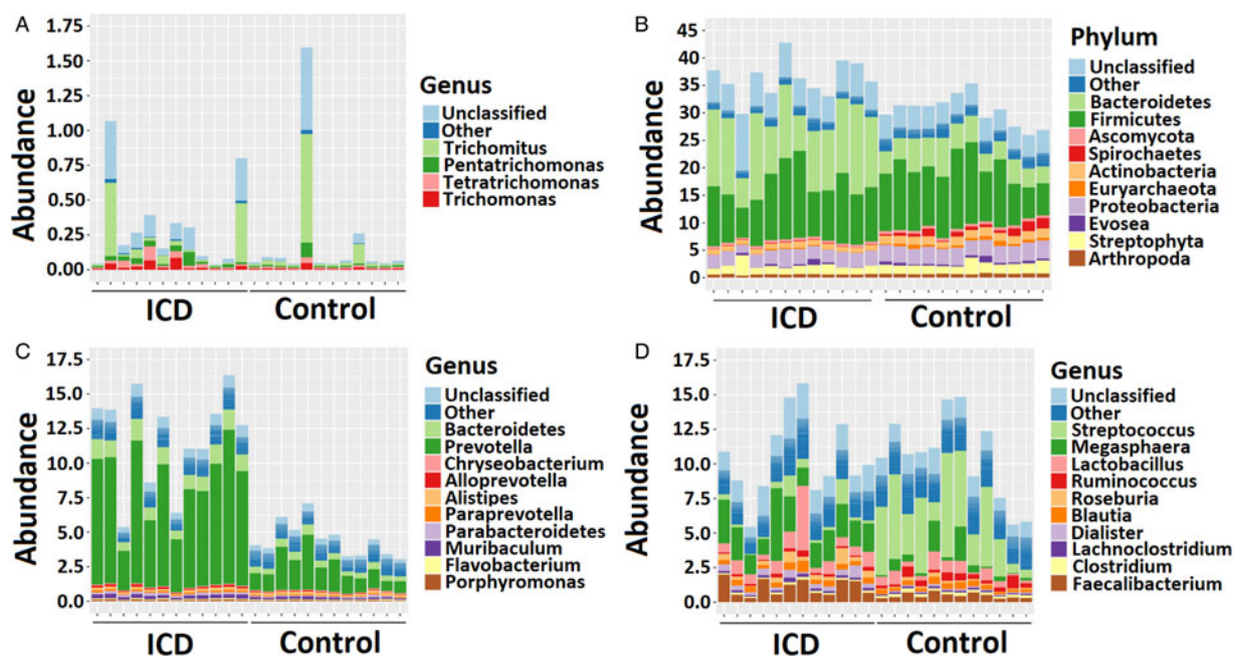


Fig. 4. Summary of microbial abundances amongst control macaques and this with idiopathic chronic diarrhoea. (A) Parasbasalia genera of interest, (B) Phyla excluding the host or Parasbasalia (C) Bacteroidetes genera and (D) Firmicutes genera. (B-D) show the abundance of the top 10 most abundant taxa (sum across all samples). Abundances are presented as a percentage of the total sequence library size. The 'other' category groups the rest of taxa not shown, and lines separate the subdivisions within these bars. The 'unclassified' category represents sequence reads which have been assigned to the relevant taxon of interest for the plot, but not to any specific phylum or genus. Samples are ordered 1–24 from left to right.

Table 1. Summary of selected Parasbasalia-like contigs of interest derived from the macaque fecal metatranscriptome

Contig name	Annotation	Pathway/function	<i>T. vaginalis</i> hit % identity (protein level)	Mean abundance (RPM)	Rank (ranked by abundance of all Parasbasalia-like contigs)
NODE_1480	Glutamate dehydrogenase	Amino acid catabolism	56.1	6031	1
NODE_1122	Phosphoenol pyruvate carboxykinase	Gluconeogenesis	55.9	4187	3
NODE_1126	Starch branching enzyme	Starch synthesis	48.0	3211	4
NODE_791	Fructose-1,6-bisphosphate aldolase	Glycolysis	65.7	3006	6
NODE_638 ^a	Glyceradehyde-3-phosphate dehydrogenase		85.3	2791	8
NODE_1712	Phosphoglucomutase/phosphomannomutase		42.4	2136	12
NODE_24	Pyruvate:ferredoxin oxidoreductase BII	Hydrogenosome metabolism	74.3	875	45
NODE_952	Ornithine carbamoyltransferase family protein	Arginine dihydrolase pathway	89.8	758	53
NODE_539	Malic enzyme	Hydrogenosome metabolism	83.6	289	200
NODE_134	Aldehyde oxidase and xanthine dehydrogenase	Purine catabolism	82.8	281	207
NODE_2461	Glucosamine-6-phosphate isomerase family protein	GlcNAc catabolism	49.8	265	222
NODE_2958	Transketolase family protein	Pentose phosphate pathway	45.9	242	239
NODE_418	Coronin	Phagocytosis	75.9	226	252
NODE_1833 ^a	Lysozyme	Bacterial cell wall degradation	70.6	134	434
NODE_2008 ^a	Lysozyme		64.8	126	462
NODE_589	Galactokinase family protein	Galactose metabolism	65.2	71	800
NODE_864	Succinyl-CoA ligase beta-chain	Hydrogenosome metabolism	78.1	60	946

^aAnnotation for the top *T. vaginalis* hit was uninformative, and so a lower hit within the top 10 hits was selected.

