Commensal bacteria and essential amino acids control food choice behavior and reproduction

Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Ana Patrícia Francisco, Gabriela Tondolo Fioreze, Margarida Anjos, Célia Baltazar, Ana Paula Elias, Pavel M. Itskov, Matthew D. W. Piper, Carlos Ribeiro

Abstract

Choosing the right nutrients to consume is essential to health and wellbeing across species. However, the factors that influence these decisions are poorly understood. This is particularly true for dietary proteins, which are important determinants of lifespan and reproduction. We show that in Drosophila melanogaster, essential amino acids (eAAs) and the concerted action of the commensal bacteria Acetobacter pomorum and Lactobacilli are critical modulators of food choice. Using a chemically defined diet, we show that the absence of any single eAA from the diet is sufficient to elicit specific appetites for amino acid (AA)-rich food. Furthermore, commensal bacteria buffer the animal from the lack of dietary eAAs: both increased yeast appetite and decreased reproduction induced by eAA deprivation are rescued by the presence of commensals. Surprisingly, these effects do not seem to be due to changes in AA titers, suggesting that gut bacteria act through a different mechanism to change behavior and reproduction. Thus, eAAs and commensal bacteria are potent modulators of feeding decisions and reproductive output. This demonstrates how the interaction of specific nutrients with the microbiome can shape behavioral decisions and life history traits.

Author summary

What animals, including humans, choose to eat has a tremendous impact on health and wellbeing. Though intake of dietary proteins and amino acids is essential for animals, excessive consumption of these nutrients is known to have detrimental effects. Many animals, therefore, execute precise control over the intake of these key nutrients. However, the factors controlling protein appetite are poorly understood. Here, we show that in the vinegar fly Drosophila melanogaster, essential amino acids and gut bacteria are key modulators of protein appetite. Lack of any one essential amino acid from the diet produces a strong and specific appetite for proteinaceous or amino acid–rich food. However, flies with an appropriate microbiome do not develop this protein appetite. Specifically, two gut
bacteria species, *Acetobacter pomorum* and *Lactobacilli*, work together to suppress protein appetite. Furthermore, we show that flies lacking dietary essential amino acids have reduced reproductive output, an effect which is also rescued by gut bacteria. Finally, based on metabolite measurements, we propose that the influence of bacteria on host physiology and behavior is not mediated by changing amino acid levels. Our study demonstrates how the interaction of specific nutrients with the microbiome can shape behavior and animal fitness and suggests that they do so through a novel mechanism.

Introduction

The appropriate intake of nutrients has a major beneficial impact on health and lifespan [1–3]. The level of dietary protein intake has emerged as a key determinant of overall mortality, fecundity, and lifespan in species ranging from humans [4] to mice [5] and *Drosophila* [6–9]. Accordingly, animals, including humans, are able to direct food choice in order to tightly control protein intake [2,3,10–13]. Despite the striking physiological and behavioral impact of nutritional proteins, how animals direct feeding decisions to ensure protein homeostasis is not understood. A major obstacle in identifying the rules governing food choice is the nutritional complexity of natural foods, which hinders the discovery of the nutritional variables controlling feeding decisions.

In *Drosophila melanogaster*, yeast is thought to cover the protein as well as most other non-caloric nutritional requirements [7]. In adult females, yeast appetite is driven by two main internal states: mating and lack of yeast [12–15]. The molecular and circuit mechanisms leading to an increase in yeast appetite upon mating have been extensively characterized. During copulation, the male-derived Sex Peptide is transferred to the female and acts on the neuronal Sex Peptide Receptor, leading to the silencing of a postmating neuronal circuit, consisting of SPSN/SAG/ociptopamine components, which projects to the central brain to change feeding preference from sugar to yeast [12,14,16]. Besides mating, the other known determinant of protein intake is removal of yeast from the diet, which leads to a strong compensatory appetite for yeast [12]. The mechanisms underlying this homeostatic change in appetite are less well understood. This is partially due to the fact that yeast is a complex food containing different nutrients, including amino acids (AAs), carbohydrates, vitamins, and sterols [17,18]. However, it is still unknown which nutrient(s), when absent, triggers flies to ingest yeast. Identifying the mechanisms controlling protein homeostasis in *Drosophila* requires untangling this nutritional complexity.

