Effects of obesity, energy restriction and neutering on the faecal microbiota of cats

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Abstract

Surveys report that 25–57% of cats are overweight or obese. The most evinced cause is neutering. Weight loss often fails; thus, new strategies are needed. Obesity has been associated with altered gut bacterial populations and increases in microbial dietary energy extraction, body weight and adiposity. This study aimed to determine whether alterations in intestinal bacteria were associated with obesity, energy restriction and neutering by characterising faecal microbiota using 16S rRNA gene sequencing in eight lean intact, eight lean neutered and eight obese neutered cats before and after 6 weeks of energy restriction. Lean neutered cats had a bacterial profile similar to obese rodents and humans, with a greater abundance (P<0.05) of Firmicutes and lower abundance (P<0.05) of Bacteroidetes compared with the other groups. The greater abundance of Firmicutes in lean neutered cats was due to a bloom in Peptostreptococcaceae. Obese cats had an 18% reduction in fat mass after energy restriction (P<0.05). Energy reduction was concurrent with significant shifts in two low-abundance bacterial genera and trends in four additional genera. The greatest change was a reduction in the Firmicutes genus, Sarcina, from 4·54 to 0·65% abundance after energy restriction. The short duration of energy restriction may explain why few bacterial changes were observed in the obese cats. Additional work is needed to understand how neutering, obesity and weight loss are related to changes in feline microbiota and how these microbial shifts affect host physiology.

Key words: Faecal microbiota; Feline nutrition; Obesity; Neutered cats; Energy restriction

Obesity is a common feline nutritional disorder, with surveys reporting between 25 and 57% of cats characterised as overweight or obese1,2,3. Obesity can be defined as an excess of body fat sufficient to impair health or body function and is generally recognised as 20–25% above ideal body weight (BW) in cats4. Obese cats face an increased risk of musculoskeletal problems, diabetes mellitus and hepatic steatosis5,6,7,8,9.

The underlying cause of obesity is an imbalance between energy intake and energy expenditure, resulting in increased energy storage as fat. Exogenous factors leading to energy imbalance include activity level, diet composition and palatability, as well as environment and lifestyle. Endogenous factors include age, sex, reproductive status, hormonal abnormalities and genetics. Of endogenous factors, neutering is the most evinced. Studies have shown that intact adult cats generally weigh less than neutered cats of the same breed and size5,6,9,10. Neutering in cats leads to increased food intake and weight gain due, in part, to changes in growth-promoting and satiety hormones11,12,13. Treatment of obesity frequently focuses on energy restriction; however, lack of owner compliance often results in failure. Therefore, additional strategies are needed to promote weight loss in cats.

One potential strategy involves manipulation of the faecal microbiota. The gut harbours a collection of viruses, bacteria, fungi and parasites collectively referred to as the faecal microbiota14. Bacteria are the most well-characterised members of the faecal microbiota and have been shown to influence host metabolism15 including the development of obesity in humans16. Undesired changes in bacterial composition or function are thought to increase BW and adiposity through a variety of mechanisms including increased inflammation17, increased energy extraction from diet18 and altered production of host satiety hormones19. In humans, obese individuals are reported to have greater proportions of Firmicutes and reduced levels of Bacteroidetes compared with lean controls20. Transplanting faecal microbiota from obese mice into germ-free mice...
recapitulated the obese phenotype in the germ-free mice, whereas germ-free mice receiving microbes from lean mice remained lean\(^{(22)}\). One factor shown to greatly alter the faecal microbiota is diet. In mice, high-fat diets have been shown to increase the Firmicutes:Bacteroidetes ratio\(^{(23)}\) and increase blood concentrations of bacterial-derived pro-inflammatory products containing pathogen-associated molecular compounds (i.e. flagellin, lipopolysaccharide)\(^{(24,25)}\). These bacterial products bind to host immune receptors and induce chronic low-grade inflammation, which over time can lead to impaired satiety hormone signalling resulting in hyperphagia\(^{(26)}\).