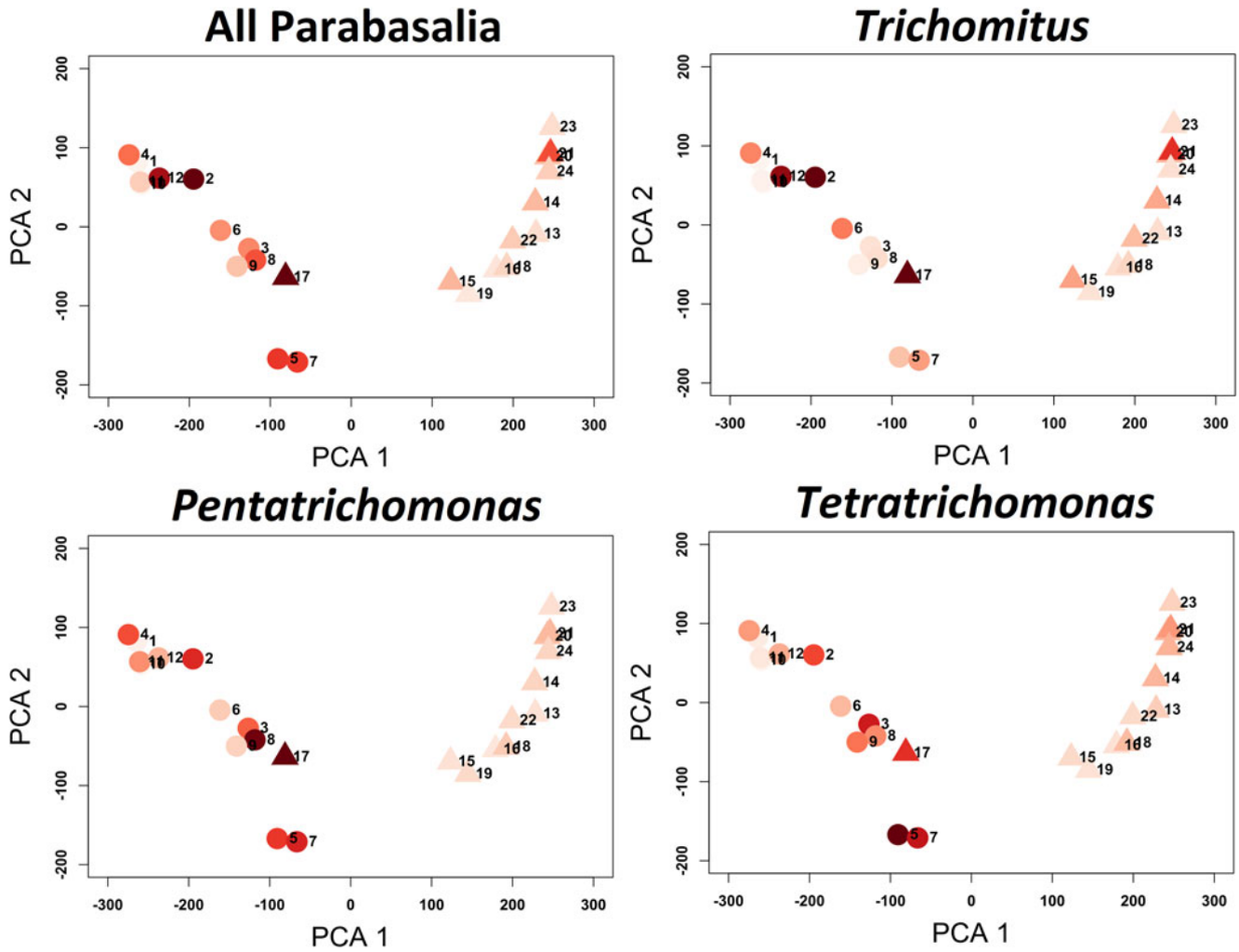


Fig. 5. Principal component analysis (PCA) plot for Aitchison distance based on non-parabaslid microbial abundances amongst macaque fecal samples. Points are shaded according to the centred log ratio normalized abundance values for all Parabasalia, *Trichomitus*, *Pentatrichomonas* and *Tetratrichomonas*, with darker shades indicating greater abundance. Triangle and circular points indicate healthy and diseased animals, respectively.

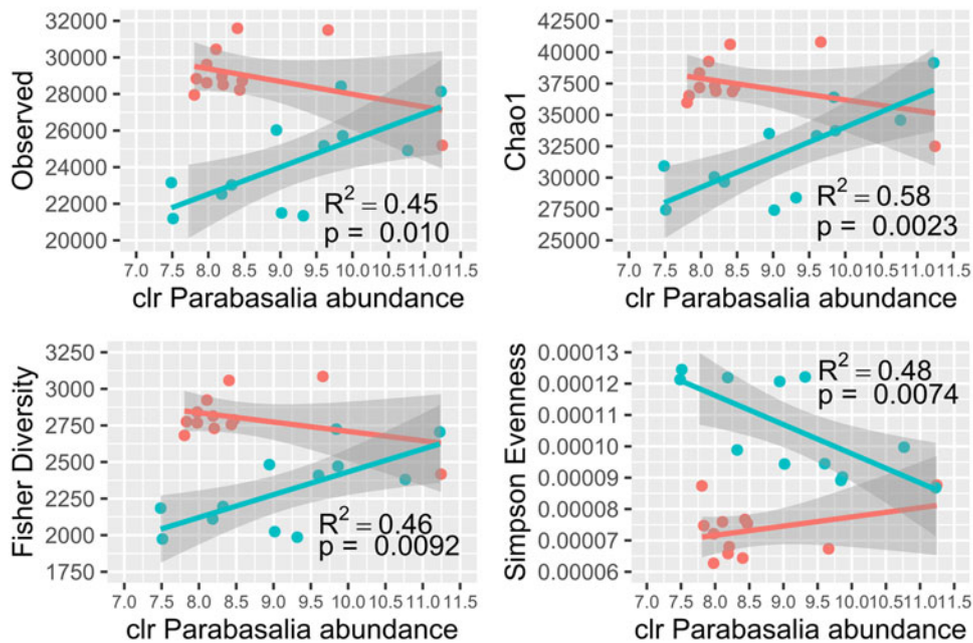


Fig. 6. Relationship between Parabasalia abundance (normalized by centred log-ratio; clr) and microbial diversity metrics. Observed refers to the total number of observed taxa. Macaques with idiopathic chronic diarrhoea (ICD) and healthy controls are shown in pink and turquoise, respectively. Lines indicate separate linear regressions fitted for the ICD and control groups, and the shaded areas indicate 95% confidence intervals. The significant linear regression *P* values (<0.05) and corresponding *R*² values derived from the ICD sample are indicated next to the corresponding line. The linear regression results for the control macaques were not significant (*P* value >0.05).

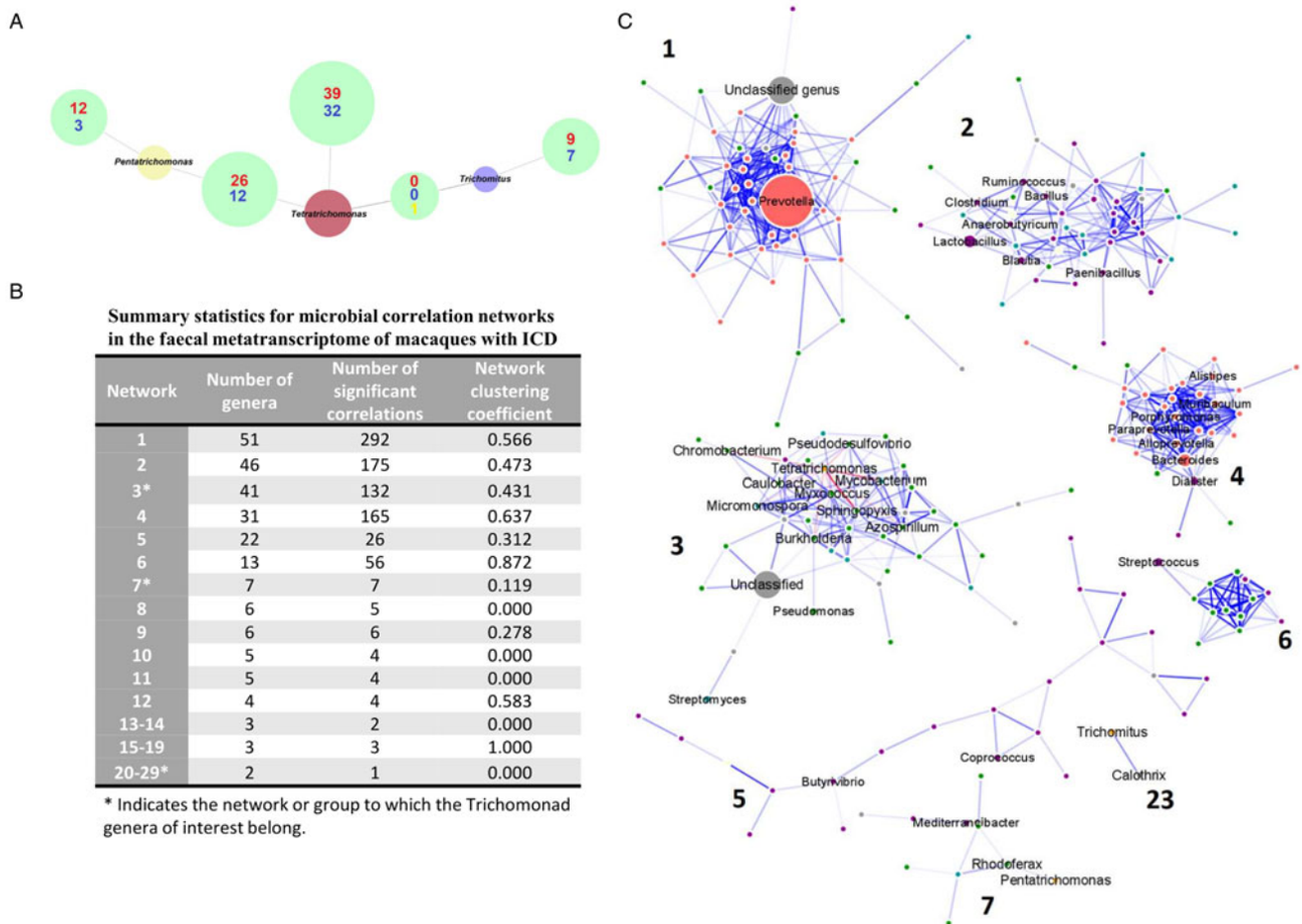


Fig. 7. Correlation analysis of microbial abundance amongst macaques with idiopathic chronic diarrhoea. (A) DiVenn (Sun *et al.*, 2019) figure showing overlap of positive (red) and negative (blue) correlations with bacteria amongst the parabasalid genera of interest. Yellow colour indicates shared correlations for which the direction of correlation differs between the groups. (B) Summary table for the microbial correlation networks. (C) Correlation network of the most abundant microbial genera (greater than 0.005% in at least 1 sample). Edges link nodes (genera) for which abundance was strongly correlated (sharing a significant correlation coefficient greater than 0.8). Node size is scaled to percentage abundance (ignoring the abundance of the 'unclassified' group), and labels for genera with less than 0.1% abundance are omitted (except for Parabasalia nodes and their immediate neighbours). Nodes are coloured according to phylum (Actinobacteria; turquoise, Bacteroides; pink, Firmicutes; purple, Proteobacteria; green and Parabasalia; orange) and edge transparency is scaled to the magnitude of the correlation coefficient. Edges linking to *Tetratrachomonas* are highlighted in red. Connected components are numbered sequentially by decreasing number of nodes and connected components with fewer than 7 nodes are not shown, except those that contain Parabasalia (21 connected components are not shown).