The interaction of microbiota with ingested nutrients has emerged as a major determinant of health and disease, including obesity [19–24]. Commensal bacteria have also been proposed to affect a wide array of brain functions [25–29] ranging from bulk food intake [30] to anxiety [31–33], neurodevelopmental disorders [34], and social behavior [35]. Despite being an intense field of research, the importance of microbe–nutrient interactions in influencing behavior remains poorly understood. In vertebrates, this task is especially challenging given the complexity of their microbiota and the large set of nutritional parameters that could influence their function. Furthermore, in the context of nutrition, research on microbiota has mainly focused on their role in carbohydrate homeostasis [21,36]. More recently, however, the importance of commensal bacteria in controlling growth [37–39] and in protecting children from malnutrition symptoms [40] indicate that the microbiome could also play a pivotal role in protein homeostasis. However, the importance of commensals in protein homeostasis and in directing food choices has not been directly addressed.
In this study we show that yeast and AA preferences are driven by dietary deprivation from essential AAs (eAAs). While the absence of a single eAA is sufficient to induce a potent yeast appetite, removal of other important nutrients from the diet does not lead to an increase in yeast preference. The fly, however, is not specialized in detecting the identity of the missing AA. Flies rendered auxotrophic for a nonessential AA (neAA) display a strong yeast appetite upon deprivation of this artificially engineered eAA. Furthermore, we show that the presence of commensal bacteria abolishes the yeast appetite and the strong decrease in egg laying induced by the removal of eAAs. Commensal bacteria also have a strong phagostimulatory effect that is likely to aid the replenishment of gut bacteria. Using gnotobiotic animals, we test the hypothesis that commensals alter feeding decisions by providing eAAs to the host. We find, however, no evidence that the decrease in eAA levels induced by dietary deprivation is ameliorated by the presence of commensals, suggesting that they may use a different mechanism to alter food choice. Our study identifies two key components driving food choice in Drosophila: eAAs and the gut bacteria species Acetobacter pomorum and Lactobacilli. Furthermore, we provide initial insights into their action on the host, highlighting the power of Drosophila for identifying key determinants underlying complex nutritional–microbial–behavioral interactions.

Results

Essential amino acids, but not other nutrients, control yeast and amino acid preference

Yeast deprivation leads to a strong compensatory appetite for yeast [12] (Fig 1B). Given the complexity of this resource, it has not been possible to identify the nutrients that, when absent, trigger flies to ingest yeast. To answer this question, we decided to manipulate each nutrient present in yeast independently using a chemically defined (holistic) diet [7] (Fig 1A) and study their effects on feeding decisions using a two-color food choice assay [12,41]. The holistic medium is able to suppress yeast appetite to the same extent as a yeast-based medium (Fig 1B and 1C), supporting the idea that it provides the necessary nutrients to support adult behavior [7]. Removal of AAs from the holistic medium induced a potent yeast appetite, indistinguishable from that observed upon yeast deprivation (Fig 1B and 1C). Removing folic acid, metals, nucleic acids, lipids, sterols, or vitamins, however, did not lead to a significant increase in yeast appetite (Fig 1C). This effect stands in strong contrast to the clear effects on lifespan and egg production of removing these nutrients [7].

To identify the nutrients that flies select when deprived of AAs, we used the holistic diet in the choice paradigm. We gave the flies a choice between a holistic base diet containing sucrose without AAs and one containing AAs but no carbohydrates. AA deprivation shifted flies’ preference from the sucrose-containing option towards the AA-containing option (Fig 1D). This shift in preference was specific to AAs since it was abolished by removal of AAs from the choice medium, whereas removing any other class of nutrients left the shift intact (Fig 1D and S1A Fig). These results suggest that similarly to yeast deprivation [42], upon AA deprivation, flies specifically select a diet containing AAs. In our paradigm, the decision to switch from eating sucrose to yeast or AAs is therefore guided by the absence of AAs, while the absence of other physiologically important dietary nutrients does not lead to an increase in yeast or AA appetite.