Manipulation of gut bacterial populations using diet or antibiotics may be a viable strategy to promote a healthy BW in cats. Studies have characterised the feline faecal microbiota\(^{(27,28)}\); however, few have examined the effect of neutering, cats. Studies have characterised the feline faecal microbiota composition in (1) lean neutered and lean intact cats, (2) lean neutered and obese neutered cats and (3) obese neutered cats before and after 6 weeks of energy restriction with the goal of identifying microbial shifts that occur with neutering or energy restriction.

**Methods**

Approval of the experimental protocol (Protocol 17261) was granted by the Institutional Animal Care and Use Committee of the University of California, Davis.

**Animals and diets**

In all, twenty-four adult (range 1–12 years; median age 6-4 years), specific pathogen free, domestic shorthair cats owned by the University of California were used in this study. There were eight obese (four male and four female); eight lean intact (four male and four female); and eight lean neutered (six male and two female) cats. A nine-point body condition score (BCS) system was used\(^{(29)}\), where a score of 5 was considered ideal, a score >5 and <7 was considered overweight and a score >7 was considered obese. All cats were group-housed in a light (14 h light–10 h dark cycle)- and temperature (18–24°C)-controlled facility at the University of California, Davis, in an enriched environment (perches, rotating toys and scratching poles) and were brushed and socialised once a day. Cats were individually housed for faecal and blood collections. Fresh water was available at all times, except before body composition determination. All cats consumed the same extruded dry-type diet for at least 8 weeks before entering and throughout the duration of the study. BW was measured at all times, except before body composition determination. All cats consumed the same extruded dry-type diet for at least 8 weeks before entering and throughout the study. BW was measured weekly and remained weight-stable long before and throughout the duration of the study, indicating that these cats were consuming food in a quantity close to the standards of maintenance requirements. The night before blood and faecal collection for body composition determination and microbe analysis, cats were BCS and moved into individual cages. Following collection of final blood and faecal samples, all cats were returned to group housing in the feline facility.

**Energy restriction of obese neutered cats**

The obese neutered cats (four male and four female; mean age 7-25 years (range 1–11 years)) were castrated or spayed 1–6 years before entering the study. Cats were briefly individually housed twice a day and fed the above-described diet ad libitum for 10 d, during which time their BW and food intake were stable. The cats were then fed 60–70 % of their previously measured energy intake for a period of 6 weeks. The target for weight loss was 0.5–1 % of BW/week. We confirmed that the diet would still meet the National Research Council’s recommended allowance for adult cats, even with up to 40 % energy restriction\(^{(31)}\). Iodine was the one nutrient that was just below the National Research Council’s recommended allowance for adult cats, but it exceeded the Association of American Feed Control Officials recommendations\(^{(30)}\). Body composition was determined, and faecal and blood samples were taken before the start and end of energy restriction. BW was measured weekly and BCS was determined every other week by the same person.

**Parameters evaluated**

**Body composition determination.** Estimation of body fat mass (FM) and lean mass was determined using the deuterium oxide (D\(_2\)O) isotopic dilution method previously described\(^{(32)}\) with modifications\(^{(33)}\). D\(_2\)O was purchased from Fisher Scientific. A basal blood sample (3 cc), without D\(_2\)O enrichment, was obtained by jugular venepuncture. Cats were fasted (12 h) before sample collection, and water was withheld from cats 2 h before collection. D\(_2\)O (0.4 g D\(_2\)O/kg BW) was administered to the cats subcutaneously and allowed to equilibrate for 3 h, after which a D\(_2\)O-enriched blood sample (3 cc) was collected. Condensed serum water samples were analysed on an ATI Mattson Infinity Series Fourier transform IR spectrometer equipped with a class 2A laser.

**Faecal collection and characterisation of faecal microbiota via bacterial 16S rRNA gene sequencing.** Fresh faecal samples for each cat were collected from the litter box once
daily over 3 consecutive days into sterile tubes, stored at −80°C and pooled. Cats were observed every 15 min by the primary author and staff at the facility, and faeces was only considered fresh if collected within 15 min of defecation. Bacterial DNA was extracted by a bead-beating method using a commercial DNA extraction kit (Mo Bio PowerSoil Kit; Qiagen) according to the manufacturer’s instructions. The bead-beating step was performed on a homogeniser for 60 s at a speed of 4 m/s. Amplification of the 16S rRNA genes was carried out using a universal bacterial primer (27F–519R) for V3–V4 region to amplify DNA in a single-step, 30-cycle PCR reaction using the HotStarTaq Plus Master Mix Kit (Qiagen) under the following conditions: 94°C for 3 min, followed by twenty-eight cycles of 94°C for 30 s, 53°C for 40 s and 72°C for 1 min, after which a final elongation step at 72°C for 5 min was performed. Following the PCR reaction, all amplicon products from different samples were pooled in equal concentrations and purified using Agencourt Ampure beads (Agencourt Biosciences). Samples were sequenced using Roche 454 FLX titanium instruments and reagents according to the manufacturer’s guidelines.

