

# Investigation of the koala (Phascolarctos cinereus) hindgut microbiome via 16S pyrosequencing

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#### 2 Abstract

As a dietary source, the foliage of *Eucalyptus* spp. is low in available protein and carbohydrate 3 while containing polyphenolic compounds that interfere with enzymatic digestion. To overcome 4 5 this, the koala (Phascolarctos cinereus) has evolved a range of anatomical and physiological adaptations to assist with digestion and absorption of nutrients from this food source. Microbial 6 7 fermentation of partially digested eucalyptus leaves is thought to be critical in this process, however, little is known about the composition and diversity of microorganisms that are associated 8 with digestive health in this native species. In this study, we performed 16S rRNA gene 9 pyrosequencing of caecum, colon and faecal pellet samples from two wild, free ranging, 10 Queensland koalas. Our results reveal a highly complex and diverse ecosystem with considerable 11 12 intra-individual variation. Although samples were dominated by sequences from the Bacteroidetes and Firmicutes phyla there was considerable variation at the genus level. This study is the first non-13 14 culture based microbiota analysis, using 454-amplicon pyrosequencing, and provides preliminary 15 data to expand our understanding of the koala hindgut.

16

#### 17 Keywords

18 16S; Koala; Microbiome; gut microbiome; 16SrRNA sequencing

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#### 20 Introduction

As the last surviving member of a family of prehistoric marsupials (*Phascolarctidae*), the koala (*Phascolarctos cinereus*) is a biologically fascinating Australian arboreal marsupial. Part of the uniqueness of this Australian marsupial comes from its unusual reliance on consumption of foliage from species of *Eucalyptus* (Cork and Sanson, 1990). To other animals, the leaves of *Eucalyptus* spp. are a nutrient-poor and toxic source of food as they are low in available proteins and carbohydrates while having high concentrations of phenolic compounds (Cork and Foley, 1997).

The koala has evolved a range of anatomical and physiological adaptations allowing it to overcome challenges associated with the consumption of *Eucalyptus* spp. leaf. Koalas possess a unique gastrointestinal tract, comprised of a foregut and a greatly enlarged hindgut, including the proximal colon and caecum (Cork and Sanson, 1990). The ratio of caecum size to body size of the koala is the largest of any known mammal, with the mean retention time of matter passing through the gastrointestinal tract thought to be one of the longest recorded for mammals (Krockenberger and Hume, 2007).

The koala's hindgut is thought to be the primary location of microbial fermentation, an essential process required for further breakdown of eucalypt leaf cells. The exact identity and role of microorganisms in this process is currently unclear. However, the important role these organisms play in koala digestive health is highlighted by the phenomenon of pap feeding. Pap is a thick faecal paste made in the caecum of female koalas, and thought to be consumed by their young to assist in inoculation of the gastrointestinal tract microflora (Osawa et al., 1993).

In an effort to characterise the role of microorganisms in the gastrointestinal tract of koalas, culturebased studies have revealed a mixture of previously described and novel microorganisms (Osawa et al., 1995; Osawa, 1990). Interestingly, a repeating theme of these studies was the identification of tannin-degrading microorganisms including *Streptococcus bovis* (Osawa, 1990) and a new species of bacteria, *Lonepinella koalarum*, within the family *Pasteurellaceae* (Osawa et al., 1995). These bacteria were found in different locations of the koala gastrointestinal tract, including the bacterial layer of the caecal wall, and in faecal specimens (Goel et al., 2005).

Despite these findings, the use of culture-based technologies alone has meant that attempts to characterise the diversity and role of the koala gastrointestinal microbiota has been very limited. Human studies have shown that 80% or more of phylotypes inhabiting the gut remain uncultured (Zoetendal et al., 2008), suggesting that there are a significant number of potentially important microorganisms that are yet to be identified in the koala gastrointestinal tract. Culture-independent 16S rRNA gene pyrosequencing has revolutionised our knowledge of the diversity and relationships

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between gastrointestinal tract microflora in humans and animals. In this study we have utilised high-throughput 16S rRNA gene pyrosequencing to perform a preliminary characterisation of the diversity and distribution of bacteria within the caecum, colon and faecal pellet of two wild koalas. Our results show a diverse and complex microbiota in the koala hindgut, and will serve as a basis for further studies to understand the impact different microbial communities have on koala health and disease.

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#### 8 Methods

#### 9 Sample collection

Caecum, colon and faecal pellet samples were obtained from two wild, free ranging Queensland 10 koalas (designated K1 and K2) admitted to the Australia Zoo Wildlife Hospital, Beerwah, 11 12 Queensland, Australia. Animal K1, a healthy 7 year old male, was admitted with severe head trauma after being hit by a car. Physical evaluation resulted in the assignment of a body condition 13 14 score (BCS) of 6 out of 10 (considered within a 'healthy range' for wild koalas). Animal K2, an 8 year old female, was admitted to the hospital after being found severely emaciated with a BCS of 2 15 16 out of 10. Subsequent evaluation resulted in the diagnosis of cystitis (presumed to be due to a chlamydial infection) and severe cachexia of unknown (or undetermined) cause. Both animals were 17 euthanized on welfare grounds and samples provided for microbiome analysis. 18

Caecum, colon and faecal pellet samples were obtained during post mortem shortly after euthanasia.
Caecal content was collected towards the blind end of the caecum, colonic content was collected
approximately 10cm preceding the rectum and a faecal pellet was milked from the rectum. Samples
were immediately stored at -20°C until processed and sequenced (no longer than 8 weeks after
collection).