Seventeen connected components had fewer than 4 genera, which in total accounted for 41 genera. This overall suggests a complex mixture of interdependent and independent bacterial genera. *Tetratrachomonas* inhabited the largest connected component of the 3 parabasalid genera of interest (network 3), with a moderate network clustering coefficient of 0.431, indicating a greatest potential interdependence with bacteria. Within network 3, the closeness centrality of *Tetratrachomonas* was 0.465, the 10th highest in the 41-node network, suggesting a relatively central hub-like position in comparison to most bacterial nodes. In contrast, *Pentatrachomonas* (network 7) and *Trichomitus* (network 23) inhabited smaller and more sparsely interconnected components, with 7 and 2 genera, respectively.

Notably, almost all bacterial genera which showed a significant negative correlation with *Tetratrachomonas* and *Pentatrachomonas* were Gram negative, but this pattern did not extend to *Trichomitus* (Table 2). *Pentatrachomonas* in particular showed a negative correlation with many bacterial genera reported to contain mucosal inhabitants which are opportunistic pathogens in various host species, including *Gemella* (Nazik *et al.*, 2018), *Moraxella* (Goldstein *et al.*, 2009), *Mannheimia* (Clawson and Murray, 2014) and *Aggregatibacter* (Karched *et al.*, 2012). Importantly, *Tetratrachomonas* showed a strong negative correlation with *Prevotella*, which was the most abundant bacterial

genus across all samples. The full list of significant correlations for *Tetratrachomonas*, *Pentatrachomonas* and *Trichomitus* amongst the macaques with ICD is shown in Table S4.

The majority of relationships identified amongst the macaques with ICD were not consistent amongst the control group. Only 1188 out of 11 606 total significant relationships for the macaques with ICD were homodirectionally concordant amongst the control animals, including 2 out of 110 relationships for *Tetratrachomonas* (*Bradyrhizobium* and *Pentatrachomonas*), 5 out of 53 for *Pentatrachomonas* (*Acinetobacter*, *Janthinobacterium*, *Mesorhizobium*, *Streptomyces* and *Tetratrachomonas*) and 2 out of 17 for *Trichomitus* (*Calothrix* and *Colwellia*). This may indicate that the microbial community structure and interdependence was dramatically different between the ICD and control conditions.

We performed a differential abundance analysis between the ICD and control groups in order to investigate any potential impact of the parabasalids on disease aetiology. Interestingly, differential abundance analysis suggested a moderate significantly higher abundance of *Tetratrachomonas* and *Pentatrachomonas* (\log_2 fold differences were 1.62 and 1.80, respectively; adjusted *P* value <0.001), but not *Trichomitus*, amongst the macaques with ICD compared with the healthy controls. The original authors ruled out several known common GI pathogens (Bacteria: *Campylobacter jejuni*, *Salmonella*, *Shigella flexneri* and

Table 2. Top significant negative correlations between parabasalid and bacterial genera by sparCC analysis

Parasite genus	Bacterial genus	Bacterial phylum	Bacterial Gram stain	Correlation coefficient	P value
<i>Tetratrichomonas</i>	<i>Leeuwenhoekella</i>	Bacteroidetes	–	–0.824	0.00
	<i>Algibacter</i>	Bacteroidetes	–	–0.800	0.00
	<i>Desulfocapsa</i>	Proteobacteria	–	–0.793	0.00
	<i>Veillonella</i>	Firmicutes	–	–0.790	0.01
	<i>Desulfitobacterium</i>	Firmicutes	+	–0.784	0.00
	<i>Alteromonas</i>	Proteobacteria	–	–0.765	0.00
	<i>Emticicia</i>	Bacteroidetes	–	–0.751	0.00
	<i>Xenorhabdus</i>	Proteobacteria	–	–0.740	0.01
	<i>Prevotella</i>	Bacteroidetes	–	–0.721	0.00
<i>Psychrobacter</i>	Proteobacteria	–	–0.718	0.00	
<i>Pentatrichomonas</i>	<i>Basfia</i>	Proteobacteria	–	–0.735	0.00
	<i>Gemella</i>	Firmicutes	+/-	–0.675	0.00
	<i>Moraxella</i>	Proteobacteria	–	–0.639	0.03
	<i>Avibacterium</i>	Proteobacteria	–	–0.638	0.00
	<i>Acidaminococcus</i>	Firmicutes	–	–0.635	0.05
	<i>Roseburia</i>	Firmicutes	+	–0.630	0.03
	<i>Aggregatibacter</i>	Proteobacteria	–	–0.617	0.04
	<i>Mannheimia</i>	Proteobacteria	–	–0.614	0.03
	<i>Chania</i>	Proteobacteria	–	–0.614	0.04
<i>Thalassospira</i>	Proteobacteria	–	–0.610	0.05	
<i>Trichomitrus</i>	<i>Dickeya</i>	Proteobacteria	–	–0.696	0.00
	<i>Kocuria</i>	Actinobacteria	+/-	–0.689	0.02
	<i>Amycolatopsis</i>	Actinobacteria	+	–0.633	0.05
	<i>Aerococcus</i>	Firmicutes	+	–0.614	0.03
	<i>Prochlorococcus</i>	Cyanobacteria	–	–0.545	0.04
	<i>Phascolarctobacterium</i>	Firmicutes	–	–0.527	0.05
	<i>Pseudonocardia</i>	Actinobacteria	+	–0.525	0.05
	<i>Desulfosporosinus</i>	Firmicutes	+	–0.508	0.05

Yersinia enterocolitica and Parasites: *Cryptosporidium*, *Giardia*) as the cause of ICD by culture and microscopy-based methods. To complement this, we searched the dataset for potentially pathogenic viral lineage amongst the taxonomic profile. We focused on a selection of 29 potential primate-infecting eukaryotic viruses which we identified by the literature search (Oberste *et al.*, 2007; Handley *et al.*, 2012; Campanini *et al.*, 2013; Janowski *et al.*, 2017; Gao *et al.*, 2018; Zhang *et al.*, 2019b; Smura *et al.*, 2020; Kang *et al.*, 2021) (Fig. S7). Abundance of these viruses was low; total abundance of all 29 viruses was less than 1.6% for all animals, and the highest individual viral abundance was for Simian enterovirus 19, at 0.52%. Notably, we did not identify any significant difference in abundance for any of the viruses comparing between diseased and control animals (Mann–Whitney U test, *P* value >0.05).

In order to further query the potential influence of parabasalids on the microbiota, we examined the relationship between the HUMAnN2-annotated functional microbial gene expression and parabasalid abundance. The mean-variance relationship of the MetaCyc pathway quantification data is shown in Figure S8. A PCA of the ICD samples based on microbial pathway abundance showed tentative segregation of samples with low and high *Tetratrichomonas* abundance (Fig. S9). We identified a significant

negative relationship between the abundances of 12 MetaCyc pathways and that of *Tetratrichomonas* amongst the macaques with ICD (Table 3), although the magnitude of the log₂ fold changes were relatively small. The strongest relationship was detected for the Superpathway of *N*-acetylglucosamine, *N*-acetylmannosamine and *N*-acetylneuraminate degradation. The majority of functional sequences could not be attributed to a particular microbial species. However, many functions corresponded to likely constitutive bacterial functions such as peptidoglycan synthesis, potentially indicating a negative relationship with bacteria which could not be classified. The analysis did not identify any significant positive relationships between *Tetratrichomonas* and MetaCyc pathway abundances, and no significant relationships were found for the abundances of both *Pentatrichomonas* and *Trichomitrus* amongst the macaques with ICD. The significant negative relationships identified for *Tetratrichomonas* could not be detected amongst the control animals.