AAs can be broadly classified as either essential or nonessential. neAAs can be synthesized by the animal, allowing animals to be largely independent from dietary uptake of these important building blocks [43]. It is currently unclear whether animals sense these two types of AAs
differently and if they have different effects on nutrient choice [44,45]. We tested this by manipulating AAs of each type independently. Removal of all eAAs from the diet induced a yeast (Fig 1E) and AA appetite (Fig 1F) that were indistinguishable from that observed upon removal of all AAs. The complete removal of neAAs, however, had no effect on nutrient choice (Fig 1E and 1F). Given that we adjust the total level of AAs to maintain a constant amount of nitrogen in the diet, these results also show that it is the identity of the AAs and not the nitrogen level in the diet that leads to changes in food choice. Intriguingly, AA deprivation induced...
a preference for both eAAs and neAAs, suggesting that the phagostimulatory power of AAs is not correlated with their nutritional importance, as indicated by previous studies [42] (S1B Fig). Taken together, these data strongly indicate that eAAs are specific mediators of protein and AA appetite and highlight the ability of animals to efficiently buffer the absence of neAAs.

The absence of any single essential amino acid can induce a potent yeast appetite

Behavioral [45], physiological [9], and molecular studies [46] have suggested that different single AAs can vary widely in their potency to suppress protein appetite and to activate nutrient-sensitive pathways. We therefore took advantage of the unique possibility to manipulate single dietary AAs afforded by the holidic diet to remove every eAA individually from the diet and test the effect on food choice. Strikingly, removal of any eAA was sufficient to induce a clear increase in yeast choice (Fig 2A). The extent to which they did so did not differ, suggesting that each eAA has a similar impact on food choice. Furthermore, we quantified the effect of removing specific AAs from the diet on the intake of sucrose and yeast extract using a method to quantify food intake [47] (the capillary feeder [CAFE] assay; Fig 2B). Consistent with our results using the two-color assay, removal of either all AAs or single eAAs (arginine or valine) led to a specific increase in yeast extract intake without affecting carbohydrate intake (Fig 2B). In agreement with previous reports [16], these data indicate that the changes in food choice induced by AA deprivation in the two-color choice assay are due to an increase in yeast appetite and not to a decrease in sucrose intake. They further indicate that single eAAs are potent...
and specific nutritional modifers of protein intake, highlighting their unique importance in controlling food choice.

Loss of peripheral synthesis of neAAs increases yeast appetite

Animals can synthesize neAA in order to compensate for their absence from the diet. For example, tyrosine (Tyr) can be synthesized from phenylalanine (Phe) through the action of phenylalanine hydroxylase, which in Drosophila is encoded by the Henna gene (Fig 3A) [48]. In humans, mutations in phenylalanine hydroxylase cause phenylketonuria, the most common

Fig 3. Loss of peripheral synthesis of nonessential amino acids (neAAs) increases yeast appetite. (A) Schematic depicting the biosynthesis of tyrosine (Tyr) in Drosophila. (B) Feeding preference of control and Henna whole-animal knockdown flies upon removal of all amino acids (AAs), all neAAs, or all neAAs with 1x and 2x Tyr added back. (C) Feeding preference of control and Henna fat body knockdown flies upon removal of either all neAAs or all AAs. (B and C) Circles represent yeast preference in single assays, with a line representing the median and whiskers representing the interquartile range. n = 15–20. Significance was tested using the Kruskal–Wallis test followed by Dunn’s multiple comparison test. Not significant (ns) p > 0.05, * p < 0.05, ** p < 0.01, *** p < 0.001. Underlying data used in this Fig are provided in S1 Data.

https://doi.org/10.1371/journal.pbio.2000862.g003
metabolic disease [49,50]. Patients with phenylketonuria suffer from elevated Phe and low Tyr titters, leading to severe complications including neurological and behavioral symptoms [51]. Strict adherence to a diet low in Phe and high in Tyr allows patients to lead an asymptomatic life, highlighting the impact of dietary AAs on human health [52].