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#### 25 DNA extraction

1 Samples were thawed at room temperature and then processed for DNA extraction using a QIA amp 2 DNA Stool Mini Kit (QIAGEN, Valencia, CA). Briefly, 1.4 mL of ASL lysis buffer was added to approximately 200 mg of sample in a tube containing ceramic beads for bead beading. The samples 3 were then homogenised using a bead-beater, for 5 minutes in one minute intervals. Subsequent steps 4 5 were followed according to the manufacturer's protocol for 'DNA isolation from stool for pathogen detection'. DNA concentration and purity were determined using a NanoDrop spectrophotometer 6 7 (ND-1000, Nanodrop Technologies, Wilmington) at wavelengths of 230, 260 and 280 nm. DNA preparations were standardised at 10ng/µL for each sample in accordance with sample preparation 8 guidelines for 454 pyrosequencing at the Australian Genome Research Facility. 9

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#### 11 Amplification and pyrosequencing of variable regions 1-3 of the 16S rRNA gene

PCR amplification of variable regions 1-3 (V1-V3) from the 16S rRNA gene and subsequent 454 12 pyrosequencing were performed commercially by the Australian Genome Research Facility 13 14 (Brisbane, Australia). Briefly, PCR amplicons were generated using eubacterial 27F and 519R 15 primers along with AmpliTaq Gold 360 mastermix (Life Technologies, Australia). Thermal cycling 16 consisted of 3 minutes at 94°C, and 34 cycles consisting of 45 seconds at 94°C, 1 minute at 50°C, and 1 minute at 72°C, followed by 7 minutes at 72°C. Resulting amplicons were measured by 17 fluorometry, normalised, measured by qPCR, normalised a second time, and then pooled in 18 equimolar ratios. This amplicon pool was then run on the GS-FLX platform using XLR70 19 chemistry (Roche, Australia). 20

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#### 22 Quality filtering of pyrosequencing reads

Initially, multiple raw sequence files (sff files) were combined into a single file containing all
sequence data, and then fasta and quality files were extracted. This was done using Roche's 454
proprietary analysis software (sff tools). Barcodes were removed from the fasta sequences using the
Quantitative Insights Into Microbial Ecology (QIIME) (Caporaso et al., 2010b) software package's

1 split library script. During this process, sequences were also filtered using quality parameters. 2 Sequences >600 nucleotides in length or <200 nucleotides in length were discarded. Additionally, 3 sequences with a mismatch in the primer sequence, an average quality score <25, a homopolymer run >6, or more than 6 ambiguous bases were also discarded. Resulting sequences had reverse 4 5 primers removed using the truncate reverse primer script in the QIIME software package. The 6 resulting fasta file was de-multiplexed using the split fasta on sample ids script in the QIIME 7 software package. Individual fasta files were then multiplexed into caecum specific, colon specific, faecal pellet specific or a single fasta file containing all sequences, using Roche's sff tools software 8 9 package. These files were used for downstream analysis via the QIIME software pipeline. During pre-processing of sequence files, quality control analysis of all samples was performed using the 10 11 PRINSEQ-lite script (Schmieder and Edwards, 2011). Fasta and quality files were used to generate 12 quality distribution graphs (data not shown) and sample statistics (average length, GC content, and 13 occurrence of N) for sequences from each sample (Supplementary Table 1).

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#### 15 Analysis of quality filtered V1-V3 pyrosequencing reads

16 Pyrosequencing reads within a 97% similarity threshold were identified with UCLUST (Edgar, 2010). The most abundant sequence within each new or reference OTU was chosen as the OTU 17 representative sequence. The PyNAST alignment algorithm (Caporaso et al., 2010a) was used to 18 19 align OTU representative sequences against the Greengenes core set alignment (DeSantis et al., 2006) with a minimum alignment length of 365 and a minimum identity of 75% (177 OTU 20 representative sequences failed to align). The OTU representative sequences were screened for 21 22 chimeras with identified sequences being excluded from the downstream analysis. The PyNAST 23 alignment of OTU representative sequences was used to infer a phylogenetic tree using FastTree 24 software (Caporaso et al., 2010b). Taxonomy was assigned to each OTU representative sequence 25 using a BLAST method along with the SILVA (108 release) reference database (Quast et al., 2013). OTU tables had chimeras and singletons excluded. An OTU table only containing OTUs with a 26

1 relative abundance >0.1% (most abundant OTU table) was created by calculating the relative 2 abundance for each sample and removing any OTUs with an abundance <0.1%. OTU tables were summarised in QIIME to generate the relative abundance of taxonomic labels to the phylum and 3 level. genus Venn diagrams generated using VENNY 4 were (http://bioinfogp.cnb.csic.es/tools/venny/index.html). Heatmaps to represent the most abundant 5 OTUs from the caecum, colon and faecal pellet samples were constructed in iTOL (Supplementary 6 7 Figure 1.) (Letunic and Bork, 2011).