Discussion

We aimed to investigate potential relationships between the host, Trichomonads and the mucosal microbiota during health and

Table 3. Microbial pathways showing a significant relationship with *Tetratrichomonas* abundance

MetaCyc ID	Description	Log ₂ fold change ^a	Average abundance (CPM)	Species stratification	Stratified abundance	Adjusted <i>P</i> value
GLCMANNANAUT-PWY	Superpathway of <i>N</i> -acetylglucosamine, <i>N</i> -acetylmannosamine and <i>N</i> -acetylneuraminate degradation	-1.28	5.19	NA		0.04
P441-PWY	Superpathway of <i>N</i> -acetylneuraminate degradation	-1.12	12.5	NA		0.03
PWY-6507	4-Deoxy-L-threo-hex-4-enopyranuronate degradation	-0.90	16.6	NA		0.03
PWY-7242	D-fructuronate degradation	-0.76	25.4	NA		0.04
GALACTUROCAT-PWY	D-galacturonate degradation I	-0.73	21.9	NA		0.04
GALACT-GLUCUROCAT-PWY	Superpathway of hexuronide and hexuronate degradation	-0.65	25.7	NA		0.04
GLUCUROCAT-PWY	Superpathway of β -D-glucuronosides degradation	-0.61	27.1	NA		0.04
PWY-5686	UMP biosynthesis	-0.44	38.7	<i>Treponema succinifaciens</i>	0.014	0.04
				<i>Bifidobacterium angulatum</i>	0.018	
				<i>Coprococcus catus</i>	0.050	
				<i>Eubacterium bifforme</i>	0.061	
PWY-6386	UDP- <i>N</i> -acetylmuramoyl-pentapeptide biosynthesis II (lysine-containing)	-0.38	41.8	<i>Bifidobacterium angulatum</i>	0.0054	0.05
				<i>Treponema succinifaciens</i>	0.023	
				<i>Phascolarctobacterium succinatutens</i>	0.051	
				<i>Eubacterium bifforme</i>	0.070	
PWY-6385	Peptidoglycan biosynthesis III (mycobacteria)	-0.37	46.9	<i>Treponema succinifaciens</i>	0.024	0.04
				<i>Phascolarctobacterium succinatutens</i>	0.130	
PWY-6387	UDP- <i>N</i> -acetylmuramoyl-pentapeptide biosynthesis I (meso-diaminopimelate containing)	-0.36	53.3	<i>Bifidobacterium angulatum</i>	0.0050	0.04
				<i>Treponema succinifaciens</i>	0.021	
				<i>Eubacterium bifforme</i>	0.053	
				<i>Phascolarctobacterium succinatutens</i>	0.130	
PEPTIDOGLYCANSYN-PWY	Peptidoglycan biosynthesis I (meso-diaminopimelate containing)	-0.35	55.9	<i>Bifidobacterium angulatum</i>	0.0059	0.04
				<i>Treponema succinifaciens</i>	0.024	
				<i>Eubacterium bifforme</i>	0.051	
				<i>Phascolarctobacterium succinatutens</i>	0.13	

^aThe coefficient resulting from the linear regression fit between pathway and *Tetratrichomonas* abundance, interpreted as the log₂ fold change in pathway abundance per unit of *Tetratrichomonas* abundance.

disease through re-analysis of an existing metatranscriptomics dataset derived from macaque fecal samples. We identified a novel combination of *Pentatrachomonas*, *Tetratrachomonas* and *Trichomitus* parasites in the macaque gut. *P. hominis* was previously reported in laboratory macaques with ICD (Laing *et al.*, 2018), and simian immunodeficiency virus (Zaragoza *et al.*, 2011). However, to our knowledge, this is the first description of *Trichomitus* and *Tetratrachomonas* spp. colonizing the macaque gut. *Tetratrachomonas* spp. (Cepicka *et al.*, 2006) and *Pentatrachomonas* spp. (Li *et al.*, 2016; Bastos *et al.*, 2018; Kim *et al.*, 2020) are common GI-inhabitants of mammals. *Trichomitus* spp. typically infect reptiles and amphibians (Viscogliosi and Müller, 1998; Delgado-Viscogliosi *et al.*, 2000), although mammalian infection has been reported (Dimasuy *et al.*, 2013). Previous reports of macaque-infecting Trichomonads have also included gastric-localized *Tritrachomonas* spp. (Kondova *et al.*, 2005). It is possible that the intestinal Trichomonad infections represent an artefact resulting from laboratory husbandry, as reports are scarce, and studies on wild macaques did not identify intestinal Trichomonads (Adhikari and Dhakal, 2018; Zhang *et al.*, 2019a). As the animals were at times housed together (Westreich *et al.*, 2019), the possibility of transmission of parasites between the laboratory animals seems likely. Any potential relationship between Trichomonads and diseases such as ICD in macaques has not been reported.

Our results suggested commonality in the expressed functional genes across the Trichomonads. We observed similar energy generation mechanisms for the macaque-infecting parabasalids as have been previously reported for *T. vaginalis*, demonstrated by a high abundance of transcripts associated with glycolysis, hydrogenosomal metabolism, amino acid catabolism (including the ADH pathway) and glycogen storage and processing (Müller, 1990; Kulda, 1999; Westrop *et al.*, 2017). We also detected BspA expression potentially attributed to the macaque-infecting parabasalids. BspAs are of interest because proteins of this family have demonstrated roles in host adhesion by bacteria as well as adhesion between bacterial cells (Sharma, 2010). Importantly, *T. vaginalis* BspAs have been implicated in host adhesion *in vitro* (Handrich *et al.*, 2019). In addition, *T. vaginalis* BspAs are differentially expressed in response to *Mycoplasma* symbionts, suggesting a potential role in parasite–bacteria interactions in modulating parasite binding to host cells (Margarita *et al.*, 2022) and possibly binding to members of the microbiota too.

Our results also indicated a potential influential interaction between Trichomonads and microbial diversity in the macaque gut, as has been reported for other hosts and mucosa (El Sayed Zaki *et al.*, 2010; Ji *et al.*, 2020; Wei *et al.*, 2020; Li *et al.*, 2021). Probable coinfection with at least 3 Trichomonad genera complicated accurate abundance estimation due to the portion of sequences which could not be assigned to a specific genus (Watts *et al.*, 2019). Thus, dissection of the relative effects for individual parasites was recalcitrant. This highlights a limitation of observational studies (Cani, 2018). Despite the greater abundance of *Trichomitus*, our results suggested *Tetratrachomonas* had the greatest abundance correlation with differences in the microbiota. *Tetratrachomonas* participated in the greatest number of significant abundance correlations, and was a central node within a densely interconnected microbial positive correlation network. Correlation networks have been effectively utilized to identify keystone species within microbial communities with biological significance (Duran-Pinedo *et al.*, 2011). An overlap in specific relationships between differing Trichomonad spp. and bacteria may be suggested by shared correlations with bacterial abundance between *Pentatrachomonas* and *Tetratrachomonas*. In addition, of particular interest, a positive abundance correlation with *Prevotella*, which we observed for *Tetratrachomonas* in the macaque gut, has been described for *T. vaginalis* in the human

UGT (Martin *et al.*, 2013; Jarrett *et al.*, 2019). Conserved interactions may result from biochemical features shared amongst the bacteria, supported by our observation that many of the bacteria negatively correlated with *Tetratrachomonas* and *Pentatrachomonas* were Gram negative. This is notable because other Trichomonads such as *Dientamoeba fragilis* (Chan *et al.*, 1993) depend on Gram-negative bacteria for *in vitro* growth. Microbial interactions identified in this study varied hugely between the diseased and healthy conditions, similarly to previous results from the healthy and diseased human oral microbiota (Duran-Pinedo *et al.*, 2011). This could suggest wholesale changes in community structure between conditions, but may also reflect unreliability in quantifying microbial abundance correlation (Weiss *et al.*, 2016; Matchado *et al.*, 2021). Although only a single sample, 1 control macaque both resembled the ICD macaques in terms of microbial profile, and showed the greatest abundance of Trichomonads, consistent with a potential parasite–bacterial interaction.