We mimicked the genetic lesion leading to phenylketonuria by knocking down the Henna gene ubiquitously (S2A Fig), thus transforming Tyr from a neAA to an eAA. This allowed us to test if the capacity to homeostatically trigger changes in food choice is related to the specific identities of the ten eAAs or if it can be driven by low levels of any AA. While removal of dietary neAAs in control animals did not lead to the induction of a yeast appetite, the same dietary manipulation in Henna knockdown animals led to a strong yeast appetite (Fig 3B and S2B Fig). This increased yeast appetite was indistinguishable from that observed upon removal of all AAs. Supplementing the diet lacking neAAs with Tyr suppressed the preference of flies for yeast in a dose-dependent manner, indicating that the phenotype was specifically due to an acute lack of Tyr and not to other detrimental effects of our genetic manipulation (Fig 3B). Importantly, the addition of proline, a neAA which is not synthesized by phenylalanine hydroxylase, did not suppress the Henna phenotype, further emphasizing the specificity of the metabolic manipulation (S2B Fig). These results strongly suggest that flies can detect the absence of any limiting AA independent of their specific identity (eAA versus neAA).

In mammals, neAAs are mainly synthesized in the liver [53,54], and it is thought that in insects, the fat body fulfills a similar role [55–57]. We tested the importance of the fat body in guiding nutrient choice by interfering with the ability of this organ to synthesize Tyr. Knockdown of Henna using a fat body driver Cg-Gal4 rendered the animal sensitive to the absence of dietary neAAs, with induction of a strong yeast appetite (Fig 3C). Henna knockdown in neurons or trachea, in contrast, did not change the behavioral sensitivity of flies to removal of all neAAs (S2C Fig), indicating that the effect observed with the fat body manipulation is tissue specific. However, Cg-Gal4 has also been shown to drive expression in hemocytes [58]. It is thus possible that this cell type also contributes to Tyr synthesis and the observed behavioral phenotype. Taken together, these data further demonstrate that AAs, be they dietary or endogenously synthesized, are able to control yeast appetite. Furthermore, our data indicate that biosynthetically active organs are important regulators of food choice, suggesting that genetic metabolic conditions such as phenylketonuria could have effects on aspects of behavior such as nutrient-specific appetites.

Commensals and amino acids control food choice

Mounting evidence indicate that commensal bacteria are important determinants of how nutrients are utilized [59,60]. As such, they modulate a large set of nutrient-sensitive traits. However, whether commensals influence the selection of specific dietary nutrients is currently unknown. We therefore set out to test the effect of commensals on nutrient choice in Drosophila. Importantly, the flies used in our experiments had a very low baseline gut microbe load (S3 Fig). This is likely due to the use of sterile media and the fact that upon serial passage to new food, adult flies lose a large part of their microbiota [61]. To test the effect of the microbiota on behavioral protein homeostasis, we removed one eAA (histidine [His]) from the holidic diet to increase the flies’ preference for yeast and examined if they would show alterations in food choice when treated with a controlled microbiota (Fig 4A) (pure culture of five Drosophila gut bacteria strains: Lactobacillus plantarum WTE [62], Lactobacillus brevis FW [62], Acetobacter pomorum [62], Commensalibacter intestini A911T [62], and Enterococcus faecalis [63]). Strikingly, in contrast to control flies, bacteria-treated flies did not show an increased yeast appetite upon
Fig 4. Commensal bacteria control food choice and egg laying. (A) Diagram depicting the strategy used to reconstitute the commensal population in the fly. (B) Yeast preference of animals kept on holidic diet with or without histidine (His). One group was not pretreated with the five commensals (untreated), the other was pretreated with the commensals (five bacteria), and the third was pretreated with inactivated commensals (inactivated five bacteria). (C) Yeast preference of animals kept on holidic diet, holidic diet without essential amino acids (eAAs), or holidic diet without eAAs pretreated with the commensal mix. (D) Yeast preference of animals kept on holidic diet with or without His and pretreated with varying concentrations of the commensal mix. (E) Yeast preference of Henna knockdown animals (tubulin > HennaIR1) kept on holidic diet lacking nonessential amino acids (neAAs), with or without pretreatment with the commensal mix, and holidic diet lacking neAAs with 2x tyrosine (Tyr) added back. (F) Number of eggs laid per female in 24 h of animals kept on holidic diet, holidic diet with all amino acids except His, isoleucine (Ile), or valine (Val), and with or without pretreatment with commensals. Black filled circles represent complete holidic medium or pretreatment with the bacteria mix. Open black circles represent flies that were not pretreated with bacteria mix. Amino acid (AA) deprivation is indicated as –His, –Ile or –Val. Data are pooled from two different rounds of experiments performed independently on different days. n = 39–40. (B, C, and E) Circles represent yeast preference in single assays, with a line representing the median and whiskers representing the interquartile range. Filled circles represent assays in which flies had been pretreated with commensals. n = 20–30. (D) Points represent median yeast preference and error bars represent the interquartile range. n = 15. (F) Circles represent eggs laid in single assays, with the line representing the mean. (B–E) Significance was tested using the Kruskal–Wallis test followed by Dunn’s multiple comparison test and in (F) using the one-way analysis of variance test followed by Bonferroni’s multiple comparison test (B–F) Not significant (ns) p > 0.05, *p < 0.05, **p < 0.01, ***p < 0.001. Underlying data used in this Fig are provided in S1 Data.
His removal (Fig 4B). The effect of commensals was not limited to His but suppressed yeast appetite upon removal of any of the ten tested eAAs (S4 Fig). The effect of commensals on food choice was so strong that the flies with a reconstituted microbiome were even buffered against the removal of all eAAs from the diet (Fig 4C).