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#### 9 Sequence diversity analysis

10 Microbial diversity was evaluated within samples ( $\alpha$  diversity) or between samples ( $\beta$  diversity) 11 using QIIME. Alpha diversity was measured with the Chao1 (richness), Phylogenetic Diversity 12 (branch length-based diversity), Shannon diversity (OTU-based diversity), Shannon equitability, 13 and observed species metrics. Rarefaction, to a sub-sampling depth (determined by the minimum 14 number of sequences in a sample) of 6,736 reads/sample, was performed on all samples. Alpha 15 diversity means and corresponding error bars representing standard error of the mean (SEM) were 16 calculated using GraphPad PRISM software (GraphPad Software, CA, USA).

Beta diversity was evaluated with UniFrac, a community dissimilarity metric based on the fraction 17 of unique branch length observed in pairs of communities in a common phylogenetic tree 18 (Lozupone et al., 2007). Rarefaction, to a sub-sampling depth (determined by the minimum number 19 of sequences in a sample) of 7485 reads/sample, was performed on all samples. The phylogenetic 20 distance (UniFrac distance) is calculated as the fraction of unshared branch lengths between the pair 21 of communities. Unweighted UniFrac distances compare microbial communities within a 22 phylogenetic context based on the presence/absence of members, while weighted UniFrac also 23 incorporates relative abundance information (Lozupone et al., 2007). UniFrac-based Principal 24 Coordinate Analysis (PCoA) was used to map the UniFrac distance matrix onto a set of orthogonal 25 26 axes capturing the greatest amount of variation in all the samples tested. Distances between samples

on a PCoA plot reflect the corresponding dissimilarities in their community membership
 (unweighted UniFrac) or community structure (weighted UniFrac).

3

#### 4 **Results**

#### 5 Characterisation of samples

6 Caecum, colon and faecal pellet samples were collected from two wild, free ranging koalas (i.e. 7 surviving on a natural eucalypt diet). Animal K1 was admitted with a serious blunt force trauma 8 injury, but was otherwise considered healthy. Animal K2 was admitted after being found extremely 9 emaciated. It was subsequently diagnosed with severe cachexia of unknown (or undetermined) 10 cause and chlamydial cystitis. Both animals were euthanized on welfare grounds, due to the severity 11 of disease and injuries present.

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#### 13 Analysis of pyrosequencing data

A total of 81,608 (47,345 for animal K1 and 34,263 for animal K2) sequences across all samples 14 15 were generated through pyrosequencing. The average length of these sequences was approximately 16 521 nucleotides, the GC content ranged from 26-67%, and there were 10,723 sequences containing an ambiguous base call (N) (the maximum percentage of N's per sequence ranged from 9-51%). 17 Demultiplexing and quality trimming of the data set resulted in a total of 72,368 (41,801 for animal 18 K1 and 30,567 for animal K2) sequences with an average length of approximately 485 nucleotides, 19 20 an average GC content of approximately 52% and 2,563 containing an ambiguous base call (N) (the 21 maximum percentage of N's per sequence being 1%) (Supplementary Table 1).

The resulting data set was subsequently clustered into operational taxonomic units (OTUs) using the algorithm UCLUST. From a total of 5277 OTUs, 1320 were identified as potential chimeras and a further 2102 sequences were singletons. Removal of chimeras and singletons left 1855 OTUs for downstream analysis (Supplementary Table 2).

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8

#### 1 Composition of hindgut microbial communities from a healthy koala (animal K1)

2 Assignment of consensus taxonomy resulted in identification of 11 different phyla being represented across the koala caecum, colon and faecal pellet samples. Bacteroidetes and Firmicutes 3 were found to be the most abundant pyhla with Bacteroidetes abundance ranging from 11.0% to 4 5 31.4% and Firmicutes abundance ranging from 40.8% to 64.8%. The abundance of Firmicutes was lowest in the caecum and highest in the colon. In the caecum the ratio of Firmicutes:Bacteroidetes is 6 7 close to 1 (1.3:1) whereas in the colon and faecal pellet the ratio is significantly increased (5.9:1 and 3.18:1 respectively) (Supplementary Table 3). Cyanobacteria, Fusobacteria, Planctomycetes, 8 Proteobacteria and Synergistetes are also predominant phyla with abundances greater than 1% 9 across the different sample types. Cyanobacteria, Fusobacteria, Planctomycetes, and Proteobacteria 10 all have relative abundances that do not appear to change across the hindgut. In contrast, there is a 11 12 significant decrease in the abundance of Synergistetes in the faecal pellet sample (6.1%) compared to the caecum (17.0%) and colon (14.8%) (Figure 1A and Supplementary Table 3). 13