16S rRNA gene data processing. The Q25 sequence data were processed using a proprietary analysis pipeline (www.mrdalab.com)\(^{34,35}\). In brief, sequences were trimmed of barcodes and primers, and then sequences <150 bp were removed, as were sequences with ambiguous base calls and homopolymer runs exceeding 6 bp. Operational taxonomic units (OTU) were generated by clustering at 3% divergence (97% similarity) from de-noised sequences, and chimeras were removed. Final OTU were taxonomically classified using BLASTn (closed reference) against a curated database generated from sequences from GreenGenes\(^{36}\) and Ribosomal Database Project (RDP-II)\(^{37}\) and National Center of Biotechnology Information (NCBI). We obtained a mean of 7701 (SEM 1362) individual sequences from GreenGenes and Ribosomal Database Project (RDP-II)\(^{37}\) and National Center of Biotechnology Information (NCBI). We obtained a mean of 7701 (SEM 1362) individual sequencing reads per sample (min. = 4423; max. = 17893). After data processing, the average number of sequences for each sample passing through to OTU classification was 4491 (SEM 351). The average number of OTU per sample was 548. Data were compiled into each taxonomic level as the percentage of sequencing depth bias. The depth cutoff (2818) was defined by the samples with the lowest number of reads. Alpha and beta diversity measures were calculated using QIIME software (QIIME 1.8.0). Raw sequences reads were deposited at NCBI’s Sequence Read Archive (http://www.ncbi.nlm.nih.gov/Traces/sra) under accession no. SRP066010.

Statistical analysis

Very low abundance taxa (<0·1%) or taxa not represented within at least 50% of the samples within a group were excluded from analysis. Partial least squares-discriminant analysis (PLS-DA) was performed on unadjusted means of genus-level microbiota abundance data. For group comparisons, ANCOVA was performed with sex and age as covariates (Fig. 2(a)–(c)). Significance of differences between lean intact \(v\) lean neutered cats and lean neutered \(v\) obese neutered cats was assessed by Tukey’s honest significant difference test while controlling for a family-wise type I error. Significance of difference between obese neutered cats before and after energy restriction was assessed by paired, two-tailed Student’s \(t\) test. A Spearman’s correlation matrix of age and body composition \(v\) bacterial genera was obtained to assess magnitudes of their correlation. All statistical analyses were performed using R. A two-sided \(P\) value of 0·05 was considered significant. A \(P\) value ≤0·1 is considered as representative of a trend.

Results

No adverse clinical changes were observed throughout the experiment. There were no significant effects of sex or age on any of the variables. Average food intake by obese cats during \textit{ad libitum} and energy restriction phases was 75·7 and 51·6 g/d, respectively.

Body weight and composition

There were no differences in BW, lean or FM between the lean intact and lean neutered cats (Table 1). The lean neutered cats had a lower \((P<0·05)\) lean body mass compared with the obese neutered cats. After 6 weeks of energy restriction, the obese cats lost, on average, 1% of BW/week, resulting in an 18% reduction in FM but not lean mass and a small but significant reduction in BW.

Microbial diversity

Alpha and beta diversity measures of the faecal microbiota were examined. Only the lean neutered group showed a difference in alpha diversity using the phylogenetic measure, Faith’s whole-tree phylogenetic diversity (Fig. 1(a), online Supplementary Table S1). This reduced diversity was not observed using non-phylogenetic measures of diversity such as Shannon, Chao or the number of observed species. Beta diversity was also evaluated. Principal coordinates analyses using unweighted and weighted UniFrac distances clearly demonstrated lack of separation of the groups, indicating no difference in beta diversity between the groups (Fig. 1(b) and (c)).

