Flexibility and resilience of Great tit (Parus major) gut microbiomes to changing diets

Kasun H. Bodawatta (✉ bodawatta@snm.ku.dk)  
Kobenhavns Universitet  https://orcid.org/0000-0002-6095-9059

Inga Freiberga  
Biologicke centrum Akademie Ved Ceske Republiky

Katerina Puzejova  
Biologicke centrum Akademie Ved Ceske Republiky

Katerina Sam  
Akademicka knihovna Jihoceske Univerzity v Ceskych Budejovicich

Michael Poulsen  
Kobenhavns Universitet

Knud A. Jønsson  
Kobenhavns Universitet

Research

Keywords: Bacterial communities, Community flexibility, Community resilience, Gut symbionts, Illumina MiSeq, 16S rRNA gene

DOI: https://doi.org/10.21203/rs.3.rs-34716/v1

License: ☒ This work is licensed under a Creative Commons Attribution 4.0 International License.  
Read Full License
Abstract

Background: Gut microbial communities play important roles in nutrient management and can change in response to host diets. The extent of this flexibility and the concomitant resilience is largely unknown in wild animals. To begin untangling the dynamics of avian-gut microbiome symbiosis associated with diet changes, we exposed *Parus major* (Great tits) fed with a standard diet (seeds and mealworms) to either a mixed (seeds, mealworms and fruits), a seed, or a mealworm diet for four weeks, and examined the flexibility of gut microbiomes to these compositionally different diets. To assess microbiome resilience (recovery potential), all individuals were subsequently reversed to a standard diet for another four weeks. Cloacal microbiomes were collected weekly and characterised through sequencing the v4 region of the 16S rRNA gene using Illumina MiSeq.

Results: Initial microbiomes changed significantly with the diet manipulation but the communities did not differ significantly between the three diet groups (mixed, seed and mealworm), despite multiple diet-specific changes of specific bacterial genera. Reverting birds to the standard diet led to only a partial recovery in gut community compositions. The majority of the bacterial taxa that increased significantly during diet manipulation decreased in relative abundances after reversion to the standard diet; however, bacterial taxa that decreased during the manipulation rarely increased after diet reversal.

Conclusions: The gut microbial response and partial resilience to dietary changes support that gut bacterial communities of *P. major* play a role in accommodating dietary changes experienced by wild avian hosts. This may be a contributing factor to the relaxed association between microbiome composition and bird phylogeny. Our findings further imply that interpretations of wild bird gut microbiome analyses from single-time point sampling, especially for omnivorous species or species that have changing seasonal diets, should be done with caution. The partial community recovery implies that ecologically relevant diet changes (e.g., seasonality and migration) open up gut niches that may be filled by previously abundant microbes or replaced by different symbiont lineages, which has important implications for the integrity and specificity of long-term avian-symbiont associations.

Background

The establishment and maintenance of symbiosis between hosts and their gut microbiota are pivotal in the evolutionary history of animals, where microbial symbionts play a multitude of roles in nutrient management, host development and host immunity [1–3]. Diet and host taxonomy are major drivers of assemblies of gut microbial communities across diverse animal taxa, including insects [4–7], spiders [8], fish [9, 10], frogs [11], mammals [12, 13], and birds [14–18]. In vertebrates, most work to understand the importance of these factors stems from mammals and birds [12, 17, 19, 20], with intriguing differences between the two [19, 20]. Although microbiome composition in mammals is affected by diet [21–26], mammalian gut microbiomes are tightly associated with host phylogeny [19, 20], often accompanied by taxon-specific diets [12]. In contrast, although bird taxonomy, such as host family, is associated with certain microbiome characteristics [15, 16], microbial compositions tend to not be strongly associated
with host phylogeny [16, 20]. In line with this, bird microbiomes also exhibit higher individual variation, conceivably caused by dietary, environmental and social factors [14, 27–31]. It has been proposed that gut adaptations associated with flight (e.g., a smaller gut and consequently a shorter retention time of food in the intestines) may explain the lack of strong phylogenetic signal in avian microbial community compositions [20], which are potentially intensified by compressions or expansions of dietary niches associated with latitudinal and altitudinal migrations [32–34], seasonality [35, 36], and breeding vs. non-breeding seasons [37–39].

A flexible gut microbiome that accommodates changing diets may be important for the evolutionary success of birds, but our knowledge of associations with diet changes remains sparse. For example, although gut microbiomes of migratory birds differ between wintering, stopover and breeding sites, likely due to changes in diet availability [30, 40–44], the impact of diet has not been explicitly investigated in these studies. Intra-specific differences in bird populations from habitats with potentially different food availabilities further support the plastic nature of bird gut microbiomes [28, 31, 45–50], as does the higher individual variation and gut bacterial diversity of omnivorous birds compared to bird species with more specialized diets (e.g., insectivores) [14]. The gap in pinpointing the impact of diet on gut microbiomes has to some extent been filled by recent diet manipulation studies of *Passer domesticus* (house sparrows) [51] and *Parus major* (great tits) [52], in which gut microbiomes respond according to dietary contents. If dietary changes are regular (e.g., seasonality and migration) we would expect gut microbiomes to cyclically change over time depending on temporal changes in food availability [24, 53]. The resilient nature (i.e., the recovery potential after dietary fluctuations) of gut microbiomes has been documented in several mammals [22, 24, 26, 53], but has not been explored in birds.

Here we examine the recovery potential of gut microbiomes of *P. major* after a diet manipulation through characterizing the gut communities using MiSeq amplicon sequencing of the 16S rRNA gene. After feeding birds a standard diet, we fed birds either a mixed diet (including fruits, seeds and mealworms), a seed diet, or a mealworm diet for four weeks (Fig. 1a). Subsequently, we reversed the diet-manipulated birds to the standard diet for four weeks to test the recovery potential of their gut microbial communities and to assess whether full or partial microbial community restoration was achieved (Fig. 1a). If gut communities are flexible and respond to the dietary contents, we expect to find both marked differences between the initial gut microbiomes and the gut microbiomes following diet manipulations, and between the three diet groups due to their differences in dietary contents (Fig. 1b). We further hypothesized that if the gut communities are resilient to dietary changes, gut microbiomes should return to their initial community structure after the reversal period (Fig. 1b). In contrast, if the communities are not resilient, we would expect compositions to be markedly different from initial communities after the diet reversal (Fig. 1b).