14 Due to the sequencing depth being limited to approximately 12,000 reads per sample, initial genus level analysis focused on the most abundant OTUs, defined as OTUs with a relative abundance 15 16 greater than 0.1% in a given sample. These OTUs were used for the generation of Figure 1B, Supplementary Figure 1 and Supplementary Table 4. Genus level OTU classification resulted in 17 18 identification of 26 different genera having a relative abundance greater than 0.1% in any given 19 sample. Alistipes (12.7%), Cloacibacillus (10.9%), Incertae sedis (17.5%), Ruminococcus (23.4%), and OTUs classified as 'Uncultured' (13.4%) were found to be the predominant OTUs across the 20 21 hindgut of the healthy animal. Incertae sedis is a term used to define a taxonomic group where its 22 broader relationships are unknown or undefined. Further investigation of the OTUs (from the OTU 23 table) classified as *Incertae sedis* found that they were either assigned to the order Clostridales or Erysipelotrichales. OTUs classified to the order Clostridales accounted for over 99% of Incertae 24 sedis OTUs (data not shown). For the remainder of the manuscript Incertae sedis will be defined as 25 "Incertae sedis (unknown Clostridales)". OTUs classified as 'Uncultured' were further investigated 26

and found to contain sequences that were classified into five different phyla (Actinobacteria,
Chloroflexi, Firmicutes, Proteobacteria, and Synergistetes) and seven different orders
(Coriobacteriales, Anaerolineales, Clostridiales, Erysipelotrichales, Burkholderiales, Pasteurellales,
and Synergistales). However, the majority (>75%) were classified to the order Clostridiales
(Supplementary Table 5).

The predominant genera (>10% relative abundance) of the caecum was Alistipes (23.4%), 6 Cloacibacillus (15.9%), Ruminococcus (18.6%), and OTUs classified as 'Uncultured' (13.7%). The 7 predominant genera of the colon were Cloacibacillus (12.1%), Incertae sedis (unknown 8 Clostridales) (35.1%), Ruminococcus (16.6%), and OTUs classified as 'Uncultured' (14.1%). 9 Finally, the predominant genera of the faecal pellet were Bacteroides (10.1%), Incertae sedis 10 (unknown Clostridales) (13.9%), Ruminococcus (35.1%), and OTUs classified as 'Uncultured' 11 12 (12.3%) (Figure 1B and Supplementary Table 4). These results demonstrate a changing microbiome across the hindgut of a healthy koala. 13

14

Comparative analysis of OTU composition across different sites of a healthy hindgut (animal K1) 15 Comparative analysis of OTUs identified in the caecum, colon and faecal pellet of the healthy koala 16 (K1), found that the majority of OTUs were shared across all three gastrointestinal sites. Of the 54 17 different genera (including OTUs with <0.1% relative abundance) identified in the gastrointestinal 18 tract of animal K1, 26 were found to be shared across the caecum, colon and faecal pellet, five were 19 found to be unique to the caecum, eight unique to the colon, and seven unique to the faecal pellet. 20 21 Furthermore, two OTUs were found to be shared exclusively between the caecum and the colon, 22 three were shared exclusively between the colon and the faecal pellet and four were shared 23 exclusively between the caecum and faecal pellet (Figure 5).

In regards to abundance, OTUs unique to the caecum, colon, and faecal pellet, along with OTUs shared exclusively between the different sites were found to make up a very small minority of the microbiome (0.4-1%). The vast majority (99%) of OTUs are shared across the different sites,

1 however, the abundance of OTUs changes across the gastrointestinal tract. Alistipes is the dominant 2 genus of the caecum (21.9%) yet its abundance in the colon (4.7%) and faecal pellet (9.5%) is 3 significantly diminished. Relative abundance of the *Cloacibacillus* genus appears to gradually decline across the gastrointestinal tract (caecum; 14.7%, colon; 11.6, faecal pellet; 4.2%). Incertae 4 5 sedis (unknown Clostridales) is the most dominant genera in the colon (33.8%), however it is 6 significantly lower in the caecum (4.2%) and faecal pellet (14.2%). In contrast, the *Ruminococcus* 7 genus makes up over a third of the faecal pellet microbiome (33.4%), and yet it is much less abundant in the colon (16.5%) and caecum (18.2%). There are also examples of dominant genera 8 maintaining a similar level of abundance across the different sites. Bacteroides, Bilophila, 9 *Fusobacterium*, and OTUs classified as 'Uncultured' appear to have a consistent relative abundance 10 across the three different gastrointestinal tract sites (Figure 5). 11

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#### 13 Richness and diversity of OTUs identified from the koala hindgut

14 Rarefaction curves indicate reasonable depth of sequence coverage, with curves beginning to level 15 off after 6000 reads (Figure 2A). Rank abundance curves, generated from complete OTU lists, 16 appear relatively steep, indicating low evenness across all samples with high ranking species having much greater abundances than the low ranking species (Figure 3A-C). Evenness for the caecum and 17 pellet appear to be comparable, however colon samples show less evenness (Figure 3D). The alpha 18 diversity metrics, Chao1 (measuring richness), Phylogenetic diversity, Shannon equitability 19 20 (measuring evenness) and Shannon Diversity (measuring diversity whilst accounting for abundance 21 and evenness) showed no significant differences between different sample types, suggesting little 22 variability across the gastrointestinal tract samples (Figure 2B-E).

In unweighted UniFrac PCoA, the first principle coordinate (PC1) captured 52.8% of variation with the second principle coordinate (PC2) capturing 17.3% of variation (Figure 4A & C). Although there were not enough samples to undertake thorough statistical analysis, samples from animal K1 clustered together tightly. In contrast, samples from animal K2 were found to cluster away from the

K1 samples, suggesting that the microbiomes of the two animals are phylogenetically distinct. In addition, samples from animal K2 did not cluster in relation to anatomical site, suggesting distinct phylogenetic diversity across different anatomical sites of the hindgut. Samples from animal K1 showed tight clustering of the caecum, colon and faecal pellet, suggesting that phylogenetic diversity of a healthy hindgut is comparable across the different sites. Both of these findings were supported by the weighted UniFrac analysis where PC1 accounted for 59.6% variation and PC2 for 29.9% variation (Figure 4B & D).