Multivariate analyses of faecal microbiota

As an initial investigation to determine whether we could identify signatures related to the effects of neutering (Fig. 2(a)), obesity (Fig. 2(b)) and energy restriction in the context of obesity (Fig. 2(c)), we performed PLS-DA using genus-level abundance data. Indeed, in each comparison, PLS-DA discriminated the groups as shown by the scores plots. The loadings show the relative contributions of specific variables to group separation in each comparison. To assess the statistical importance of the variables driving the separation of the groups, we calculated variable importance in projection scores and used scores above the 90th percentile as a cutoff for the most significant contributors (online Supplementary Table S2). Notably, different genera were identified as discriminatory for each comparison. When comparing lean intact \(v\) lean neutered
cats, we identified the genera *Bacteroides*, *Eubacterium*, *Faecalibacterium*, *Phascolarctobacterium* and *Sutterella* as important discriminators of the intact vs. neutered state. In obesity, we identified *Prevotella*, *Acidaminococcus* and *Phascolarctobacterium* as important in distinguishing lean vs. obese neutered cats. In the case of energy restriction, *Acidaminococcus*, *Bacillus*, *Dorea*, *Phascolarctobacterium*, *Sarcina* and *Staphylococcus* were the key contributors to the distinction of the same cat before and after energy restriction.

**Phylum-level faecal microbiota.** The overall mean phylum-level proportions observed in the cat faecal microbiota from all groups in decreasing order of abundance were as follows: *Firmicutes* (65.8%), *Bacteroidetes* (25.2%), *Proteobacteria* (3.52%), *Actinobacteria* (2.20%) and *Fusobacteria* (0.3%). The majority of change in the faecal microbiota was observed between the lean neutered and obese neutered cats. The lean neutered cats had significantly greater proportions of the phylum *Firmicutes* (*P < 0.05*) and significantly lower proportions of *Bacteroidetes* (*P < 0.05*) (Table 2) compared with obese neutered cats. There was also a trend (*P < 0.10*) towards the lean neutered cats showing this same shift in *Firmicutes* and *Bacteroidetes* populations compared with the lean intact cats. There were no other phylum-level differences between lean intact and lean neutered cats or obese neutered cats before and after energy restriction.

**Family-level faecal microbiota.** A total of eighteen bacterial families were identified in the faecal samples. Within the *Firmicutes* phylum, *Lachnospiraceae*, *Peptostreptococcaceae*, *Veillonellaceae* and *Ruminococcaceae* were the predominant families identified in cat faeces (Table 3). The greater abundance of *Firmicutes* and reduced proportions of *Bacteroidetes* in the lean neutered cats compared with obese neutered cats was driven by significantly greater proportions of *Peptostreptococcaceae* (*P = 0.015*) and reduced proportions of *Prevotellaceae* (*P = 0.05*). An unidentified family within the *Bacteroidiales* showed a trend for an increase in the obese neutered cats (*P = 0.077*). There were notable family-level differences between lean intact and lean neutered cats; however, these did not reach statistical significance. There was a trend for 2-fold greater abundance of *Peptostreptococcaceae* (*P = 0.057*) in the lean neutered cats compared with lean intact cats. In addition, there was a tendency towards decreased *Clostridiaceae* (*P = 0.065*) in obese cats after energy restriction.