**Results**

After analysing and quality filtering of sequences with DADA2 [54] within Qiime2 [55], we acquired a total of 2,965,765 16S rRNA gene sequences (mean ± SE: 20,313 ± 3407) from cloacal swabs from all nine
weeks (initial first week, four weeks of diet manipulation and four weeks of diet reversal) of the experiment. Sequences were identified using the SILVA 132 database [56] and assigned to 2,537 amplicon sequence variances (ASVs) with 100% similarity (Table S1). The majority of the subsequent analyses was only conducted on data from week 1 (initial microbiomes), week 5 (microbiomes after the diet manipulation) and week 9 (reversed microbiomes), since our main interest was to investigate the end result of the diet manipulation and the diet reversal. Samples from week 1 (18 samples after removing samples that failed during sequencing), week 5 (mixed diet: six samples, seed diet: five samples and mealworm diet: five samples) and week 9 (12 samples) had 1,709,192 bacterial 16S rRNA gene sequences (mean ± SE: 37,156.35 ± 5478.41) belonging to 1,290 ASVs (Table S2). Overall, Firmicutes dominated the cloacal microbiomes with 75% of the ASVs, followed by Proteobacteria (10.13%), Tenericutes (10.79%) and Bacteroidetes (1.83%). Only 0.64% of the sequences were unidentified at the phylum level. The initial gut microbiomes of captively raised and wild caught adults fed with identical diets (see Methods) did not differ significantly (permutational multivariate analysis of variance (PERMANOVA)10,000 permutations: $F_{1,16} = 0.9828$, $R^2 = 0.0579$, $p = 0.4175$). Furthermore, the sex of the birds did not have a significant impact on the initial microbial community composition (PERMANOVA10,000 permutations: $F_{1,16} = 1.119$, $R^2 = 0.0653$, $p = 0.2806$), thus we pooled individuals irrespective of the sex in subsequent analyses. Due to high variation in the number of sequences per sample (min: 1,045 sequences and max: 224,835 sequences), we rarefied the original data set (Table S2) to have 1,045 sequences per sample (Table S3) and performed alfa and beta diversity analysis on both the complete and rarefied data sets.

**ASV richness and diversity decrease due to diet manipulation**

Bacterial richness (total number of ASVs) was significantly lower in all diet groups and in the reversed diet gut communities compared to the initial communities (Kruskal-Wallis: $H = 24.89$, df = 4, $p < 0.0001$; Fig. 2a, Table S4). The Shannon's diversity index was also significantly lower in all diet groups compared to the initial gut communities but did not differ between the initial and the reversed diet communities (Kruskal-Wallis: $H = 22.375$, df = 4, $p < 0.0001$; Fig. 2b, Table S4). We found no significant difference in richness and diversity between the three diet groups at the end of the diet manipulation period (Fig. 2, Table S4). Only 27 ASVs were shared among all diet groups and many ASVs tend to be lost or reduced to below detection after the diet manipulation and reversal (Figure S1, Figure S2. The rarefied data set provided similar results for bacterial richness (Kruskal-Wallis: $H = 27.49$, df = 4, $p < 0.0001$; Figure S3a, Table S5) and Shannon's diversity index (Kruskal-Wallis: $H = 22.19$, $p < 0.0001$; Figure S3b, Table S5).

**Bacterial community changed with diet contents**

A PERMANOVA analysis demonstrated a significant impact of diet on the gut bacterial community structure, accounting for 27.1% of the observed variance (PERMANOVA10,000 permutations: $F_{4,41} = 3.805$, $R^2 = 0.271$, $p < 0.001$; Fig. 3a). Pairwise PERMANOVAs showed significant deviation of bacterial
communities of birds on the mixed and the mealworm diets from the initial community structure (Table 1). Gut communities of birds on the seed diet were more variable between bird individuals and did not differ significantly from the initial diet (Fig. 3a, Table 1). The bacterial community structure did not differ significantly between the three diet groups (Table 1). There was a significant difference between the microbiome composition of the initial and the reversed communities (Table 1, Fig. 3a), despite similarities in relative abundances of bacterial phyla and major genera between these two communities (Figs. 3b and 4). We observed significant differences in individual variation (average distance to group centroid) in microbial communities in birds on different diets (Permutation test for multivariate dispersions \(10,000\) permutations: \(F = 4.857, \text{df} = 4, p = 0.0049\)). Pairwise comparisons revealed that gut communities of individuals on the mealworm diet exhibit significantly lower average distances to the group centroid compared to individuals on the initial, the seed and the reversed diets (Figure S4), indicating that the mealworm diet leads to little individual variation in gut microbial communities.

Community-level analyses on the rarefied data set produced similar results, demonstrating the importance of diet on microbial community structures (PERMANOVA with \(10,000\) permutations: \(F_{4,41} = 5.439, R^2 = 0.3467, p = 0.001\); Figure S3c). For the complete data set (including all nine weeks), we observed a week-by-week change in gut communities, with the biggest difference occurring after four weeks (Figure S5). Over the diet manipulation (Figure S5 a, c, e) and diet reversal (Figure S5 b, d, f) periods the microbial communities changed more gradually.

The dominant bacterial phyla changed notably in response to different diets (Fig. 3b) compared to gut communities of birds on the initial diet. Initial communities were dominated by *Firmicutes* (53.45%), followed by *Proteobacteria* (30.62%), *Bacteroidetes* (6.94%) and *Tenericutes* (3.41%). In all three diet treatments, we found a decrease in *Proteobacteria*. In the mixed and the seed diet we found an increase of *Tenericutes* (Fig. 3b). In the mealworm diet treatment, *Firmicutes* vastly dominated the gut microbiota (97.5%) (Fig. 3b). However, the gut communities recovered to initial proportions of bacterial phyla after the diet reversal (Fig. 3b).

The macronutrient content of the four diets (standard, mixed, seed, and mealworm) were not markedly different, except for the mealworm diet (Fig. 3c). The standard and mixed diets were similar in their macronutrient content, except for decreased amounts of proteins and fat in the latter. The seed diet had higher fat and fibre proportions than the mixed and mealworm diet (Fig. 3c), while the mealworm diet had higher proportion of protein and less carbohydrates than in other three diets (Fig. 3c).
Table 1
Results of pair-wise PERMANOVAs with 10,000 permutations comparisons of gut bacterial communities between the initial diet, the three diet manipulation treatments, and the reversed diet (on original data set).

| Diet pairs                  | F statistic | $R^2$   | Adjusted p value |
|-----------------------------|-------------|---------|------------------|
| Initial vs. mixed diet      | 4.894       | 0.1819  | 0.0009*          |
| Initial vs. mealworm diet   | 6.261       | 0.2297  | 0.0019*          |
| Initial vs. seed diet       | 2.127       | 0.0921  | 0.1369           |
| Mixed vs. mealworm diet     | 1.001       | 0.1001  | 1.000            |
| Mixed vs. seed diet         | 1.929       | 0.1765  | 0.3579           |
| Mealworm vs. seed diet      | 3.015       | 0.2737  | 0.0819           |
| Reversed vs. mixed diet     | 4.465       | 0.2182  | 0.0009*          |
| Reversed vs. mealworm diet  | 6.258       | 0.2944  | 0.0019*          |
| Reversed vs. seed diet      | 2.181       | 0.1269  | 0.0189*          |
| Initial vs. reversed diet   | 3.768       | 0.1186  | 0.0009*          |

Differential abundances of bacteria varied according to diet treatment

Relative abundances of the top 25 bacterial genera differed between the initial diet, the three diet manipulation treatments, and the reversed diet (Fig. 4). These 25 genera also accounted for 96.1% of the overall sequences. We found more variation between individual birds in the initial and the reversed bacterial communities than in the mixed and the mealworm diet treatments, whereas the high variation between communities in the seed diet remained (Figs. 4 and S4).