Previous studies have suggested the interaction between Trichomonads and the vaginal microbiota is bidirectional. The microbial profile can influence the ability of Trichomonads to colonize the mucosa (Rathod *et al.*, 2011), and the presence of Trichomonad can perturb the microbial profile (Fichorova *et al.*, 2013; Wei *et al.*, 2020). However, the direction of influence between Trichomonads and the microbiota in the macaque gut could not be determined in the absence longitudinal data. In the macaque, Parabasalia expression of potential microbial-targeting genes such as lysozyme could indicate predation, as has been demonstrated for *T. vaginalis* (Pinheiro *et al.*, 2018). This could provide a mechanistic basis for negative correlations between parasite and microbial abundance. However, we could only reliably attribute lysozyme-encoding transcripts to *Trichomitus*, whereas *Tetratrachomonas* was the only Trichomonad genus correlated with bacterial functional expression. Negative correlation of *Tetratrachomonas* abundance with bacterial degradation pathways for monosaccharides such as GlcNAc and Sia5NAc, potentially derived from mucin glycoproteins (Yurewicz *et al.*, 1987) or microbial cells (Pinheiro *et al.*, 2018), could indicate nutritional competition. This is supported by the detected expression of GlcNAc-targeting glycosyl hydrolases and potentially associated catabolic enzymes by the Trichomonads.

The absence of several known GI bacteria pathogens and microbial parasites was confirmed by culture and microscopy-based methods, and thus may be excluded as causative agents of ICD in the macaques. We performed an additional search for potentially pathogenic viruses amongst the datasets. However, we did not identify any clear differences for any putative host-infecting virus when comparing between diseased and control animals, suggesting viral infection may not be the primary cause of ICD. A greater abundance of reads classified as originating from the *Campylobacter* genus amongst the animals with ICD was originally reported (Westreich *et al.*, 2019), and so the potential presence of other pathogens in this genus cannot be ruled out. Our results did not establish a causal link between Trichomonads and ICD in macaques. The higher abundance of *Pentatrachomonas* and *Tetratrachomonas* could indicate a causal role in disease. High *P. hominis* abundance in macaques with ICD was previously reported, but not causally linked to disease (Laing *et al.*, 2018). However, abundance of these parasites appeared to promote a more diverse (control-like) microbiota. Trichomonads were positively correlated with microbial diversity, which was also higher in healthy animals, and has been considered characteristic of healthy human gut (Malard *et al.*, 2021). This contrasts with previous work which suggested that the presence of *T. gallinae* and *Tritrichomonas musculus* decreases GI microbial diversity (Ji *et al.*, 2020; Wei *et al.*, 2020). Notably, we did not detect any correlation between Trichomonads and the

abundance of bacterial genes underlying mucin degradation or fucose utilization, the previously proposed determinants of macaque ICD (Westreich *et al.*, 2019). It is feasible that the ICD state provides a beneficial environment for Trichomonad colonization, within which the parasites exert a disruptive influence. This is supported by the observation that Trichomonad–microbial interactions appeared to be highly dependent on disease state.

Our results revealed a relatively low parasite abundance in the macaque fecal samples, highlighting the need for greater sequencing depth or selective target enrichment (Gaudin and Desnues, 2018) to quantitatively study the parasite transcriptome *in vivo*. Reference sequences from closely related parasite strains would also have greatly facilitated analysis (Breitwieser *et al.*, 2019). As is typical for a diverse *in vivo* metatranscriptome (Li *et al.*, 2019), a large proportion of sequences could not be assigned to a specific phylum.

In summary, these metatranscriptomics analyses of Trichomonads in the macaque gut have provided the first *in vivo* insight into Trichomonad mucosal colonization, which validates numerous *in vitro* studies (Müller, 1990; Kulda, 1999; Westrop *et al.*, 2017; Handrich *et al.*, 2019). Our findings support previous reports of Trichomonad–microbiota interactions (Ji *et al.*, 2020; Wei *et al.*, 2020; Bierlein *et al.*, 2021), and demonstrate that such interactions vary between parasite species and are highly context-dependent. Longitudinal studies, or those involving experimental Trichomonad infection, could be used to investigate causality and underlying mechanisms in the parasite–microbiota–disease interrelationship.

Supplementary material. The supplementary material for this article can be found at <https://doi.org/10.1017/S0031182022001688>

Data availability. This work presents a re-analysis of metatranscriptomics data generated by Westreich *et al.* (2019), DOI: <https://doi.org/10.1186/s40168-019-0664-z>. Original data are available from the NCBI SRA database (Leinonen *et al.*, 2011) under accession numbers SRX3517701–SRX3517724. All the supplementary files containing *de novo* assemblies of the original sequence data and alignments used for phylogenetics are available *via* figshare (<https://figshare.com/s/5d6f50cb71ed2ffc82fb>).

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Conflict of interest. The authors declare that there are no conflicts of interest.

Ethical standards. No experimental data collection requiring ethical approval was performed during the course of this work.