To test if the bacteria were merely acting as food or if they needed to be metabolically active, we tested the effect of inactivated bacteria on food choice. Inactivation abolished the ability of the bacteria to alter feeding decisions, suggesting that their activity is essential to drive the change in behavior (Fig 4B). The behavioral effect was specific to commensal bacteria, as it was not observed when a non-commensal bacterium (Escherichia coli) was used in the experiments (S5A Fig). That bacteria do not simply act as food is further supported by the fact that even when diluted, they are able to lower the preference for yeast (Fig 4D). Furthermore, to show that the effect of the bacteria is not confined to flies fed on the synthetic diet, we pretreated flies with decreasing amounts of yeast to induce a yeast appetite. As expected, flies fed with decreasing amounts of yeast showed an increase in yeast preference (S5B Fig). Similarly to the effect on AA-deprived flies, bacteria pretreatment reduced the yeast appetite of yeast-deprived flies when compared to non-pretreated controls (S5B Fig). This indicates that the effect of commensal bacteria on food choice is generalizable to ecologically relevant AA sources. Commensals are therefore strong modifiers of food choice behavior by buffering the animal from the effect of dietary lack of eAAs.

Commensal bacteria can only buffer the absence of essential amino acids

Given the strong effect of commensals on food choice, we set out to test if they would also be able to suppress the food choice phenotype in flies which are impaired in Tyr synthesis. Surprisingly, addition of the five bacteria to the diet was not able to suppress the yeast appetite induced by neAA deprivation in Henna knockdown flies, while Tyr supplementation was able to suppress this appetite (Fig 4E). The bacteria mix was still able to reduce the yeast appetite induced by His deprivation in Henna knockdown flies, indicating that the bacteria are effective in this genetic background (S6 Fig). This indicates that the microbiota exerts its effect specifically in the context of eAA but not neAA depletion. This experiment also strongly suggests that the commensal pretreatment does not alter food choice in an indirect way (e.g., by suppressing the ability of flies to choose yeast or only increasing their preference for sucrose) and that the effect of the bacteria is not due to them serving as food, as these effects should also lead to a decreased yeast preference index in these flies. Commensal bacteria are therefore strong modifiers of food choice specifically in the context of eAA deficiencies.