Significantly differentially abundant bacterial genera (using the original data set) between diet treatments were identified using DESeq2 [57]. Bacterial taxa that significantly increased during the diet manipulation also differed between the three diets (Fig. 4, Table S6). Individuals on the mixed diet experienced significant increases in four genera (e.g., Enterococcus, Mangrovibacter, Listeria and Pediococcus) and significant decreases in 20 genera compared to their initial gut microbial communities (Table S6.1). Diet reversal of mixed diet individuals led to a significant increase in 49 genera and a significant reduction in five genera, including Enterococcus and Mangrovibacter (Table S6.2). The seed diet led to significant increases in 13 bacterial genera (e.g., Fusobacterium, Blautia and Ruminococcus), and four genera significantly decreased from initial communities (Table S6.3). Diet reversal of seed diet fed individuals led to significant decreases in 20 genera and an increase in one genus (Jeotgalicoccus) (Table S6.4). Fifteen bacterial genera increased in individuals on the mealworm diet, while five genera decreased significantly (Table S6.5). After the diet reversal of individuals on the mealworm diet two genera decreased significantly (Ureaplasma and Enterococcus) while eight genera increased (Table S6.6).
Intriguingly, only 13% (± 6.5 SE) of the same bacterial genera that decreased during the diet manipulation increased significantly after the diet reversal (Fig. 4). However, around half of the bacterial genera (52.2% ± 23.1 SE) that increased significantly due to the diet manipulation decreased after the diet reversal (Fig. 4, Table S6).

**Discussion**

Exploration of host-gut microbiome dynamics of wild hosts is important to better understand the flexibility, resilience and long-term associations of symbiotic interactions under dietary changes. By examining the flexibility and resilience of wild omnivorous passerine bird gut microbiomes through diet manipulation, our findings document a rapid and significant impact of diet, aligning with other studies on passerine birds [51, 52]. We observed a significant deviation in microbial community structure from the initial gut microbiomes after diet manipulation, but we found no significant differences in gut communities between the three diet groups (Fig. 3). Despite this, there were noteworthy diet-associated trends in the bacterial community variation and differentially abundant bacterial genera. Furthermore, gut microbiomes partially recovered after the diet reversal, implying that the association between *P. major* and their gut communities is somewhat resilient to diet-induced changes.

**P. major gut microbiomes respond to diet changes, but not as flexibly as predicted**

The deviation of gut microbial communities from the initial diet to three different diet groups exemplifies the ability of *P. major* to respond to dietary changes. However, the magnitude of the microbial response varied among diets, suggesting that responses occur according to both the physical (ratios of different diet components) and the macronutrient compositions. Although the macronutrient content of standard, mixed and seed diets did not differ markedly, differences in gut microbial communities indicate that the physical content of diets can impact the gut microbiomes (Figs. 3 and 4). A diet that is both compositionally and macronutritionally different, such as the mealworm diet (Fig. 1a and 3c), has the strongest and most consistent impact on microbiome structure (Figs. 3a and S4).

The differences in individual variation in gut microbiomes between seed and mealworm diets provided some intriguing insights into gut microbiome dynamics (Figure S4). Microbiomes of birds fed with mealworms experienced significantly lower individual variation than birds fed with a standard or a seed diet (Figure S4). The reduced individual variation in insectivores has been demonstrated in wild birds [14] and in another diet manipulation study of *P. major* [52], suggesting that a strictly insectivorous diet (mealworms) might be associated with a specific and more narrow set of gut microbes [58] compared to the seed diet. This is potentially associated with more homogeneous nutrient availability in the mealworm diet [58] compared to the more heterogeneous nutrient availability from multiple types of seeds (e.g., sunflower seeds [59], millet [60], and wheat [61]). Additionally, the taxonomic diversity of initial gut microbiomes of *P. major* in this study differed markedly from the other diet manipulation study [52], where *Proteobacteria* dominated the gut microbiomes, indicating regional differences. However, the observed similarity in the overall community-level response of gut microbiomes for individuals on seed or
insect diets in the two studies, suggest that despite the regional differences of *P. major* gut microbiomes, microbial communities respond in similar ways to similar dietary changes. The microbes responding to these differences may depend on the taxonomic composition of the starting microbiome. This species-level regional variation in gut microbiomes and the flexible nature of passerine microbiomes to dietary changes further support the lack of [16, 20] or weak [62] association between bird gut microbiome structure and host phylogeny.

We also documented significant diet-specific responses of a few bacterial genera in different diet groups (Figs. 1, 3 and Table S6). The seed diet led to a significant increase in relative abundance of the genera *Fusobacterium* (*Fusobacteria*), *Blautia* (*Firmicutes*), and *Ruminococcus* (*Firmicutes*). Some *Fusobacterium* members are animal pathogens [63], but their consistent presence in wild bird guts [2, 14, 64, 65], especially in herbivorous species [64–66], suggests a possible symbiotic role. Bacteria from the genus *Blautia* may facilitate the metabolism of plant secondary metabolites [67, 68], consistent with an increase in the relative abundance of this genus in birds feeding on seeds. *Ruminococcus* bacteria (e.g., the *R. gauvrauii* and *R. gnatus* groups) that increased significantly with seed diet are within the family *Lachnospiraceae* and their possible functions are poorly resolved [69]. However, the *Ruminococcus gnatus* group has been proposed to degrade mucins [69], and the mucin levels in digestive tracts tend to increase on plant-based diets with high fibre contents [70]. These bacteria may thus be opportunistic and utilize the increased amount of mucins.

Birds on the mealworm diet experienced a significant increase in the relative abundance of e.g., *Rombutisia* and *Akkermansia* that are generally presumed to be associated with protein metabolism [71]. Relative increases in the *Firmicutes* orders *Lactobacillales*, *Bacillales* and *Clostridiales* (Table S6.5) in birds on the mealworm diet are consistent with a previous study on insectivorous passerines [14]. Most of these lineages are believed to play roles related to protein fermentation and degradation of toxic by-products from protein metabolism [68, 72–74], thus enabling hosts to sustain an insect diet. Furthermore, a study has demonstrated the prebiotic effects of mealworms on mice gut microbiomes, where mealworm exuviae led to an increase in lactic acid bacteria (e.g., family *Lactobacillaceae*) [75], similar to what we observe in *P. major* fed mealworms (Figs. 1a, 3c and 4). However, understanding how macronutrient content affects wild bird gut microbiome lineages and their associated functional roles, requires further studies with nutritionally markedly different diets and data to decipher their association with gut microbial processes.

Although we cannot rule out that foodborne microbes could impact gut microbial community structures, as has been shown for a small fraction of lactic-acid bacteria from fermented foods in humans [26], we find this unlikely to be a main driver of our results. We did observe new bacterial ASVs in the three diet groups and reversed microbiomes compared to initial communities (Figures S1 and S2), suggesting that new lineages colonise bird gut microbiomes along with dietary changes. However, the most abundant ASVs observed in manipulated gut microbiomes were shared with the initial communities (Table S1 and Figure S2), supporting that major changes in gut microbiomes are accounted for by pre-existing community members. Previous studies on wild birds have demonstrated low bacterial diversity and
community composition differences in the midgut region (stomach and small intestine) compared to the crop microbiota. This suggests that the highly-acidic conditions in the midgut region acts as a barrier for environmental and foodborne bacteria [14, 66, 76, 77]. This is further evident when we look at previously published mealworm gut microbiomes [78], where only a small fraction of the microbiome consists of bacterial genera that we identified in the gut microbiomes of mealworm-fed *P. major*. Furthermore, cloacal swabs appear to adequately capture the microbial diversity of entire digestive tracts [79], suggesting that the observed differences in the cloacal microbial communities are unlikely to be driven by foodborne microbes.