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# 9 Comparative analysis of hindgut OTU composition from a healthy (animal K1) versus diseased 10 (animal K2) koala

11 Animal K2 was severely emaciated and suffering from severe cachexia of unknown (or 12 undetermined) cause. Phylogenetic analysis of OTUs (with a relative abundance greater than 0.1%) 13 identified in the caecum, colon and faecal pellet demonstrate a clear difference in OTU presence 14 and abundance between the two animals (Supplementary Figure 1).

Comparative analysis revealed that from 60 caecal OTUs, classified to the genus level, 23 were 15 16 exclusive to animal K2, 11 were exclusive to animal K1 and 26 were shared (Supplementary Figure 2A). Of the 11 caecum OTUs that were exclusive to animal K1, there were three with a relative 17 abundance greater than 1% (Bilophila, Cloacibacillus, and Fusobacterium). The total abundance of 18 animal K1 caecum exclusive OTUs accounted for 7.7% of the total caecal microbiome. For animal 19 K2 caecum exclusive OTUs (23), there were only two with a relative abundance greater than 1% 20 (Enterobacter and Candidatus Tammella). OTUs found only in the caecum of animal K2 accounted 21 22 for 6.3% of the total caecal microbiome. Interestingly, this exclusive group of OTUs included genera of known gastrointestinal pathogens (Enterobacter and Escherichia-shigella) at low levels 23 of abundance. Although *Escherichia* and *Shigella* are two separate genera the extremely high level 24 of genetic relatedness between the different species means that 16S classification databases are 25 unable to resolve which genus the 454 pyrosequencing reads fall into. Therefore, the genus level 26

classification of OTUs is described as; '*Escherichia-shigella*'. Although the 26 shared OTUs make
up the majority of the caecal microbiome (92-93%), there were distinct differences in relative
abundance between the two animals. The relative abundance of *Alistipes, Cloacibacillus,* and *Ruminococcus* was much greater in animal K1, whereas the relative abundance of *Desulfovibrio* and *Parabacteroides* were much higher in animal K2 (Supplementary Table 6).

6 In the colon samples, 23 OTUs were shared between animals, 16 were exclusive to animal K1 and 19 exclusive to animal K2 (Supplementary Figure 2B). The OTUs exclusive to animal K1 make up 7 7.3% of the colon microbiome. Only two colon exclusive OTUs had a relative abundance greater 8 than 1% (Bilophila and Fusobacterium). For animal K2, two colon exclusive OTUs were found to 9 have an abundance greater than 1% (Candidatus Tammella and Flavonifractor), with the colon 10 exclusive OTUs accounting for only 3.4% of the total microbiome. Interestingly, *Enterobacter* and 11 Escherichia-shigella were not identified in the colon. In regard to the shared OTUs, similar to the 12 caecum, the genera *Cloacibacillus* and *Ruminococcus* were found to be in greater abundance in 13 14 animal K1, whereas the relative abundance of *Desulfovibrio* and *Parabacteroides* were higher in 15 animal K2 (Supplementary Table 6).

In the faecal pellet samples, 26 OTUs were shared between the animals, 14 were exclusive to 16 animal K1 and 16 exclusive to animal K2 (Supplementary Figure 2C). Exclusive OTUs make up 17 10.0% and 6.4% of the total faecal microbiomes for animals K1 and K2 respectively. *Bilophila*, 18 Cloacibacillus, and Fusobacterium were found to be the most abundant (>1%), exclusive OTUs, in 19 animal K1's faecal pellet. In contrast, only a single OTU, exclusive to animal K2 faecal pellet, was 20 found to be greater than 1% (*Enterobacter*). In regards to OTUs shared between the two animals; 21 Alistipes, Bacteroides, and Ruminococcus appear significantly more dominant in animal K1 22 23 whereas *Blautia*, *Incertae sedis* (unknown Clostridales), and *Parabacteroides* appear significantly more dominant in animal K2 (Supplementary Table 6). 24

25

26 Discussion

High throughput pyrosequencing of the 16S rRNA gene has revealed a diverse and complex
microbiota in the hindgut of the koala. Microbiota composition was found to vary across the
different anatomical sites and between the two different animals.

Analysis of microbial diversity found no significant differences when measuring the alpha diversity 4 5 of the different samples using several metrics (Phylogenetic diversity, Chao1, Equitability, and 6 Shannon) (Figure 2B-E) and a rank abundance analysis (Figure 3). Interestingly, UniFrac analysis 7 (beta diversity) found that samples from animal K1 clustered together tightly, suggesting phylogenetic diversity is comparable across the different, hindgut, gastrointestinal sites of the 8 healthy koala (Figure 4). This is similar to findings from the human gastrointestinal tract (Stearns et 9 al., 2011) and the Brazilian Nelore steer (de Oliveira et al., 2013). In contrast, samples from animal 10 K2 clearly clustered away from the corresponding K1 samples and away from one another, 11 suggesting that the sick animal has a distinct phylogenetic diversity profile (Figure 4). 12