Commensal bacteria increase egg laying upon essential amino acid deprivation

Stem cell proliferation and differentiation is limited by the availability of eAAs [64–66]. In Drosophila this is most evident in the context of egg production, in which depletion of eAAs strongly reduces egg laying [7,9]. We therefore tested if commensals would also be able to affect egg laying. As shown above, depriving animals of three different eAAs (His, Ile, or Val) led to the induction of yeast appetite, which was strongly suppressed in animals pretreated with commensals (S4 Fig). Removal of any of these three eAAs significantly decreased egg laying when compared to the complete holidic diet (Fig 4F). Flies treated with the five bacteria, however, laid a significantly higher number of eggs in the context of a diet without single eAAs when compared to the flies without bacterial pretreatment (Fig 4F). The microbiota is
therefore not only able to buffer the effect of removing eAAs in the context of food choice but also in terms of physiological traits such as the reduction of egg laying triggered by removing one eAA. Given the importance of fecundity for the fitness of the animal, this effect therefore suggests that in the adult fly the host–bacteria interaction is mutualistic.

Commensal bacteria specifically increase yeast appetite in the context of eAA depletion

To separately analyze the effects of the microbiome on yeast and sugar feeding, we chose to use the flyPAD assay [67]. Furthermore, to fully control the microbial conditions of the experiments, we performed the flyPAD experiments starting with flies kept in an axenic state, which ensured that our microbiome reconstitutions resulted in gnotobiotic flies. Flies kept on a full holidic diet and pretreated with the five bacteria did not show an increase in yeast feeding when compared to the germ-free controls (Fig 5A), but they showed an increase in sucrose feeding (Fig 5B). The sucrose effect is reminiscent of previous reports that the gut microbiota can increase food intake in flies [60]. In agreement with the data generated using the CAFE assay, both His and Ile deprivation led to a specific increase in yeast feeding in the axenic flies (Fig 5A), while sucrose feeding was unaltered (Fig 5B). Compared with germ-free flies, the gnotobiotic flies pretreated with the five gut bacteria showed a highly significant decrease in yeast feeding, corroborating the hypothesis that the microbiota suppresses yeast appetite (Fig 5A).

In contrast to the effect on yeast appetite, the increase in sugar feeding observed in the flies treated with bacteria was more variable. While a significant effect was observed in the His-deprived flies, Ile-deprived flies did not show an increase in sucrose feeding (Fig 5B). These data support the idea that commensal bacteria specifically change food choice by decreasing yeast appetite in eAA-deprived flies.

It has been indicated that the microbiota competes with their host for the availability of sugars in the diet [36]. To test this hypothesis and rule out that the effect on yeast choice is due to the observed increase in sugar intake, we increased the amount of sugar in the holidic diet. Adding increasing amounts of sucrose to the diet decreased the sugar appetite, suggesting that the bacteria were indeed reducing the sugar available to the fly from the diet (Fig 5D). Importantly, neither the levels of sucrose in the diet nor the level of sugar feeding affected the yeast appetite, showing that these macronutrient appetites are independently regulated (Fig 5C and 5D). Therefore, while a decrease in the sugar content of the food could account for the previously reported increase in food intake caused by gut bacteria [60], this effect is not related to the changes in food choice we describe. Gut bacteria therefore use an independent mechanism to specifically reduce the yeast appetite of the host.