**Wild-bird gut microbiomes are partially resilient to diet-induced changes**

For the first time, we illustrate the resilience of wild bird gut microbiomes to diet changes. However, the recovered bacterial communities did not completely mirror the initial gut communities, despite the recuperated bacterial diversity. There were differences in relative abundances of dominant bacterial genera compared to initial community compositions, and ASV richness was significantly lower in reversed than initial gut communities (Figs. 2, 4 and S1). Intriguingly, we found a consistent pattern that the majority of the taxa that significantly increased in abundance during specific diet treatments decreased again when birds were returned to the standard diet, while taxa that decreased significantly during the diet manipulation rarely recovered following diet reversal (Table S6, Fig. 4).

Partial recovery suggests that ecologically relevant diet changes render available gut niches, that are subsequently filled by either the original microbes filling these niches or that niches are replaced by novel symbiont lineages. Lineages filling particular niches may have the opportunity to flourish by outcompeting bacteria playing similar roles due to functional redundancies inherent in complex gut communities [68, 80, 81]. To understand whether this indeed plays a role, exploration of microbial gene expression between initial and reversed gut bacterial communities will be needed. Our results also demonstrate that *P. major* microbiomes are resilient to natural dietary changes that individuals might experience due to seasonality (e.g., seed-dominated diet during winter and insect-dominated diet in the spring) [36], but these recovery trajectories can be impacted by the competition among bacterial lineages that fill the same functional niches. A partial restoration of gut bacterial communities after dietary changes imply that natural diet changes could lead to microbial symbiont losses (Figure S1), further suggesting that the nature of digestive tract microbiotas in wild birds is plastic to some extent. Hosts may utilize other mechanisms, such as acquiring bacteria from the environment [40] or hosting bacteria with similar metabolic capacities [68, 80] to maintain stability after disruptions of their symbiotic associations. The flexible yet resilient nature of avian gut microbiomes may thus provide an additional level of plasticity for bird hosts to cope with natural (i.e., seasonality, migration, and interspecific competition) [32–39] and anthropogenic (i.e., habitat degradation and invasive species) [82, 83] dietary fluctuations, as predicted by many avian gut microbiome studies [28, 30, 40–49].
Conclusions

Our study documents responses of bird gut microbiomes to diet changes and the resilient nature of gut bacterial communities, supporting the important role of gut bacterial communities in accommodating the dietary breadth of wild bird hosts. The extent of these changes implies an extra level of plasticity in hosts to adjust their dietary niche. This malleable nature of bird gut microbiomes might play a role in observed relaxed association of bird gut microbial communities and host phylogenies. Furthermore, changes that bird gut microbiomes experience due to dietary fluctuations raise concerns about drawing conclusions on wild bird gut microbiomes based on single time points, as they imply that seasonal and regional fluctuations in food sources most likely affect gut microbial compositions that would go unnoticed in the absence of sampling across time and space. The partial recovery of gut microbiomes also suggests that niches that are opened due to diet changes are subsequently either filled by the microbes originally filling these niches or by other symbiont lineages with similar functions or metabolic properties. This suggests that symbiont loss and replacement may be prominent in bird gut microbiomes, with implications for the integrity, specificity, and long-term dynamics of avian-gut microbial associations.

Methods

Bird collection and captive raring

Fifteen *P. major* chicks were collected from nests at the age of 10 days in the surrounding of Ceske Budejovice, Czech Republic on 20th of May 2018 and hand raised in captivity in a breeding room at the Faculty of Sciences, University of South Bohemia. Individuals were fed with a standard diet including 10 mealworms, 2 g insect cake (a bread-like diet made from: Nutribird a21, commercial chicken food (Country’s Best Show 1 crumble), eggs, wheat flower, sugar and sunflower margarine, mixed and baked for ~ 40 min. at ~ 180 °C) and 2tsp of moistened mixed seeds (Living World Premium Mix for Cockatiels & Lovebirds) per day. Furthermore, vitamin water (Acidomid exot®) was provided three times per week. During June and July, the birds were used in a behavioural test (30 min per day), during which they were tested for behavioural responses to plant volatile compounds. The behavioural tests did not affect food intake of the birds and their feeding regime. Five additional adults were captured 20th of September 2018 and kept in captivity for five days on the same standard diet until the diet manipulation started. All birds were kept in individual cages (0.7 × 0.4 × 0.5 m) that were cleaned daily.

The sex of the birds was determined through extracting DNA from 10 µl of blood using the Genomic DNA Mini Kit (Geneaid Biotech Ltd., New Taipei, Taiwan) and conducting PCR using avian-specific sex primers P2 and P8 [84] following an established protocol [85]. Heterogametic females (two bands after PCR) and homogametic males (one band) were identified through visual inspection of agarose gels.

Feeding experiment

Birds (15 captively reared and five wild-caught) were randomly divided into three compositionally different diet groups on the 26th of September 2018 and each group was offered either a mixed diet (six
mealworms + two berries + 2tsp mixed seeds), a mealworm diet (6 g mealworms \([40–50]\) + 4 g insect cake) or a seed diet (3tsp mixed seeds) (Table 2, Fig. 1a). Before the introduction of different diets, a cloacal swab (using a Copan mini swab™) was collected from each individual and stored in 60 µl RNAlater® to investigate the composition of gut microbiota prior to diet manipulation. Birds were under the three different diets for four weeks and cloacal swabs were collected each week. At the end of the fourth week, all birds were reverted to the standard diet for an additional four weeks to investigate whether microbial communities recovered to their initial composition (Fig. 1a). Cloacal swabs were again collected weekly. All the swab samples in RNAlater® were stored at -20 °C until the DNA extractions.
Table 2
Individual IDs of *P. major* and their corresponding rearing conditions, diet treatments and sex.

| Individual ID | Rearing conditions* | Diet treatment | Sex |
|---------------|---------------------|----------------|-----|
| MO1           | Captive reared      | Mixed diet     | F   |
| MO2           | Captive reared      | Mixed diet     | F   |
| MO3           | Captive reared      | Mixed diet     | M   |
| MO4           | Captive reared      | Mixed diet     | F   |
| MO5           | Captive reared      | Mixed diet     | F   |
| MW1           | Wild caught         | Mixed diet     | F   |
| MW2           | Wild caught         | Mixed diet     | F   |
| SO1           | Captive reared      | Seed diet      | M   |
| SO2           | Captive reared      | Seed diet      | F   |
| SO3           | Captive reared      | Seed diet      | F   |
| SO4           | Captive reared      | Seed diet      | F   |
| SO5           | Captive reared      | Seed diet      | F   |
| SW1           | Wild caught         | Seed diet      | F   |
| SW2           | Wild caught         | Seed diet      | M   |
| WO1           | Captive reared      | Mealworm diet  | M   |
| WO2           | Captive reared      | Mealworm diet  | F   |
| WO3           | Captive reared      | Mealworm diet  | F   |
| WO4           | Captive reared      | Mealworm diet  | M   |
| WO5           | Captive reared      | Mealworm diet  | F   |
| WW1           | Wild caught         | Mealworm diet  | M   |

* captive reared: individuals were taken from wild nests and reared in the breeding room; wild caught: individuals were captured as adults a week before the diet manipulation experiment was initiated.