**Genus-level faecal microbiota.** *Blautia*, *Bacteroides*, *Catenibacterium*, *Clostridium*, *Megasphaera*, *Oscillospira*, *Prevotella*, *Ruminococcus* and *Sarcina* were the predominant genera identified in cat faeces (Table 4). Most of the statistically significant differences observed were between lean neutered and obese neutered cats. Significant changes were greater abundances in the *Bacteroidetes* *Prevotella* and reduced proportions in the *Firmicutes* *Blautia* and *Clostridium* in the obese neutered cats. In the lower abundance genera, there was an increase in *Acidaminococcus*, *Bulleidia* and *Phascolarctobacterium* and a trend for increased *Faecalibacterium* (*P = 0.069*) in the obese neutered group. After energy restriction, the *Bacteroidetes* and *Firmicutes* groups showed a trend for an increase in the obese neutered cats. In the lower abundance genera, there was an increase in *Acidaminococcus*, *Bulleidia* and *Phascolarctobacterium* and a trend for increased *Faecalibacterium* (*P = 0.069*) in the obese neutered group. After energy restriction.
restriction, there was a significant decrease in Acidaminococcus and a significant increase in Staphylococcus. Although statistically insignificant at the 0.05 level, several trends were noted including decreases in Bulleidia (P = 0.058) and Sarcina (P = 0.091) and increases in Bacillus (P = 0.059) and Lactobacillus (P = 0.055) after energy restriction. Prevotella was less abundant in lean neutered cats (P < 0.05) compared with obese neutered cats, and the relative absence of this bacteria was the main contributor to the reduced abundance of Bacteroidetes in this group. Blautia, Clostridium and Lactobacillus were the main bacteria contributing to the greater abundance of Firmicutes observed in lean neutered cats. A notable trend was the 1.9-fold greater abundance of Clostridium (P = 0.057) in the lean neutered cats compared with the lean intact cats. While not meeting the P ≤ 0.1 cutoff as a trend, the lean intact and lean neutered group include a nearly 2.5-fold reduced abundance of Prevotella (P = 0.104); however, these differences were not statistically significant because of high variability among individual cats (Table 4). Correlations among bacterial genera and age, BW, lean and FM are presented in Fig. 3. Several bacteria significantly correlated with age and body composition.

Discussion
To our knowledge, this is the first study comparing gut microbial diversity in lean intact, lean neutered and obese
neutered cats before and after energy restriction. The goal of this study was to identify bacterial signatures that distinguish these groups from one another and determine how these bacterial shifts relate to changes in body composition. Inclusion of the lean neutered cats made this study especially unique because cats usually gain weight after neutering, and therefore this group of cats is less common. Post-neutering weight gain is variable, with 6 months post-neutering BW gain ranging from 3 to 53% (14); however, the reasons behind this variation have not been fully elucidated. Increased food intake (hyperphagia), due at least in part to neutering-induced hormonal alterations and not decreased energy expenditure, has been identified as the main driver of post-neutering weight gain (9,11,14). Interestingly, previous studies in mice have demonstrated a relationship between the faecal microbiota and FM (21), hyperphagia (38) and sex hormones (39). Understanding this complex relationship may prove invaluable in the prevention and/or treatment of post-neutering weight gain.

Fig. 2. Partial least squares-discriminant analysis reveals discriminating characteristics of genus-level microbiota with respect to neutering between lean intact (●) v. lean neutered (○) (a), obesity (lean neutered (○) v. obese neutered (●)) (b) and energy restriction (obese neutered (●) before v. after energy restriction (▲, energy-restricted (ER)-obese)) (c). Inset in each panel displays the scores plot (clustering based on group assignment), with the coloured ellipses representing the 95% confidence of the populations as calculated based on Hotelling’s T2 test; each symbol represents an individual cat. Discrimination of the groups in the scores plot was explained by the variance in the variables indicated in the loadings plot in each panel.
highlighted the importance of this genus in the discrimination of the groups. Previous studies have found *Clostridium* to positively correlate with carbohydrate oxidation and negatively correlate with fat oxidation\(^{44}\). We also observed the genus *Clostridium* to negatively correlate with lean body mass and positively correlate with FM in the lean intact and lean neutered cats. Taken together, these results imply that members of *Clostridium* may influence host macronutrient metabolism and body composition.

Compared with the obese neutered cats, the lean neutered cats had significantly more Firmicutes and less Bacteroidetes, which is in contrast to that commonly reported for obese mice\(^{45}\) and humans\(^{22,46}\). This seemingly contradictory observation lends support to the notion that shifts at lower taxonomic levels (i.e. family or genus) may be more relevant rather than broad phylum-level changes. At the family level the main difference was >2-fold reduced abundance in Peptostreptococcaceae (*P*=0.015) and an almost 2.5-fold reduced abundance in Prevotellaceae (*P*=0.05) in obese neutered cats compared with lean neutered cats. Peptostreptococcaceae has been found to negatively correlate with life span in mice and decrease with energy restriction\(^{47}\). Another study found that feeding rats a high-fat diet for 4 weeks increased Peptostreptococcaceae and decreased Prevotellaceae. To understand how these changes in the microbiota relate to phenotypic changes, correlation analyses revealed a negative correlation between *Roseburia* and FM in the lean and obese neutered cats. *Roseburia* was previously shown to negatively correlate with fasting hyperglycaemia, glucose intolerance, hepatic TAG accumulation and hypercholesterolaemia\(^{48}\). *Roseburia* is known to produce butyrate\(^{49}\), which has been shown to have a number of health benefits including reducing BW gain and increasing insulin sensitivity, as well as satiety hormones\(^{50,51}\). Results from these studies imply that these bacteria may interact with the host to influence metabolism and may therefore warrant further investigation in relation to weight maintenance.