References

- Adhikari PP and Dhakal P (2018) Prevalence of gastro-intestinal parasites of rhesus macaque (*Macaca mulatta* Zimmermann, 1780) and hanuman langur (*Semnopithecus entellus* Dufresne, 1797) in Devghat, Chitwan, Nepal. *Journal of Institute of Science and Technology* **22**, 7.
- Altschul SF, Gish W, Miller W, Myers EW and Lipman DJ (1990) Basic local alignment search tool. *Journal of Molecular Biology* **215**, 403–410.
- Andrews S (2018) FastQC. Vol. 2018 Babraham bioinformatics. <https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>
- Arroyo R, Cárdenas-Guerra RE, Figueroa-Angulo EE, Puente-Rivera J, Zamudio-Prieto O and Ortega-López J (2015) *Trichomonas vaginalis* cysteine proteinases: iron response in gene expression and proteolytic activity. *Biomedical Research International* **2015**, 946787.
- Bastos BF, Brener B, de Figueiredo MA, Leles D and Mendes-de-Almeida F (2018) *Pentatrichomonas hominis* infection in two domestic cats with chronic diarrhoea. *Journal of Feline Medicine and Surgery Open Reports* **4**, 2055116918774959.
- Beghini F, McIver LJ, Blanco-Míguez A, Dubois L, Asnicar F, Maharjan S, Mailyan A, Manghi P, Scholz M, Thomas AM, Valles-Colomer M, Weingart G, Zhang Y, Zolfo M, Huttenhower C, Franzosa EA and Segata N (2021) Integrating taxonomic, functional, and strain-level profiling of diverse microbial communities with bioBakery 3. *Elife* **10**, e65088. doi: 10.7554/eLife.65088.
- Bello-Ortí B, Howell KJ, Tucker AW, Maskell DJ and Aragon V (2015) Metatranscriptomics reveals metabolic adaptation and induction of virulence factors by *Haemophilus parasuis* during lung infection. *Veterinary Research* **46**, 102.
- Benson DA, Clark K, Karsch-Mizrachi I, Lipman DJ, Ostell J and Sayers EW (2015) GenBank. *Nucleic Acids Research* **43**, D30–D35.
- Bierlein M, Hedgespeth BA, Azcarate-Peril MA, Stauffer SH and Gookin JL (2021) Dysbiosis of fecal microbiota in cats with naturally occurring and experimentally induced *Tritrichomonas foetus* infection. *PLoS One* **16**, e0246957.
- Blondel VD, Guillaume JL, Lambiotte R and Lefebvre E (2008) Fast unfolding of communities in large networks. *Journal of Statistical Mechanics – Theory and Experiment* **12**, P10008. doi: 10.1088/1742-5468/2008/10/p10008.
- Breitwieser FP, Lu J and Salzberg SL (2019) A review of methods and databases for metagenomic classification and assembly. *Briefings in Bioinformatics* **20**, 1125–1136.
- Bricheux G, Coffe G, Bayle D and Brugerolle G (2000) Characterization, cloning and immunolocalization of a coronin homologue in *Trichomonas vaginalis*. *European Journal of Cell Biology* **79**, 413–422.
- Campanini G, Rovida F, Meloni F, Cascina A, Ciccocioppo R, Piralla A and Baldanti F (2013) Persistent human cosavirus infection in lung transplant recipient, Italy. *Emerging Infectious Diseases* **19**, 1667–1669.
- Caní PD (2018) Human gut microbiome: hopes, threats and promises. *Gut* **67**, 1716–1725.
- Capella-Gutiérrez S, Silla-Martínez JM and Gabaldón T (2009) trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. *Bioinformatics (Oxford, England)* **25**, 1972–1973.
- Carlton JM, Hirt RP, Silva JC, Delcher AL, Schatz M, Zhao Q, Wortman JR, Bidwell SL, Alsmark UC, Besteiro S, Sicheritz-Ponten T, Noel CJ, Dacks JB, Foster PG, Simillion C, Van de Peer Y, Miranda-Saavedra D, Barton GJ, Westrop GD, Müller S, Dessi D, Fiori PL, Ren Q, Paulsen I, Zhang H, Bastida-Corcuera FD, Simoes-Barbosa A, Brown MT, Hayes RD, Mukherjee M, Okumura CY, Schneider R, Smith AJ, Vanacova S, Villalvazo M, Haas BJ, Perlea M, Feldblyum TV, Utterback TR, Shu CL, Osoegawa K, de Jong PJ, Hrdy I, Horvathova L, Zubacova Z, Dolezal P, Malik SB, Logsdon JM, Henze K, Gupta A, Wang CC, Dunne RL, Upcroft JA, Upcroft P, White O, Salzberg SL, Tang P, Chiu CH, Lee YS, Embley TM, Coombs GH, Mottram JC, Tachezy J, Fraser-Liggett CM and Johnson PJ (2007) Draft genome sequence of the sexually transmitted pathogen *Trichomonas vaginalis*. *Science (New York, N.Y.)* **315**, 207–212.
- Caspi R, Billington R, Ferrer L, Foerster H, Fulcher CA, Keseler IM, Kothari A, Krummenacker M, Latendresse M, Mueller LA, Ong Q, Paley S, Subhraveti P, Weaver DS and Karp PD (2016) The MetaCyc database of metabolic pathways and enzymes and the BioCyc collection of pathway/genome databases. *Nucleic Acids Research* **44**, D471–D480.
- Cepicka I, Hampl V, Kulda J and Flegr J (2006) New evolutionary lineages, unexpected diversity, and host specificity in the parabasalid genus *Tetratrichomonas*. *Molecular Phylogenetics and Evolution* **39**, 542–551.
- Chan FT, Guan MX and Mackenzie AM (1993) Application of indirect immunofluorescence to detection of *Dientamoeba fragilis* trophozoites in fecal specimens. *Journal of Clinical Microbiology* **31**, 1710–1714.
- Clawson ML and Murray RW (2014) Pathogen variation across time and space: sequencing to characterize *Mannheimia haemolytica* diversity. *Animal Health Research Reviews* **15**, 169–171.