**Nutrient contents of different diets**

Macronutrient contents of 100 g of different diets were analysed at the Eurofins Steins Laboratory (Vejen, Denmark). Amount of crude proteins, fats and fibre were calculated using multiple chemistry protocols at the Steins Laboratory. The amount of Nitrogen-free extract (a proxy for total Carbohydrates) were calculated through subtracting proteins, fats, fibre, ash (i.e. minerals) and water from 100 g. Ash (which
include molecules that are not associated with macronutrients) was measured by burning the whole sample under 550 °C.

**Molecular methods**

DNA from all the swabs were extracted using the Qiagen DNeasy Blood and Tissue® kits (Qiagen, Germany), to sequence the bacterial communities using the V4 region of the 16S rRNA gene. Manufacture’s extraction protocol was followed exactly, except for an extended incubation period (~14 hours) during the lysis step and use of 70 µl of heated (56 °C) AE buffer during the elution step. Initial PCRs were conducted using two primers targeting the V4 region of the 16S rRNA: ‘SB711 (5’-CAAGCAGAAGACGGCATACGAGATTCAGCGTTAGTCAGTCAGCCGGACTACHVGGGTWTCTAAT-3’) and ‘SA504 (5’-AATGATACGGCGACCACTACATAGGGGTWTCTAAT-3’) [cf. 14] and following a well establish PCR protocol for the primer pair [79]. PCR products were visualized on a 2% agarose gel. DNA from positively amplified samples from the initial PCRs along with three negative controls were sent to the Microbial System Molecular Biology lab at the University of Michigan, where the samples were sequenced (using same primers) on the Illumina MiSeq platform.

**Data analysis**

Out of the 176 samples, 169 (96%) were successfully sequenced and the negative controls did not amplify or sequence, demonstrating no contaminations during DNA extractions. Sequences were analysed using DADA2 pipeline [54] within QIIME2 [55]. Bacterial 16S rRNA gene sequence were assigned to taxonomy using SILVA 132 bacterial reference library [56], and archaeal, mitochondrial and chloroplast sequences were removed subsequently. Sequences with 100% similarity were categorized into the same Amplicon sequence variances (ASVs). ASVs with fewer than 10 sequences and samples with fewer than 1,000 sequences were removed from the data analysis. Further analyses were mainly conducted on samples from week one (initial gut microbiota), week five (gut microbiota after diet manipulation) and week nine (gut microbiota after diet reversal), since we were primarily interested in the final responses of gut microbiota to specific diets and their ability to recover after the diet reversal (Table S2). Samples from other weeks were only analysed to investigate the trajectories of changes in bacterial communities and data is presented in Supplementary table 1.

The data was analysed using R 3.5.3 [86, 87]. Shannon diversity index (accounting both bacterial richness and abundances) for samples were calculated using the package vegan [88]. Non-metric multidimensional scaling (NMDS) plots, permutational multivariate analysis of variance (PERMANOVA with 10,000 permutations) and pair-wise PERMANOVA comparisons were conducted using vegan [88] and the wrapper package pairwiseAdonis [89] using Bray Curtis distances between communities to identify statistical differences in gut bacterial communities and to visualize community level differences. We further investigated the individual variation in gut microbial communities through calculating the distance of each community to the centroid of their diet group, using the betadisper function in the package vegan in R [88]. Groups with a smaller average distance to the centroid show less individual
variation. Statistical differences in these distances were tested using permutest (with 10,000 permutations) in the package vegan. We investigated the differential abundances of bacterial genera that increased significantly under different diets (in the non-rarefied data set) using the DESeq2 package [57]. Differential abundances of bacterial genera were investigated separately for each diet group with their appropriate initial and reversed bacterial communities. To assure that the sequencing depth of different samples did not impact the final outcome of our study, we rarefied the original data set (Table S2) using the sample with the lowest number of sequences using the vegan [88] and phyloseq [90] packages (Table S3). Then we performed similar alpha and beta diversity analysis to compare the results of original data set with the rarefied data set.

**Abbreviations**

ASV
amplicon sequence variance

**Declarations**

*Ethic approval and consent to participate*

This research was conducted under the ethical approval of the University of South Bohemia. The wild *P. major* individuals were captured under the permit OOZP/5218/2019/R La from the city council of Ceske Budejovice, Czech Republic and the experiment on animals were conducted under MZP/2019/630/1464 permit issued by the Ministry of Environment, Czech Republic.

*Consent for publication*

Not applicable

*Availability of data and material*

The data generated and analysed are available from the GenBank SRA database (PRJNA548757).

*Competing interests*

There are no competing interests.

*Funding*

We thank the Carlsberg Foundation for financial support through a Distinguished Associate Professor Fellowship to KAJ (CF17-0248). Work of KS was supported by the Grant Agency of the Czech Republic (18-23794Y) and KP by the Grant Agency of the University of South Bohemia (GAJU n.048/2019/P).

*Author contributions*
KHB, MP, KAJ: Developed the idea and designed the experiments
KS, IF, KP: Took care of the birds, carried out the feeding experiment and collected cloacal swabs
KHB: Carried out the molecular work and data analyses
KHB, KS, MP, KAJ: Wrote the manuscript

Acknowledgements

We thank Bela Arora and Xiangyan Su for comments on initial drafts of the manuscript and Radka Pialkova for sexing the birds.