Flies eat more vigorously from food containing commensal bacteria

Flies rely on the continuous replenishment of their microbiome through feeding [61]. If commensal bacteria provide protection against eAA depletion, one might expect flies to prefer ingesting food containing commensals. We therefore set out to compare the appetite of flies towards food with or without commensal bacteria using the flyPAD (Fig 5E). In agreement with our hypothesis, flies ate more vigorously from a food source containing the commensal bacteria when compared to the same food without commensals (Fig 5E). Flies are therefore able to increase feeding behavior when bacteria are present in the food. This suggests that flies are able to actively modulate their feeding behavior to replenish or modify their microbiota in order to profit from the physiological benefits of the commensals.
Fig 5. Commensal bacteria decrease yeast appetite upon essential amino acid (eAA) deprivation and are phagostimulators. Numbers of sips on yeast (A) or sucrose (B) of axenic flies prefed on holidic medium with different eAAs and with or without the commensal mix as measured using the flyPAD. $n = 70–82$. Numbers of sips on yeast (C) or sucrose (D) of axenic flies prefed on holidic medium containing different eAAs and different sucrose concentrations with or without the commensal mix as measured using the flyPAD. $n = 25–37$. (E) On the left, a diagram depicting the strategy used to test the phagostimulatory power of commensal bacteria. flyPAD drawing from [67]. On the right, numbers of sips of flies feeding from holidic base medium containing amino acids (AAs) with or without the commensal mix as measured using the flyPAD. $n = 37–39$. (A–E) Boxes represent upper and lower quartiles with the median. In this and other Figs, empty boxes represent non–bacteria-treated conditions and filled boxes represent bacteria-pretreated conditions. (A–D) Filled black circles represent a complete holidic medium or pretreatment with the bacteria mix. Open black circles represent flies not pretreated with bacteria mix. AA deprivation is indicated as –histidine (–His) or –isoleucine (–Ile) and in (C–D) sucrose concentration is indicated as values of 50 mM, 100 mM, or 200 mM. (A–C) Significance was tested using the Kruskal–Wallis test followed by Dunn’s multiple comparison test, in (D) using the Mann–Whitney test, and in (E) using the unpaired t test. Not significant (ns) $p > 0.05$, * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Underlying data used in this Fig are provided in S1 Data.

https://doi.org/10.1371/journal.pbio.2000862.g005
**Acetobacter pomorum** acts together with *Lactobacilli* to modify food choice

Our data suggest that specific bacteria directly act on host physiology and behavior and provide evidence contrary to a generalized effect of bacterial material. We therefore decided to use the gnotobiotic model to identify which bacteria in the mix were producing the change in feeding behavior in eAA-deprived animals. To do so, we first removed each species separately from the mix and tested if the reduced sets could suppress the yeast appetite of His-deprived flies. While removal of *Acetobacter pomorum* (*Ap*) abolished the capacity of the mix to suppress yeast appetite, removal of any of the other four species had no effect (Fig 6A). *Ap* alone, however, is not sufficient to change yeast appetite, indicating that it acts in concert with other bacteria in the mix. Given that *Lactobacilli* act together with *Ap* to alter metabolite composition in flies [68], we decided to test if *Ap* together with *Lactobacillus plantarum* (*Lp*) or *Lactobacillus brevis* (*Lb*) are sufficient to alter yeast appetite. Indeed, the combination of *Ap* with either *Lp* or *Lb* is sufficient to suppress the yeast appetite induced by deprivation from either His or Ile (Fig 6A and S7 Fig). This result also explains why removing either *Lp* or *Lb* from the five-bacteria mix had no effect, as these species seem to act redundantly. Furthermore, neither *Lp*, *Lb*, nor the combination of both change feeding behavior, highlighting the specificity of the combined *Ap–Lactobacilli* effect on yeast appetite (Fig 6A).

The same approach allowed us to conclude that *Ap* and *Lp* act together to increase sugar appetite (Fig 6B). In contrast to the effect on yeast appetite, the *Ap–Lb* combination has no effect on carbohydrate consumption (Fig 6B). This reinforces the previous data showing that yeast appetite is independent of sugar appetite. Taken together, these data show that *Acetobacter pomorum* can act together with either *Lactobacillus plantarum* or to a certain extent with *Lactobacillus brevis* to change food selection.