We found 6 weeks of energy restriction in obese cats to have little impact on the faecal microbiota, with only a few changes in bacterial taxa. This may have been due, in part, to the short period of energy restriction in this study. Short-term studies in cats that have observed drastic changes in the microbiota usually are related to shifts to the diet composition, indicating that diet strongly shapes the faecal microbiota\(^{41,52,53}\). In our study, the same diet was used during the weight-loss phase for the obese cats, demonstrating that reducing energy intake by 30–40% for 6 weeks was not enough to induce significant changes in the faecal microbiota. Nevertheless, weight loss was significant, achieving the target weight loss rate of approximately 1% BW per week and inducing significant changes in body composition. Our goal was not to promote marked weight loss, but to evaluate the effect of a moderate energy restriction on changes in faecal microbiota early in the weight loss process that could be driving physiological responses to weight change. We wanted to determine what changes occurred during initial weight loss rather than waiting to see what happens after significant weight loss had already been achieved. Understanding the changes that occur initially during weight loss may aid in identifying targets that may help promote greater weight loss.

### Table 2. Predominant bacterial phyla (expressed as a percent abundance) in the faeces of lean intact, lean neutered and obese neutered cats before and after 6 weeks of energy restriction (mean values with their standard errors; n=8/group)

<table>
<thead>
<tr>
<th>Phylum</th>
<th>Lean intact</th>
<th>Lean neutered after energy restriction</th>
<th>Obese neutered after energy restriction</th>
<th>ANCOVA Tukey's post hoc pairwise comparisons on ANCOVA-adjusted means</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean SEM</td>
<td>Mean SEM</td>
<td>Mean SEM</td>
<td>Overall Mean, s</td>
</tr>
<tr>
<td>Firmicutes</td>
<td>63 ± 36</td>
<td>96 ± 92</td>
<td>36 ± 13</td>
<td>0.08</td>
</tr>
<tr>
<td>Bacteroidetes</td>
<td>27 ± 38</td>
<td>24 ± 36</td>
<td>51 ± 28</td>
<td>0.13</td>
</tr>
<tr>
<td>Actinobacteria</td>
<td>9 ± 72</td>
<td>8 ± 92</td>
<td>40 ± 11</td>
<td>0.95</td>
</tr>
<tr>
<td>Proteobacteria</td>
<td>2 ± 85</td>
<td>3 ± 92</td>
<td>1 ± 72</td>
<td>0.89</td>
</tr>
<tr>
<td>Fusobacteria</td>
<td>0 ± 38</td>
<td>0 ± 38</td>
<td>2 ± 85</td>
<td>0.89</td>
</tr>
<tr>
<td>Unclassified</td>
<td>3 ± 00</td>
<td>2 ± 00</td>
<td>3 ± 00</td>
<td>0.89</td>
</tr>
</tbody>
</table>
Table 3. Bacterial families (expressed as a percent abundance) in the faeces of lean intact, lean neutered and obese neutered cats before and after 6 weeks of energy restriction.