References

1. Flint HJ, Scott KP, Louis P, Duncan SH. The role of the gut microbiota in nutrition and health. Nat Rev Gastro Hepat. 2012;9(10):577–89.
2. Waite DW, Taylor MW. Exploring the avian gut microbiota: current trends and future directions. Front Microbiol. 2015;6:673.
3. Hanning I, Diaz-Sanchez S. The functionality of the gastrointestinal microbiome in non-human animals. Microbiome. 2015;3:51.
4. Sabree ZL, Moran NA. Host-specific assemblages typify gut microbial communities of related insect species. Springerplus. 2014;3:138.
5. Colman DR, Toolson EC, Takacs-Vesbach CD. Do diet and taxonomy influence insect gut bacterial communities? Mol Ecol. 2012;21(20):5124–37.
6. Dietrich C, Kohler T, Brune A. The cockroach origin of the termite gut microbiota: patterns in bacterial community structure reflect major evolutionary events. Appl Environ Microbiol. 2014;80(7):2261–9.
7. Anderson KE, Russell JA, Moreau CS, Kautz S, Sullam KE, Hu Y, Basinger U, Mott BM, Buck N, Wheeler DE. Highly similar microbial communities are shared among related and trophically similar ant species. Mol Ecol. 2012;21(9):2282–96.
8. Kennedy SR, Tsau S, Gillespie R, Krehenwinkel H. Are you what you eat? A highly transient and prey-influenced gut microbiome in the grey house spider *Badumna longinqua*. Mol Ecol. 2020;29(5):1001–15.
9. Liu H, Guo X, Gooneratne R, Lai R, Zeng C, Zhan F, Wang W. The gut microbiome and degradation enzyme activity of wild freshwater fishes influenced by their trophic levels. Sci Rep. 2016;6:24340.
10. Sullam KE, Rubin BE, Dalton CM, Kilham SS, Flecker AS, Russell JA. Divergence across diet, time and populations rules out parallel evolution in the gut microbiomes of Trinidadian guppies. ISME J. 2015;9(7):1508–22.
11. Chang CW, Huang BH, Lin SM, Huang CL, Liao PC. Changes of diet and dominant intestinal microbes in farmland frogs. Bmc Microbiol. 2016;16:33.
12. Muegge BD, Kuczynski J, Knights D, Clemente JC, Gonzalez A, Fontana L, Henrissat B, Knight R, Gordon JI. Diet Drives Convergence in Gut Microbiome Functions Across Mammalian Phylogeny and Within Humans. Science. 2011;332(6032):970–4.

13. Delsuc F, Metcalf JL, Wegener Parfrey L, Song SJ, Gonzalez A, Knight R. Convergence of gut microbiomes in myrmecophagous mammals. Mol Ecol. 2014;23(6):1301–17.

14. Bodawatta KH, Sam K, Jonsson KA, Poulsen M. Comparative Analyses of the Digestive Tract Microbiota of New Guinean Passerine Birds. Front Microbiol. 2018;9:1830.

15. Hird SM, Sanchez C, Carstens BC, Brumfield RT. Comparative Gut Microbiota of 59 Neotropical Bird Species. Front Microbiol. 2015;6:1403.

16. Capunitan DC, Johnson O, Terrill RS, Hird SM. Evolutionary signal in the gut microbiomes of 74 bird species from Equatorial Guinea. Mol Ecol. 2020;29(4):829–47.

17. Grond K, Sandercock BK, Jumpponen A, Zeglin LH. The avian gut microbiota: community, physiology and function in wild birds. J Avian Biol 2018, 49(11).

18. Waite DW, Taylor MW. Characterizing the avian gut microbiota: membership, driving influences, and potential function. Front Microbiol. 2014;5:223.

19. Youngblut ND, Reischer GH, Walters W, Schuster N, Walzer C, Stalder G, Ley RE, Farnleitner AH. Host diet and evolutionary history explain different aspects of gut microbiome diversity among vertebrate clades. Nat Commun. 2019;10(1):2200.

20. Song SJ, Sanders JG, Delsuc F, Metcalf J, Amato K, Taylor MW, Mazel F, Lutz HL, Winker K, Graves GR, et al: Comparative Analyses of Vertebrate Gut Microbiomes Reveal Convergence between Birds and Bats. mBio 2020, 11(1).

21. Amato KR, Leigh SR, Kent A, Mackie RI, Yeoman CJ, Stumpf RM, Wilson BA, Nelson KE, White BA, Garber PA. The Gut Microbiota Appears to Compensate for Seasonal Diet Variation in the Wild Black Howler Monkey (Alouatta pigra). Microb Ecol. 2015;69(2):434–43.

22. Maurice CF, Knowles SCL, Ladau J, Pollard KS, Fenton A, Pedersen AB, Turnbaugh PJ. Marked seasonal variation in the wild mouse gut microbiota. Isme Journal. 2015;9(11):2423–34.

23. Carmody RN, Gerber GK, Luevano JM, Gatti DM, Somes L, Svenson KL, Turnbaugh PJ. Diet Dominates Host Genotype in Shaping the Murine Gut Microbiota. Cell Host Microbe. 2015;17(1):72–84.

24. Ren TT, Boutin S, Humphries MM, Dantzer B, Gorrell JC, Coltman DW, McAdam AG, Wu M. Seasonal, spatial, and maternal effects on gut microbiome in wild red squirrels. Microbiome. 2017;5:163.

25. Hang I, Rinttila T, Zentek J, Kettunen A, Alaja S, Apajalahti J, Harmoinen J, de Vos WM, Spillmann T. Effect of high contents of dietary animal-derived protein or carbohydrates on canine faecal microbiota. Bmc Vet Res. 2012;8:90.