Taxonomic classification of OTUs led to the identification of 11 different phyla across the three 13 14 different anatomical sites. Only 0.3% of sequences were not classified to a known phylum. The hindgut ecosystem of the koala is predominately colonised by two phyla, Bacteroidetes and 15 16 Firmicutes. These two phyla, which comprised 77% of our sequences, are also the most ubiquitous phyla found across the vertebrate gut (Ley et al., 2008). The ratio of Firmicutes to Bacteroidetes 17 (FB ratio) has been associated with a number of different variables in humans, including age 18 (Mariat et al., 2009), obesity (Ley et al., 2006) and diet (De Filippo et al., 2010). Generally, a high 19 FB ratio has been associated with adults aged 20-45 years, obese people and a diet consisting of 20 animal protein and low levels of fibre (De Filippo et al., 2010; Ley et al., 2006; Mariat et al., 2009). 21 22 In the current study we found that the FB ratio changes considerably across the koala hindgut. In the 23 caecum the ratio is close to 1, whereas the ratio increases significantly in the colon ( $\sim 6:1$ ) and then drops in the faecal pellet ( $\sim$ 3:1) (Supplementary Table 3). Although the FB ratio has been found to 24 vary across different vertebrates, in the majority of species it is closer to 3 (Lev et al., 2008). Our 25 finding suggests that the bacterial fermentation process undertaken in the caecum may differ to that 26

of the colon, with the caecum requiring a higher proportion of Bacteroidetes. Interestingly, a recent
study investigating the microbiota of the marsupial macropod (kangaroos and wallabies) also
identified a low FB ratio in the forestomach region where microbial fermentation of ingested plant
material occurs (Gulino et al., 2013).

At a deeper level of taxonomic classification, six genera were found to represent 85% of all 5 6 sequences. These genera include Alistipes, Bacteroides, Cloacibacillus, Incertae sedis (unknown 7 Clostridales), Ruminococcus, and OTUs classified as 'Uncultured' (Supplementary Table 5). Although a variety of dominant genera were identified in the hindgut, there were significant 8 differences in the abundance of different microbial communities for the caecum, colon and faecal 9 pellet (Figure 1B, and 5 Supplementary Table 4 and Supplementary Figure 1). The caecum of 10 animal K1 (healthy) was dominated by Alistipes (23%), Ruminococcus (19%) and Cloacibacillus 11 12 (16%). The identification of *Ruminococcus* in the caecum is not surprising due to it containing species associated with efficient degradation of plant cell walls (Flint et al., 2008). Unsurprisingly, 13 14 the highest abundance of *Ruminococcus* is found in the faecal pellet (which is indicative of the lower section of the hind gut) where it can take advantage of dietary residue (plant cell wall) that 15 16 has evaded digestion in the upper gastrointestinal tract (Flint et al., 2008). Cloacibacillus is a recently defined genus with a species isolated from the gastrointestinal tract of a pig, which is 17 capable of degrading mucin (Looft et al., 2012). However, little is known about this genus in 18 regards to its role in the digestive tract. Given the considerable abundance of this genus in both the 19 caecum and colon (12%), it may warrant further characterisation. In the lower hindgut (colon and 20 faecal pellet), there appears to be a considerable shift to the Firmicutes phylum (as observed 21 through the FB ratio), which is associated with a significant increase in the abundance of 22 23 *Ruminococcus* and other Firmicutes (OTUs classified as both *Incertae sedis* (unknown Clostridales) and 'Uncultured'). Further analysis is required to clearly elucidate the dominant communities found 24 in the colon due to genus level classification not clearly resolving the Clostridiales identified at 25 these sites. 26

1 Alistipes is a recently defined genus created from the re-classification of Bacteroides putredinis 2 (Rautio et al., 2003). More recently it has been associated with an important role in the metabolism 3 of d-pinitol (Zhao et al., 2013). A study with *Bacillus subtilis* has found that this bacterium can use pinitol as a sole carbon source, thought to be via the *myo*-inositol catabolism pathway. Pinitol is an 4 5 inositol-related compound, a 3-O-methyl of d-chiro-inositol. The *myo*-inositol catabolic pathway 6 involves step-wise reactions that ultimately yield dihydroxyacetone phosphate and acetyl-CoA, thus 7 acting as an energy source (Morinaga et al., 2006). Both pinitol and *myo*-inositol are cyclitols. Significantly, cyclitols have been found to be key osmotica (a substance that acts to supplement 8 osmotic pressure) in higher plants, and may provide a putative link between adaptation of eucalypts 9 to aridity (Hasegawa et al., 2000). A metabolite profiling study of Eucalyptus trees has found that 10 there are numerous species (including species used as a food source by koalas (e.g. Eucalyptus 11 12 saligna)) containing high levels of myo-inositol (Merchant et al., 2006), suggesting a possible link to the abundance of *Alistipes* in the koala caecum. Interestingly, the abundance of *Alistipes* is 13 14 significantly reduced in the colon and faecal pellet (caecum; 21.9%, colon; 4.7%, & faecal pellet; 15 9.5%). Although the benefit of *Alistipes* to the host is unclear, it was also found to be significantly 16 reduced (9%) in the caecum of animal K2 (diagnosed with severe cachexia of unknown cause).