| Family                  | Mean (% of Faeces) | SEM | Mean (% of Faeces) | SEM | Mean (% of Faeces) | SEM | Overall test (P) | Lean intact v. Lean neutered | Lean intact v. Obese neutered | Obese neutered v. Obese after energy restriction |
|-------------------------|--------------------|-----|--------------------|-----|-------------------|-----|----------------|-----------------------------|-------------------------------|--------------------------------|--------------------------------|
| **Firmicutes**          | 49.53              | 1.52| 50.67              | 1.52| 52.61             | 1.52| 0.04           |    |                 |                               |                               |
| **Bacteroidetes**       | 4.09               | 0.23| 3.72               | 0.23| 2.81              | 0.23| 0.002          |    |                 |                               |                               |
| **Proteobacteria**      | 8.47               | 0.53| 9.13               | 0.53| 10.24             | 0.53| 0.005          |    |                 |                               |                               |
| **Fusobacteria**        | 0.45               | 0.07| 0.53               | 0.07| 0.58              | 0.07| 0.23           |    |                 |                               |                               |
| **Epsilonbacteriota**   | 0.32               | 0.02| 0.37               | 0.02| 0.41              | 0.02| 0.22           |    |                 |                               |                               |
| **Actinobacteria**      | 0.24               | 0.01| 0.25               | 0.01| 0.27              | 0.01| 0.66           |    |                 |                               |                               |

†: ANCOVA for group comparisons in mean, adjusted for age and sex as covariates for first three independent groups (lean intact, lean neutered and obese neutered).
‡: ANCOVA for pairwise comparisons on ANCOVA-adjusted mean.
§: Unknown family within the order Clostridales.
¶: Unknown family within the class Clostridiales.
**Significance assessed by two-tailed paired Student t-test.**
Table 4. Bacterial genera (expressed as a percent abundance) in the faeces of lean intact, lean neutered and obese neutered cats before and after 6 weeks of energy restriction (Mean values with their standard errors; n 8/group)

<table>
<thead>
<tr>
<th>Gammaproteobacteria</th>
<th>Lean intact</th>
<th>Lean neutered</th>
<th>Obese neutered</th>
<th>ANCOVA</th>
<th>Tukey's post hoc pair comparisons</th>
<th>Paired test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall test (P)*</td>
<td>0.374</td>
<td>0.700</td>
<td>0.344</td>
<td>0.587</td>
<td>0.322</td>
<td></td>
</tr>
<tr>
<td>Lean intact v. lean neutered†</td>
<td>0.008</td>
<td>0.005</td>
<td>0.101</td>
<td>0.005</td>
<td>0.101</td>
<td></td>
</tr>
<tr>
<td>Lean neutered v. obese neutered†</td>
<td>0.041</td>
<td>0.043</td>
<td>0.043</td>
<td>0.043</td>
<td>0.043</td>
<td></td>
</tr>
<tr>
<td>Obese before v. obese after energy restriction†</td>
<td>0.049</td>
<td>0.049</td>
<td>0.049</td>
<td>0.049</td>
<td>0.049</td>
<td></td>
</tr>
</tbody>
</table>

* ANCOVA for group comparisons in mean, adjusted for age and sex as covariates for first three independent groups (lean intact, lean neutered and obese neutered).
† Tukey's honest significant difference test for post hoc pairwise comparisons on ANCOVA-adjusted means.
‡ Significance assessed by two-tailed paired Student's t test.
Fig. 3. Spearman’s correlation matrix of age and body composition v. bacterial genera. Only significant correlations are shown ($P \leq 0.05$). The coloured bar below the plot indicates positive or negative correlation (Spearman’s $\rho$ rank correlation coefficient) and size of the square indicates strength of correlation (i.e. larger square indicates strong relationship).
loss. Longer-term weight-loss trials examining the faecal microbiota at multiple time points may be a useful approach to determine which bacteria change with weight loss.

In conclusion, the present study reports changes in the faecal microbial population in lean and obese and intact and neutered domestic cats. We observed the greatest alterations in the faecal microbiota when we compared the lean cats with obese cats. We were also able to detect shifts as a result of neutering, but only minor changes elicited by energy restriction in obese cats. Multivariate analyses using PLS-DA discriminated the groups when we specifically examined the effects of neutering, obesity or energy restriction in the context of obesity and identified the genera that contributed to the distinction of those groups. Correlations among faecal bacteria and body composition were observed, which were consistent with previously published findings. Additional work is needed to understand the mechanisms behind how neutering, obesity and weight loss induce changes to the feline microbiota and how these in turn affect host physiology. This information can then potentially be leveraged to develop probiotic supplements that can favourably affect host metabolism and body composition.

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Supplementary material

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