26. David LA, Maurice CF, Carmody RN, Gootenberg DB, Button JE, Wolfe BE, Ling AV, Devlin AS, Varma Y, Fischbach MA, et al. Diet rapidly and reproducibly alters the human gut microbiome. Nature. 2014;505(7484):559–63.
27. Cao J, Hu Y, Liu F, Wang Y, Bi Y, Lv N, Li J, Zhu B, Gao GF. Metagenomic analysis reveals the microbiome and resistome in migratory birds. Microbiome. 2020;8(1):26.
28. Loo WT, Garcia-Loor J, Dudaniec RY, Kleindorfer S, Cavanaugh CM. Host phylogeny, diet, and habitat differentiate the gut microbiomes of Darwin’s finches on Santa Cruz Island. Sci Rep. 2019;9(1):18781.
29. Grond K, Santo Domingo JW, Lanctot RB, Jumpponen A, Bentzen RL, Boldenow ML, Brown SC, Casler B, Cunningham JA, Doll AC, et al. Composition and Drivers of Gut Microbial Communities in Arctic-Breeding Shorebirds. Front Microbiol. 2019;10:2258.
30. Wang W, Wang F, Li L, Wang A, Sharshov K, Druzyaka A, Lancuo Z, Wang S, Shi Y. Characterization of the gut microbiome of black-necked cranes (Grus nigricollis) in six wintering areas in China. Arch Microbiol 2020.
31. Hird SM, Carstens BC, Cardiff SW, Dittmann DL, Brumfield RT. Sampling locality is more detectable than taxonomy or ecology in the gut microbiota of the brood-parasitic Brown-headed Cowbird (Molothrus ater). PeerJ. 2014;2:e231.
32. Araújo PM, Viegas I, Rocha AD, Villegas A, Jones JG, Mendonça L, Ramos JA, Masero JA, Alves JA. Understanding how birds rebuild fat stores during migration: insights from an experimental study. Sci Rep-Uk. 2019;9:10065.
33. Pierce BJ, McWilliams SR. Seasonal Changes in Composition of Lipid Stores in Migratory Birds: Causes and Consequences. The Condor. 2005;107(2):269–79.
34. Boyle WA, Conway CJ, Bronstein JL. Why do some, but not all, tropical birds migrate? A comparative study of diet breadth and fruit preference. Evol Ecol. 2010;25:219–36.
35. Witmer MC. Annual Diet of Cedar Waxwings Based on U.S. Biological Survey Records (1885–1950) Compared to Diet of American Robins: Contrasts in Dietary Patterns and Natural History. Auk. 1996;113(2):414–30.
36. Vel’ky M, Kanuch P, Kristin A. Food composition of wintering great tits (Parus major): habitat and seasonal aspects. Journal of Vertebrate Biology. 2011;60(3):228–36.
37. Bakour S, Moulaï R. Dietary analysis across breeding seasons of Eleonora’s Falcon Falco eleonorae on the western coast of Algeria. Ostrich. 2019;90(1):63–72.
38. Westerberg K, Brown R, Eagle G, Votier SC. Intra-population variation in the diet of an avian top predator: generalist and specialist foraging in Great Black-backed Gulls Larus marinus. Bird Study. 2019;66(3):390–7.
39. Ankney CD, Scott DM. Changes in Nutrient Reserves and Diet of Breeding Brown-Headed Cowbirds. Auk. 1980;97(4):684–96.
40. Lewis WB, Moore FR, Wang SA. Characterization of the gut microbiota of migratory passerines during stopover along the northern coast of the Gulf of Mexico. J Avian Biol. 2016;47(5):659–68.
41. Wu Y, Yang Y, Cao L, Yin H, Xu M, Wang Z, Liu Y, Wang X, Deng Y. Habitat environments impacted the gut microbiome of long-distance migratory swan geese but central species conserved. Sci Rep. 2018;8(1):13314.
42. Risely A, Waite DW, Ujvari B, Hoye BJ, Klaassen M. Active migration is associated with specific and consistent changes to gut microbiota in Calidris shorebirds. J Anim Ecol. 2018;87(2):428–37.
43. Dong Y, Xiang X, Zhao G, Song Y, Zhou L. Variations in gut bacterial communities of hooded crane (Grus monacha) over spatial-temporal scales. Peerj. 2019;7:e7045.
44. Zhang F, Xiang X, Dong Y, Yan S, Song Y, Zhou L. Significant Differences in the Gut Bacterial Communities of Hooded Crane (Grus monacha) in Different Seasons at a Stopover Site on the Flyway. Animals. 2020;10(4):701.
45. Liu G, Gong Z, Li Q. Variations in gut bacterial communities between lesser white-fronted geese wintering at Caizi and Shengjin lakes in China. Microbiologyopen. 2020;00:e1037.
46. Juan PAS, Hendershot JN, Daily GC, Fukami T. Land-use change has host-specific influences on avian gut microbiomes. The ISME Journal. 2020;14:318–21.
47. Phillips JN, Berlow M, Derryberry EP. The Effects of Landscape Urbanization on the Gut Microbiome: An Exploration Into the Gut of Urban and Rural White-Crowned Sparrows. Frontiers in Ecology and Evolution 2018, 6.
48. Teyssier A, Rouffae LO, Saleh Hudin N, Strubbe D, Matthysen E, Lens L, White J. Inside the guts of the city: Urban-induced alterations of the gut microbiota in a wild passerine. Sci Total Environ. 2018;612:1276–86.
49. Gillingham MAF, Bechet A, Cezilly F, Wilhelm K, Rendon-Martos M, Borghesi F, Nissardi S, Baccetti N, Azafzaf H, Menke S, et al. Offspring Microbiomes Differ Across Breeding Sites in a Panmictic Species. Front Microbiol. 2019;10:35.
50. Murray MH, Lankau EW, Kidd AD, Welch CN, Ellison T, Adams HC, Lipp EK, Hernandez SM. Gut microbiome shifts with urbanization and potentially facilitates a zoonotic pathogen in a wading bird. Plos One. 2020;15(3):e0220926.
51. Teyssier A, Matthysen E, Hudin NS, de Neve L, White J, Lens L: Diet contributes to urban-induced alterations in gut microbiota: experimental evidence from a wild passerine. Proc Biol Sci 2020, 287(1920):20192182.
52. Davidson G, Wiley N, Cooke AC, Johnson CN, Fouhy F, Reichert MS, Hera I, Crane JMS, Kulahtci IG, Ross RP, et al: Diet-induced changes to host gut microbiota is linked to foraging innovation in a wild bird In. bioRxiv; 2020.
53. Hicks AL, Lee KJ, Couto-Rodriguez M, Patel J, Sinha R, Guo C, Olson SH, Seimon A, Seimon TA, Ondzie AU, et al. Gut microbiomes of wild great apes fluctuate seasonally in response to diet. Nat Commun. 2018;9(1):1786.
54. Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJ, Holmes SP. DADA2: High-resolution sample inference from Illumina amplicon data. Nat Methods. 2016;13(7):581–3.
55. Bolyen E, Rideout JR, Dillon MR, Bokulich NA, Abnet CC, Al-Ghalith GA, Alexander H, Alm EJ, Arumugam M, Asnicar F, et al. Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. Nat Biotechnol. 2019;37(8):852–7.
56. Quast C, Pruesse E, Yilmaz P, Gerken J, Schweer T, Yarza P, Peplies J, Glockner FO. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Res. 2013;41(D1):D590–6.

57. Love MI, Huber W, Anders S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 2014;15(12):550.

58. Finke MD. Complete Nutrient Composition of Commercially Raised Invertebrates Used as Food for Insectivores. Zoo Biology. 2002;21:269–85.

59. Muhammad Anjum F, Nadeem M, Issa Khan M, Hussain S. Nutritional and therapeutic potential of sunflower seeds. British Food Journal. 2012;114(4):544–52.

60. Barbeau WE, Hilu KW. Protein, calcium, iron, and amino acid content of selected wild and domesticated cultivars of finger millet. Plant Foods Hum Nutr. 1993;43(2):97–104.

61. Escarnot E, Jacquemin J, Agneessens R, Paquot M. Comparative study of the content and profiles of macronutrients in spelt and wheat, a review. Biotechnology Agronomy Society Environment. 2012;16(2):243–56.

62. Trevelline BK, Sosa J, Hartup BK, Kohl KD: A bird’s-eye view of phylosymbiosis: weak signatures of phylosymbiosis among all 15 species of cranes. Proc Biol Sci 2020, 287(1923):20192988.

63. Hofstad T: The Genus Fusobacterium. In: The Prokaryotes - A handbook on the biology of bacteria. Edited by Dworkin M, Falkow S, Rosenberg E, Schleifer KH, Stackebrandt E, vol. 2. New York: Springer-Verlag; 2006: 1016–1027.

64. Bennett DC, Tun HM, Kim JE, Leung FC, Cheng KM. Characterization of cecal microbiota of the emu (Dromaius novaehollandiae). Vet Microbiol. 2013;166(1–2):304–10.

65. Wang W, Zheng SS, Li LX, Yang YS, Liu YB, Wang AZ, Sharshov K, Li Y. Comparative metagenomics of the gut microbiota in wild greylag geese (Anser anser) and ruddy shelducks (Tadorna ferruginea). Microbiologyopen. 2019;8(5):e00725.

66. Drovetski SV, O'Mahoney M, Ransome EJ, Matterson KO, Lim HC, Chesser RT, Graves GR. Spatial Organization of the Gastrointestinal Microbiota in Urban Canada Geese. Sci Rep-Uk. 2018;8:3713.

67. Burapan S, Kim M, Han J. Demethylation of Polymethoxyflavones by Human Gut Bacterium, Blautia sp MRG-PMF1. J Agr Food Chem. 2017;65(8):1620–9.