Besides a significant decrease in caecal abundance of *Alistipes*, animal K2 also appeared to have a 17 significant reduction in both Ruminococcus (K1; 18.3%, K2; 0.47%) and Cloacibacillus (K1; 18 14.7%, K2; 0.08%). The reduction in Ruminococcus could be indicative of reduced bacterial 19 fermentation of cellulose, meaning reduced availability of essential nutrients. Furthermore, there 20 was also a significant increase in abundance of *Parabacteroides* and *Desulfovibrio*. Interestingly, 21 species from both genera are known to be opportunistic pathogens of humans (Goldstein et al., 22 23 2003; Nakano et al., 2008), with the *Desulfovibrio* genus also containing species that are strict animal pathogens (Lawson and Gebhart, 2000). As well as identifying potential opportunistic 24 25 pathogens, comparative analysis of animals K1 and K2 also identified the presence of known gut

pathogens (OTUs classified as either *Escherichia* or *Shigella*) that were exclusively found in the
 gastrointestinal tract of animal K2 (Supplementary Table 6).

3 The comparative analysis of animals K1 and K2 also found that the genus Akkermansia is completely absent from the caecum, colon and pellet of the emaciated animal, K2 (Supplementary 4 5 Table 6). A recent, elegant study by Everard et al. clearly defined a causal role for the species 6 Akkermansia muciniphila in controlling gut inflammation, the gut mucosal barrier and gut peptide 7 secretion (Everard et al., 2013). The abundance of the genus Akkermansia in the healthy animal (K1) is much lower than that found in humans. However, this bacterium is normally found in the 8 mucus layer of the gut and thus the caecal content sampling technique may not have been extensive 9 enough to reveal the true abundance of this important bacterial community. Future studies should 10 incorporate sampling of the host tissue to ensure characterisation of important communities lining 11 the gut epithelium. 12

The koala's sole diet of eucalypt leaf is high in tannins which limit the availability of energy and 13 14 nutrients. Adaptation to this diet has resulted in the koala gastrointestinal tract harbouring tannin 15 degrading bacteria (Goel et al., 2005). Using culturing techniques, Osawa et al. identified the novel 16 tannin degrading bacterium, Streptococcus bovis and Lonepinella koalarum, from the koala caecum and faeces (Osawa et al., 1995; Osawa, 1990). Surprisingly, both bacterial genera were not 17 identified during the initial analysis of this study. Further investigation, via BLAST alignment of 18 either L. koalarum or S. bovis 16S rRNA gene against an OTU fasta file, led to the identification of 19 OTUs that were highly homologous to both 16S sequences (data not shown). OTUs found to be 20 21 homologous to the S. bovis 16S sequence were generally classified to the Firmicutes phylum and as 22 Incertae sedis at the genus level. OTUs homologous to the L. koalarum 16S sequence were 23 classified to the Proteobacteria phylum, Pasteurellales order, and as 'Uncultured' at the genus level. These findings suggest that both S. bovis and L. koalarum are not present in the SILVA database 24 25 used to assign classification. Interestingly, the abundance of these OTUs was very low (particularly L. koalarum, <0.2%), highlighting the importance of rare or low abundant OTUs at a functional 26

1 level. Characterisation of less abundant OTUs is essential as abundance does not necessarily reflect

2 important and complex functionality (Arumugam et al., 2011).

Although this is a preliminary, observational study, it must be noted that limitations of sample size make meaningful intra-animal and inter-animal comparisons very difficult. In addition, variations in diet (based on differing geographical locations of the sampled animals) and sex may have also impacted on the microbiome differences noted in this study. Future studies should attempt to sample larger numbers of animals from multiple geographical locations and attempt to account for factors such as sex, age and diet in order to define a core hindgut microbiome.

In conclusion, this study characterised the predominant bacterial communities across the hindgut of the koala showing significant intra-individual variation, particularly in regards to abundance of bacterial communities in the caecum versus the faecal pellet, and differences between a healthy and diseased koala gastrointestinal tract microbiome. Future studies are needed to further define a core 'healthy' koala gastrointestinal tract microbiome at greater sequencing depth and functional capacity in order to develop a complete picture of the koala microbiome and the role it may play in health and disease.

16

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8

#### 9 Figure Captions

10 **Figure 1.** 

Distribution of 16S rRNA gene sequences (Variable 1-3 regions) from koala hindgut samples. A. Depicts the distribution of samples classified to the level of phyla. B. Depicts the distribution of samples classified to the level of genus. A complete list of relative abundance counts for each phyla and genera can be found in Supplementary tables 3 and 4.

15 **Figure 2.** 

Alpha diversity based on 16S rRNA gene sequences (Variable 1-3 regions) from koala hindgut samples. **A.** Rarefaction curves for the 6 different hindgut samples measured using the observed species metric. **B.** Phylogenetic Diversity measurements (the minimum total branch length of the phylogenetic tree that incorporates all OTUs in a sample). **C.** Chao 1, estimated richness measurement for all hindgut samples. **D.** Shannon equitability index, a measure of evenness, for all hindgut samples. **E.** Shannon diversity index, characterising species diversity by accounting for abundance and evenness, for all hindgut samples.

23 **Figure 3.** 

Rank abundance curves. A. Rank abundance curves for caecal OTUs. B. Rank abundance curves for
colon OTUs. C. Rank abundance curves for faecal pellet OTUs. D. Rank abundance curves for the
caecum, colon and faecal pellet. Averages were taken for each sample site, i.e. caecum rank

1 abundance values were averaged from both animals to produce the caecum curve. For all graphs,

2 the y axis displays OTU relative abundance on a logarithmic scale, and the x axis lists the OTUs in

3 rank order of descending abundance.