- Costello EK, Sun CL, Carlisle EM, Morowitz MJ, Banfield JF and Relman DA (2017) Candidatus *Mycoplasma girerdii* replicates, diversifies, and co-occurs with *Trichomonas vaginalis* in the oral cavity of a premature infant. *Scientific Reports* 7, 3764.
- Csardi G and Nepusz T (2006) The igraph software package for complex network research. Vol. Complex Systems pp. 1695. InterJournal.
- Delgado-Viscogliosi P, Viscogliosi E, Gerbod D, Kulda J, Sogin ML and Edgcomb VP (2000) Molecular phylogeny of parabasalids based on small subunit rRNA sequences, with emphasis on the Trichomonadinae subfamily. *Journal of Eukaryotic Microbiology* 47, 70–75.
- de Miguel N, Lustig G, Twu O, Chattopadhyay A, Wohlschlegel JA and Johnson PJ (2010) Proteome analysis of the surface of *Trichomonas vaginalis* reveals novel proteins and strain-dependent differential expression. *Molecular and Cellular Proteomics* 9, 1554–1566.
- Dessi D, Margarita V, Cocco AR, Marongiu A, Fiori PL and Rappelli P (2019) *Trichomonas vaginalis* and *Mycoplasma hominis*: new tales of two old friends. *Parasitology* 146, 1150–1155.
- Dimasuy KG, Lavilla OJ and Rivera WL (2013) New hosts of *Simplicimonas similis* and *Trichomitus batrachorum* identified by 18S ribosomal RNA gene sequences. *Journal of Parasitology Research* 2013, 831947.
- Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M and Gingeras TR (2013) STAR: ultrafast universal RNA-seq aligner. *Bioinformatics (Oxford, England)* 29, 15–21.
- Duboucher C, Noël C, Durand-Joly I, Gerbod D, Delgado-Viscogliosi P, Jouvesshomme S, Leclerc C, Cartolano GL, Dei-Cas E, Capron M and Viscogliosi E (2003) Pulmonary coinfection by *Trichomonas vaginalis* and *Pneumocystis* sp. as a novel manifestation of AIDS. *Human Pathology* 34, 508–511.
- Duran-Pinedo AE, Paster B, Teles R and Frias-Lopez J (2011) Correlation network analysis applied to complex biofilm communities. *PLoS One* 6, e28438.
- El Sayed Zaki M, Raafat D, El Emshaty W, Azab MS and Goda H (2010) Correlation of *Trichomonas vaginalis* to bacterial vaginosis: a laboratory-based study. *Journal of Infection in Developing Countries* 4, 156–163.
- Fichorova RN, Buck OR, Yamamoto HS, Fashemi T, Dawood HY, Fashemi B, Hayes GR, Beach DH, Takagi Y, Delaney ML, Nibert ML, Singh BN and Onderdonk AB (2013) The Villain Team-Up or how *Trichomonas vaginalis* and bacterial vaginosis alter innate immunity in concert. *Sexually Transmitted Infections* 89, 460–466.
- Franzosa EA, McIver LJ, Rahnavard G, Thompson LR, Schirmer M, Weingart G, Lipson KS, Knight R, Caporaso JG, Segata N and Huttenhower C (2018) Species-level functional profiling of metagenomes and metatranscriptomes. *Nature Methods* 15, 962–968.
- Friedman J and Alm EJ (2012) Inferring correlation networks from genomic survey data. *PLoS Computational Biology* 8, e1002687.
- Gao F, Bian L, Hao X, Hu Y, Yao X, Sun S, Chen P, Yang C, Du R, Li J, Zhu F, Mao Q and Liang Z (2018) Seroepidemiology of coxsackievirus B5 in infants and children in Jiangsu province, China. *Human Vaccines & Immunotherapeutics* 14, 74–80.
- Gaudin M and Desnues C (2018) Hybrid capture-based next generation sequencing and its application to human infectious diseases. *Frontiers in Microbiology* 9, 2924.
- Goldstein E, Murphy TF and Parameswaran GI (2009) *Moraxella catarrhalis*, a human respiratory tract pathogen. *Clinical Infectious Diseases* 49, 124–131.
- Gouy M, Guindon S and Gascuel O (2010) SeaView version 4: a multiplatform graphical user interface for sequence alignment and phylogenetic tree building. *Molecular Biology and Evolution* 27, 221–224.
- Handley SA, Thackray LB, Zhao G, Presti R, Miller AD, Droitt L, Abbink P, Maxfield LF, Kambal A, Duan E, Stanley K, Kramer J, Macri SC, Permar SR, Schmitz JE, Mansfield K, Brechley JM, Veazey RS, Stappenbeck TS, Wang D, Barouch DH and Virgin HW (2012) Pathogenic simian immunodeficiency virus infection is associated with expansion of the enteric virome. *Cell* 151, 253–266.
- Handrich MR, Garg SG, Sommerville EW, Hirt RP and Gould SB (2019) Characterization of the BspA and Pmp protein family of trichomonads. *Parasites & Vectors* 12, 406.
- Janowski AB, Krishnamurthy SR, Lim ES, Zhao G, Brechley JM, Barouch DH, Thakwalakwa C, Manary MJ, Holtz LR and Wang D (2017) Statoviruses, A novel taxon of RNA viruses present in the gastrointestinal tracts of diverse mammals. *Virology* 504, 36–44.
- Jarrett OD, Srinivasan S, Richardson BA, Fiedler T, Wallis JM, Kinuthia J, Jaoko W, Mandaliya K, Fredricks DN and McClelland RS (2019) Specific vaginal bacteria are associated with an increased risk of *Trichomonas vaginalis* acquisition in women. *Journal of Infectious Diseases* 220, 1503–1510.
- Ji F, Zhang D, Shao Y, Yu X, Liu X, Shan D and Wang Z (2020) Changes in the diversity and composition of gut microbiota in pigeon squabs infected with *Trichomonas gallinae*. *Scientific Reports* 10, 19978.
- Juliano C, Cappuccinelli P and Mattana A (1991) In vitro phagocytic interaction between *Trichomonas vaginalis* isolates and bacteria. *European Journal of Clinical Microbiology & Infectious Diseases: Official Publication of the European Society of Clinical Microbiology* 10, 497–502.
- Kang HJ, Yoon Y, Lee YP, Kim HJ, Lee DY, Lee JW, Hyeon JY, Yoo JS, Lee S, Kang C, Choi W and Han MG (2021) A different epidemiology of enterovirus A and enterovirus B co-circulating in Korea, 2012–2019. *Journal of the Pediatric Infectious Diseases Society* 10, 398–407.
- Karched M, Furgang D, Planet PJ, DeSalle R and Fine DH (2012) Genome sequence of *Aggregatibacter actinomycetemcomitans* RHAA1, isolated from a rhesus macaque, an Old World primate. *Journal of Bacteriology* 194, 1275–1276.
- Kim JY, Whon TW, Lim MY, Kim YB, Kim N, Kwon MS, Kim J, Lee SH, Choi HJ, Nam IH, Chung WH, Kim JH, Bae JW, Roh SW and Nam YD (2020) The human gut archaeome: identification of diverse haloarchaea in Korean subjects. *Microbiome* 8, 114.
- Kondova I, Simon MA, Klumpp SA, MacKey J, Widmer G, Domingues HG, Persengiev SP and O'Neil SP (2005) Trichomonad gastritis in rhesus macaques (*Macaca mulatta*) infected with simian immunodeficiency virus. *Veterinary Pathology* 42, 19–29.
- Kopylova E, Noé L and Touzet H (2012) SortMeRNA: fast and accurate filtering of ribosomal RNAs in metatranscriptomic data. *Bioinformatics (Oxford, England)* 28, 3211–3217.
- Kulda J (1999) Trichomonads, hydrogenosomes and drug resistance. *International Journal for Parasitology* 29, 199–212.
- Lahti L and Shetty S (2017) microbiome R package. Bioconductor. <https://doi.org/10.18129/B9.bioc.microbiome>
- Laing ST, Merriam D, Shock BC, Mills S, Spinner A, Reader R and Hartigan-O'Connor DJ (2018) Idiopathic colitis in rhesus macaques is associated with dysbiosis, abundant enterochromaffin cells and altered T-cell cytokine expression. *Veterinary Pathology* 55, 741–752.
- Law CW, Chen Y, Shi W and Smyth GK (2014) Voom: precision weights unlock linear model analysis tools for RNA-seq read counts. *Genome Biology* 15, R29.
- Leinonen R, Sugawara H and Shumway M and Collaboration INSD (2011) The sequence read archive. *Nucleic Acids Research* 39, D19–D21.
- Letunic I and Bork P (2019) Interactive Tree of Life (iTOL) v4: recent updates and new developments. *Nucleic Acids Research* 47, W256–W259.
- Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R and Subgroup GPPD (2009) The sequence alignment/map format and SAMtools. *Bioinformatics (Oxford, England)* 25, 2078–2079.
- Li WC, Ying M, Gong PT, Li JH, Yang J, Li H and Zhang XC (2016) *Pentatrichomonas hominis*: prevalence and molecular characterization in humans, dogs, and monkeys in Northern China. *Parasitology Research* 115, 569–574.
- Li F, Hitch TCA, Chen Y, Creevey CJ and Guan LL (2019) Comparative metagenomic and metatranscriptomic analyses reveal the breed effect on the rumen microbiome and its associations with feed efficiency in beef cattle. *Microbiome* 7, 6.
- Li T, Liu ZH, Zhang Z, Bai HH, Zong XN, Wang FJ and Fan LY (2021) Comparative analysis of the vaginal microbiome of Chinese women with *Trichomonas vaginalis* and mixed infection. *Microbial Pathogenesis* 154, 104790. doi: 10.1016/j.micpath.2021.104790.
- Lin H and Peddada SD (2020) Analysis of compositions of microbiomes with bias correction. *Nature Communications* 11, 3514.
- Malard F, Dore J, Gaugler B and Mohty M (2021) Introduction to host micro-ecology in health and disease. *Mucosal Immunology* 14, 547–554.
- Malik SB, Brochu CD, Bilic I, Yuan J, Hess M, Logsdon JM and Carlton JM (2011) Phylogeny of parasitic parabasalids and free-living relatives inferred from conventional markers vs Rpb1, a single-copy gene. *PLoS One* 6, e20774. doi: 10.1371/journal.pone.0020774.
- Margarita V, Bailey NP, Rappelli P, Diaz N, Dessi D, Fettweis JM, Hirt RP and Fiori PL (2022) Two different species of *Mycoplasma* endosymbionts can influence *Trichomonas vaginalis* pathophysiology. *Mbio* 13, e0091822.
- Maritz JM, Land KM, Carlton JM and Hirt RP (2014) What is the importance of zoonotic Trichomonads for human health? *Trends in Parasitology* 30, 333–341.