68. Rowland I, Gibson G, Heinken A, Scott K, Swann J, Thiele I, Tuohy K. Gut microbiota functions: metabolism of nutrients and other food components. Eur J Nutr. 2018;57(1):1–24.

69. La Reau AJ, Suen G. The Ruminococci: key symbionts of the gut ecosystem. J Microbiol. 2018;56(3):199–208.

70. Montagne L, Piel C, Lalles JP. Effect of diet on mucin kinetics and composition: nutrition and health implications. Nutr Rev. 2004;62(3):105–14.

71. Amaretti A, Gozzoli C, Simone M, Raimondi S, Righini L, Perez-Brocal V, Garcia-Lopez R, Moya A, Rossi M. Profiling of Protein Degraders in Cultures of Human Gut Microbiota. Front Microbiol. 2019;10:2614.
72. Leblanc DJ: Enterococcus. In: *The Prokaryotes - A handbook on the biology of bacteria*. Edited by Dworkin M, Falkow S, Rosenberg E, Schleifer KH, Stackebrandt E, vol. 4, 3 edn. New York: Springer-Verlag; 2006: 175–204.

73. Holzapfel WH, Franz CMAP, Ludwig W, Back W, Dicks LMT: The Genera Pediococcus and Tetragenococcus. In: *The Prokaryotes - A handbook on the biology of bacteria*. Edited by Dworkin M, Falkow S, Rosenberg E, Schleifer KH, Stackebrandt E, vol. 4, 3 edn. New York: Springer-Verlag; 2006: 229–266.

74. Teuber M, Geis A: The Genus Lactococcus. In: *The Prokaryotes - A handbook on the biology of bacteria*. Edited by Dworkin M, Falkow S, Rosenberg E, Schleifer KH, Stackebrandt E, vol. 4, 3 edn. New York: Springer-Verlag; 2006: 205–228.

75. Kwon GT, Yuk HG, Lee SJ, Chung YH, Jang HS, Yoo JS, Cho KH, Kong H, Shin D. Mealworm larvae (*Tenebrio molitor* L.) exuviae as a novel prebiotic material for BALB/c mouse gut microbiota. Food Sci Biotechnol. 2020;29(4):531–7.

76. Kropackova L, Pechmanova H, Vinkler M, Svobodova J, Velova H, Tesicky M, Martin JF, Kreisinger J. Variation between the oral and faecal microbiota in a free-living passerine bird, the great tit (*Parus major*). Plos One. 2017;12(6):e0179945.

77. Drovettski SV, O'Mahoney MJV, Matterson KO, Schmidt BK, Grant BR. Distinct microbiotas of anatomical gut regions display idiosyncratic seasonal variation in an avian folivore. BMC Animal Microbiome. 2019;1:2.

78. Przemieniecki SW, Kosewska A, Ciesielski S, Kosewska O. Changes in the gut microbiome and enzymatic profile of *Tenebrio molitor* larvae biodegrading cellulose, polyethylene and polystyrene waste. Environ Pollut. 2020;256:113265.

79. Bodawatta KH, Puzejova K, Sam K, Poulsen M, Jonsson KA. Cloacal swabs and alcohol bird specimens are good proxies for compositional analyses of gut microbial communities of Great tits (*Parus major*). BMC Animal Microbiome. 2020;2:9.

80. Moya A, Ferrer M. Functional Redundancy-Induced Stability of Gut Microbiota Subjected to Disturbance. Trends Microbiol. 2016;24(5):402–13.

81. Trosvik P, de Muinck EJ. Ecology of bacteria in the human gastrointestinal tract-identification of keystone and foundation taxa. Microbiome. 2015;3:44.

82. Isaksson C, Anderson S. Carotenoid diet and nestling provisioning in urban and rural great tits *Parus major*. J Avian Biol. 2007;38(5):564–72.

83. Lambertucci SA, Trejo A, Di Martino S, Sánchez-Zapata JA, Donázar JA, Hiraldo F. Spatial and temporal patterns in the diet of the Andean condor: ecological replacement of native fauna by exotic species. Anim Conserv. 2009;12(4):338–45.

84. Griffiths R, Double MC, Orr K, Dawson R.JG. A DNA test to sex most birds. Mol Ecol. 1998;7(8):1071–5.

85. Lezalova-Pialkova R. Molecular evidence for extra-pair paternity and intraspecific brood parasitism in the Black-headed Gull. J Ornithol. 2011;152(2):291–5.
86. **R: A language and environment for statistical computing.** [https://www.R-project.org/](https://www.R-project.org/) – Accessed January 2020.

87. RStudio. **Integrated Development for R** [http://www.rstudio.com/].

88. vegan. **Community Ecology Package. R package version 2.5-4** [https://CRAN.R-project.org/package=vegan](https://CRAN.R-project.org/package=vegan) - Accessed January 2020.

89. pairwiseAdonis. **Pairwise multilevel comparison using adonis**
https://github.com/pmartinezarbizu/pairwiseAdonis - Accessed January 2020.

90. McMurdie PJ, Holmes S. **phyloseq: An R Package for Reproducible Interactive Analysis and Graphics of Microbiome Census Data.** *Plos One* 2013, 8(4).

**Figures**
Figure 1

a. Contents of standard, mixed, seed and mealworm diets, with a schematic timeline of the experiment. b. Predicted flexible (toward three diet groups) and resilient (recovery after the diet reversal) microbiome response to dietary changes in Parus major. Filled grey circles represent recovery of gut microbiomes if the communities are resilient, while open grey circles represent potential outcome of non-resilient microbiomes.
Figure 2

Mean ASV richness (a) and Shannon's diversity index (b) of gut microbial communities in initial, after the diet manipulation and after the diet reversal. Letters above each boxplot represent the pairwise differences between different groups (Dunn's post-hoc test). Different letters indicate significant differences.
Figure 3

a. Non-Metric Multidimensional Scaling (NMDS) plot of bacterial communities for initial, mixed, mealworm, seed, and reversed diets (ellipses indicate 95% CI; stress = 0.166). b. Relative abundances of bacterial phyla in gut microbiomes under different diets. c. Relative abundance of macronutrients in 100g of each diet.
Figure 4

Most abundant bacterial genera changed on new diets but partially recover after diet reversal. Relative abundances of the 25 most common bacterial genera in the cloacal swabs of P. major on the initial diet, after diet manipulation and after diet reversal are shown in the bar graphs. Unclassified genera are indicated with a “U”, following the closest taxonomic level classification (i.e. family or order). Individual birds are represented with their name codes on the x-axis. Top ten significantly differentially abundant
genera (divided between comparison groups) are given between panels. Positive log2-fold values represent increase of certain genera in bottom panel compared to the top panel. Empty columns indicate the samples that did not sequence or fail the quality filtering step.

**Supplementary Files**

This is a list of supplementary files associated with this preprint. Click to download.

- FigureS5.pdf
- figureS4.pdf
- FigureS3.pdf
- FigureS2.pdf
- FigureS1.pdf
- TableS6.xlsx
- TableS3.xlsx
- TableS2.xlsx
- TableS1.xlsx