4 **Figure 4.** 

5 Principal Coordinate Analysis of unweighted and weighted UniFrac distances for 16S rRNA gene 6 sequences (Variable 1-3 regions) from koala hindgut samples. Beta diversity patterns were explored 7 using Principal Coordinate Analysis (PCoA). Samples were rarefied to 7485 reads/sample for all figures. A. Depicts unweighted clustering of samples from animal K2 (sick animal, represented by 8 circular symbols) and animal K1 (healthy animal, represented by square symbols). B. Depicts 9 10 weighted clustering of samples from animal K2 (sick animal, represented by circular symbols) and animal K1 (healthy animal, represented by square symbols). C. Depicts unweighted clustering of 11 12 samples from the caecum (represented by square symbols), colon (represented by triangle symbols) and the faecal pellet (represented by circular symbols). **D.** Depicts weighted clustering of samples 13 14 from the caecum (represented by square symbols), colon (represented by triangle symbols) and the 15 faecal pellet (represented by circular symbols).

16 **Figure 5.** 

17 A venn diagram depicting the intra-individual variation of OTUs along the hindgut of a healthy

18 koala (K1). The corresponding table lists the relative abundance of OTUs at various sites.









Caecum			Colon
5		8	
			þ
	4	3	
	б		
	Faecal Pelle	et	

ACCEP	Relati	ive Abundan	ce (%)			Relative Abundance (%)			
Genus	CAECUM	COLON	PELLET	Site	Genus	CAECUM	COLON	PELLET	Site
Porphyromonas	0.042	0.020	NA	CAECUM & COLON	4c0d-2	1.132	0.489	1.578	SHARED
Actinobacillus	0.021	NA	0.008	CAECUM & PELLET	Akkermansia	0.042	0.065	0.259	SHARED
Barnesiella	0.021	NA	0.042	CAECUM & PELLET	Alistipes	21.874	4.709	9.526	SHARED
Coprococcus	0.021	NA	0.050	CAECUM & PELLET	Alkaliflexus	0.566	0.195	0.100	SHARED
Saccharofermentans	0.094	NA	0.017	CAECUM & PELLET	Allisonella	0.136	0.013	0.017	SHARED
Butyrivibrio	NA	0.026	0.017	COLON & PELLET	Anaerotruncus	0.063	0.033	0.134	SHARED
Clostridium	NA	0.026	0.092	COLON & PELLET	Anaerovorax	0.231	0.091	0.050	SHARED
Faecalibacterium	NA	0.020	0.033	COLON & PELLET	Bacteroides	8.228	5.621	9.760	SHARED
Mucilaginibacter	0.031	NA	NA	CAECUM ONLY	Bilophila	2.316	1.264	1.570	SHARED
Rikenella	0.063	NA	NA	CAECUM ONLY	Blautia	0.304	0.046	0.267	SHARED
S24-7	0.063	NA	NA	CAECUM ONLY	Cloacibacillus	14.726	11.587	4.199	SHARED
Sarcina	0.042	NA	NA	CAECUM ONLY	Desulfovibrio	0.440	0.137	0.334	SHARED
Vadinha17	0.021	NA	NA	CAECUM ONLY	Dysgonomonas	0.105	0.130	0.109	SHARED
Aerococcus	NA	0.078	NA	COLON ONLY	Fusobacterium	4.287	5.243	3.615	SHARED
Anaerostipes	NA	0.046	NA	COLON ONLY	Incertae sedis	4.161	33.837	14.176	SHARED
Cft112h7	NA	0.013	NA	COLON ONLY	Megasphaera	0.063	0.254	0.058	SHARED
Dorea	NA	0.065	NA	COLON ONLY	No BLAST Hit	0.157	0.410	0.301	SHARED
Enterobacter	NA	0.410	NA	COLON ONLY	Oxalobacter	0.063	0.033	0.142	SHARED
Phocaeicola	NA	0.013	NA	COLON ONLY	Parabacteroides	0.346	0.221	0.225	SHARED
Planomicrobium	NA	0.098	NA	COLON ONLY	Phascolarctobacterium	4.570	0.879	2.738	SHARED
Slackia	NA	0.202	NA	COLON ONLY	Pseudobutyrivibrio	0.084	0.085	0.050	SHARED
Chloroplast	NA	NA	0.017	PELLET ONLY	Roseburia	0.472	1.133	0.292	SHARED
Enterorhabdus	NA	NA	0.067	PELLET ONLY	Rs-k70 termite group	0.073	0.020	0.033	SHARED
Geobacillus	NA	NA	0.017	PELLET ONLY	Ruminococcus	18.248	16.453	33.353	SHARED
Gordonibacter	NA	NA	0.042	PELLET ONLY	Uncultured	14.873	14.987	13.884	SHARED
Parasutterella	NA	NA	0.017	PELLET ONLY	Vadinha49	2.002	0.970	2.763 Page	e 29 of 29 SHARED
Propionibacterium	NA	NA	0.017	PELLET ONLY	Vc2.1 bac22	0.021	0.078	0.033	SHARED