- Martin DH, Zozaya M, Lillis RA, Myers L, Nsuami MJ and Ferris MJ (2013) Unique vaginal microbiota that includes an unknown Mycoplasma-like organism is associated with *Trichomonas vaginalis* infection. *Journal of Infectious Diseases* **207**, 1922–1931.
- Martínez-Herrero MDC, Garijo-Toledo MM, González F, Bilic I, Liebhart D, Ganas P, Hess M and Gómez-Muñoz MT (2019) Membrane associated proteins of two *Trichomonas gallinae* clones vary with the virulence. *PLoS One* **14**, e0224032.
- Marzano V, Mancinelli L, Bracaglia G, Del Chierico F, Vernocchi P, Di Girolamo F, Garrone S, Tchidjou Kuekou H, D'Argenio P, Dallapiccola B, Urbani A and Putignani L (2017) “Omic” investigations of protozoa and worms for a deeper understanding of the human gut “parasitome”. *PLoS Neglected Tropical Diseases* **11**, e0005916.
- Matchado MS, Lauber M, Reitmeier S, Kacprowski T, Baumbach J, Haller D and List M (2021) Network analysis methods for studying microbial communities: a mini review. *Computational and Structural Biotechnology Journal* **19**, 2687–2698.
- Matthews HM (1986) Carbohydrate utilization by *Trichomonas gallinae* – effects on growth and metabolic-activity under nongrowth conditions. *Journal of Parasitology* **72**, 170–174.
- McMurdie PJ and Holmes S (2013) Phyloseq: an R package for reproducible interactive analysis and graphics of microbiome census data. *PLoS One* **8**, e61217.
- Müller M (1990) Biochemistry of *Trichomonas vaginalis*. In Honigberg BM (ed.), *Trichomonads Parasitic in Humans*. New York: Springer-Verlag, pp. 156.
- Nazik S, Cingöz E, Şahin AR and Ateş S (2018) Evaluation of cases with *Gemella* infection: cross-sectional study. *Journal of Infectious Disease and Epidemiology* **4**, 063. doi: doi.org/10.23937/2474-3658/1510063.
- Nguyen LT, Schmidt HA, von Haeseler A and Minh BQ (2015) IQ-TREE: a fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. *Molecular Biology and Evolution* **32**, 268–274.
- Nurk S, Meleshko D, Korobeynikov A and Pevzner PA (2017) metaSPAdes: a new versatile metagenomic assembler. *Genome Research* **27**, 824–834.
- Oberste MS, Maher K and Pallansch MA (2007) Complete genome sequences for nine simian enteroviruses. *Journal of General Virology* **88**, 3360–3372.
- O'Leary NA, Wright MW, Brister JR, Ciufu S, Haddad D, McVeigh R, Rajput B, Robertse B, Smith-White B, Ako-Adjei D, Astashyn A, Badretdin A, Bao Y, Blinkova O, Brover J, Chetvernin V, Choi J, Cox E, Ermolaeva O, Farrell CM, Goldfarb T, Gupta T, Haft D, Hatcher E, Hlavina W, Joardar VS, Kodali VK, Li W, Maglott D, Masterson P, McGarvey KM, Murphy MR, O'Neill K, Pujar S, Rangwala SH, Rausch D, Riddick LD, Schoch C, Shkeda A, Storz SS, Sun H, Thibaud-Nissen F, Tolstoy I, Tully RE, Vatsan AR, Wallin C, Webb D, Wu W, Landrum MJ, Kimchi A, Tatusova T, DiCuccio M, Kitts P, Murphy TD and Pruitt KD (2016) Reference sequence (RefSeq) database at NCBI: current status, taxonomic expansion, and functional annotation. *Nucleic Acids Research* **44**, D733–D745.
- Pereira-Neves A and Benchimol M (2007) Phagocytosis by *Trichomonas vaginalis*: new insights. *Biology of the Cell* **99**, 87–101.
- Pinheiro J, Biboy J, Vollmer W, Hirt RP, Keown JR, Artuyants A, Black MM, Goldstone DC and Simoes-Barbosa A (2018) The protozoan *Trichomonas vaginalis* targets bacteria with laterally acquired NlpC/P60 peptidoglycan hydrolases. *Mbio* **9**, 17.
- Rathod SD, Krupp K, Klausner JD, Arun A, Reingold AL and Madhivanan P (2011) Bacterial vaginosis and risk for *Trichomonas vaginalis* infection: a longitudinal analysis. *Sexually Transmitted Diseases* **38**, 882–886.
- Rendon-Maldonado JG, Espinosa-Cantellano M, Gonzalez-Robles A and Martinez-Palomo A (1998) *Trichomonas vaginalis*: In vitro phagocytosis of lactobacilli, vaginal epithelial cells, leukocytes, and erythrocytes. *Experimental Parasitology* **89**, 241–250.
- Shannon P, Markiel A, Ozier O, Baliga NS, Wang JT, Ramage D, Amin N, Schwikowski B and Ideker T (2003) Cytoscape: a software environment for integrated models of biomolecular interaction networks. *Genome Research* **13**, 2498–2504.
- Sharma A (2010) Virulence mechanisms of *Tannerella forsythia*. *Periodontology* **2000** **54**, 106–116.
- Shen W and Xiong J (2019) TaxonKit: a cross-platform and efficient NCBI taxonomy toolkit. *bioRxiv*, 513523. doi: 10.1101/513523.
- Sievers F, Wilm A, Dineen D, Gibson TJ, Karplus K, Li W, Lopez R, McWilliam H, Remmert M, Söding J, Thompson JD and Higgins DG (2011) Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. *Molecular Systems Biology* **7**, 539.
- Smura T, Blomqvist S, Kolehmainen P, Schuffenecker I, Lina B, Böttcher S, Diedrich S, Löve A, Brytting M, Hauzenberger E, Dudman S, Ivanova O, Lukasev A, Fischer TK, Midgley S, Susi P, Savolainen-Kopra C, Lappalainen M and Jääskeläinen AJ (2020) Aseptic meningitis outbreak associated with echovirus 4 in Northern Europe in 2013–2014. *Journal of Clinical Virology: The Official Publication of the Pan American Society for Clinical Virology* **129**, 104535.
- Sommer U, Costello CE, Hayes GR, Beach DH, Gilbert RO, Lucas JJ and Singh BN (2005) Identification of *Trichomonas vaginalis* cysteine proteases that induce apoptosis in human vaginal epithelial cells. *Journal of Biological Chemistry* **280**, 23853–23860.
- Sun L, Dong S, Ge Y, Fonseca JP, Robinson ZT, Mysore KS and Mehta P (2019) DiVenn: an interactive and integrated web-based visualization tool for comparing gene lists. *Frontiers in Genetics* **10**, 421.
- Suyama M, Torrents D and Bork P (2006) PAL2NAL: robust conversion of protein sequence alignments into the corresponding codon alignments. *Nucleic Acids Research* **34**, W609–W612.
- Suzek BE, Wang Y, Huang H, McGarvey PB, Wu CH and Consortium U (2015) UniRef clusters: a comprehensive and scalable alternative for improving sequence similarity searches. *Bioinformatics (Oxford, England)* **31**, 926–932.
- Viscogliosi E and Müller M (1998) Phylogenetic relationships of the glycolytic enzyme, glyceraldehyde-3-phosphate dehydrogenase, from parabasalid flagellates. *Journal of Molecular Evolution* **47**, 190–199.
- Wang CH, Zhang C and Xing XH (2016) Xanthine dehydrogenase: an old enzyme with new knowledge and prospects. *Bioengineered* **7**, 395–405.
- Watts GS, Thornton JE, Youens-Clark K, Ponsoero AJ, Slepian MJ, Menashi E, Hu C, Deng W, Armstrong DG, Reed S, Cranmer LD and Hurwitz BL (2019) Identification and quantitation of clinically relevant microbes in patient samples: comparison of three k-mer based classifiers for speed, accuracy, and sensitivity. *PLoS Computational Biology* **15**, e1006863.
- Wei YX, Gao J, Kou YB, Meng LY, Zheng XP, Liang M, Sun HX, Liu ZZ and Wang YG (2020) Commensal bacteria impact a protozoan's integration into the murine Gut microbiota in a dietary nutrient-dependent manner. *Applied and Environmental Microbiology* **86**, 12.
- Weiss S, Van Treuren W, Lozupone C, Faust K, Friedman J, Deng Y, Xia LC, Xu ZZ, Ursell L, Alm EJ, Birmingham A, Cram JA, Fuhrman JA, Raes J, Sun F, Zhou J and Knight R (2016) Correlation detection strategies in microbial data sets vary widely in sensitivity and precision. *The ISME Journal* **10**, 1669–1681.
- Westreich ST, Ardeshir A, Alkan Z, Kable ME, Korf I and Lemay DG (2019) Fecal metatranscriptomics of macaques with idiopathic chronic diarrhoea reveals altered mucin degradation and fucose utilization. *Microbiome* **7**, 41.
- Westrop GD, Wang L, Blackburn GJ, Zhang T, Zheng L, Watson DG and Coombs GH (2017) Metabolomic profiling and stable isotope labelling of *Trichomonas vaginalis* and *Trichomonas foetus* reveal major differences in amino acid metabolism including the production of 2-hydroxyisocaproic acid, cystathionine and S-methylcysteine. *PLoS One* **12**, e0189072.
- Wood DE, Lu J and Langmead B (2019) Improved metagenomic analysis with Kraken 2. *Genome Biology* **20**, 257.
- Yurewicz EC, Matsuura F and Moghissi KS (1987) Structural studies of sialylated oligosaccharides of human midcycle cervical mucin. *Journal of Biological Chemistry* **262**, 4733–4739.
- Zaragoza MM, Sankaran-Walters S, Canfield DR, Hung JK, Martinez E, Ouellette AJ and Dandekar S (2011) Persistence of gut mucosal innate immune defenses by enteric α -defensin expression in the simian immunodeficiency virus model of AIDS. *Journal of Immunology* **186**, 1589–1597.
- Zhang Q, Han S, Liu K, Luo J, Lu J and He H (2019a) Occurrence of selected zoonotic fecal pathogens and first molecular identification of. *Biomedical Research International* **2019**, 2494913.
- Zhang W, Kataoka M, Doan HY, Ami Y, Suzuki Y, Takeda N, Muramatsu M and Li TC (2019b) Characterization of a novel simian sapelovirus isolated from a cynomolgus monkey using PLC/PRF/5 cells. *Scientific Reports* **9**, 20221.