**Commensal bacteria do not seem to change the levels of eAAs in the host**

The ability of the commensal bacteria to compensate for the effect of eAA deprivation on yeast appetite and egg laying suggests that the bacteria could supply the host with eAAs, thus buffering the animal from the absence of these important nutrients in the diet. Such an effect would be reminiscent of the role of the *Buchnera* endosymbiont in aphids, which allows this insect to thrive while feeding on sap, which contains very low amounts of AAs [69]. We tested this hypothesis by depriving flies from three different eAAs (His, Ile, and Val) and comparing the levels of free AAs in the heads of flies that had been either pretreated or not with the five-bacteria mix. We decided to focus on the AA levels in heads to avoid effects due to changes in the number of eggs carried by the fly and because of evidence that nutrient sensing could act at the level of the brain of the fly to change food preference [12]. His, Ile, or Val deprivation lead to a drastic decrease in the levels of these three AAs in head extracts (Fig 7A), which is likely to cause the previously observed increases in yeast appetite (Fig 2). This effect was specific to the manipulated AAs, as the levels of nonmanipulated AAs neither increased nor decreased (Fig 7A). AA-satiated flies treated with the bacterial mix did not show an increase in His, Ile, or Val. Surprisingly, deprived flies continued having very low titers of the measured eAA independent of the bacterial pretreatment (Fig 7A). This stands in contrast to the clear effect of the bacterial pretreatment on yeast preference and egg laying (Fig 4 and S4 Fig). Our failure to observe changes in eAA levels induced by bacterial pretreatment opens the intriguing possibility that the commensal bacteria modify food choice and egg laying through an AA-independent mechanism (Fig 7B).
Fig 6. *A. pomorum* and *Lactobacilli* are sufficient to modify yeast appetite. Numbers of sips on yeast (A) or sucrose (B) as measured using the flyPAD of axenic flies prefed on complete holidic medium or holidic medium without histidine (His) and pretreated with different bacterial mixes. Filled black circles represent the complete holidic medium or presence of specific bacteria in the pretreatment mix. Open black circles represent the absence of specific bacteria in the pretreatment mix. His deprivation is indicated as –histidine (–His). Boxes represent upper and lower quartiles with the median. \( n = 46–120 \). Significance was tested using the Kruskal–Wallis test followed by Dunn’s multiple comparison test. Not significant (ns) \( p > 0.05 \), *** \( p < 0.001 \). Underlying data used in this Fig are provided in S1 Data.

https://doi.org/10.1371/journal.pbio.2000862.g006
Commensal bacteria do not seem to change the levels of essential amino acids (eAAs) in the host. (A) Histidine (His), isoleucine (Ile), and valine (Val) concentrations in the heads of flies prefed on complete holidic medium (green) or holidic medium lacking His, Ile, or Val (blue), without (empty columns) or with (filled columns) commensals pretreatment. The columns represent the mean and the error bars the standard error of the mean of three independent experiments. Filled black circles represent complete holidic medium or pretreatment with the bacteria mix. Open black circles represent no pretreatment with bacteria mix. Amino acid (AA) deprivation is indicated as –histidine (–His), –isoleucine (–Ile), or –valine (–Val). Significance was tested using the unpaired *t* test with Bonferroni correction. Not significant (ns) *p* > 0.05, *p* < 0.05, **p* < 0.01. (B) Model of the impact of eAAs on food choice and reproduction, depending on the presence of the microbiota of the host. The nervous system is highlighted in turquoise, AAs in orange, and commensal bacteria in purple. Arrow weight from the proboscis to the food drops indicate amount of feeding, and the number of eggs reflect the reproductive output. The orange and purple arrows indicate potential effects of eAAs and metabolites, respectively, at the level of the nervous and reproductive systems. Metabolite X refers to a hypothetical metabolite mimicking the presence of AAs. Underlying data used in this Fig are provided in S1 Data.

https://doi.org/10.1371/journal.pbio.2000862.g007
Discussion
The complexity of natural foods makes it very difficult to identify the nutrients which guide behavior and physiology. Furthermore, while the microbiota has been suggested to affect behavior, until now its influence on specific nutrient appetites has not been explored. The present study identifies two factors as strong and specific modulators of feeding decisions and reproduction: eAAs and commensal bacteria (Fig 7B).

Amino acids as potent modulators of protein appetite
Multiple nutrients, including AAs, metals, vitamins, and sterols have been shown to be nutritional modulators of life history traits [7]. Given that in an ecological setting, yeast is likely to be an important nutritional source of these nutrients, it is therefore surprising that the animal only develops a yeast appetite upon the restriction of AAs. One reason might be that animals have not evolved strategies to regulate the intake of all nutrients separately but just specific ones. This could be explained by the fact that in an ecological setting, animals do not need to react to the lack of each nutrient independently as these distinct nutrients are found together in nature in the form of food. If yeast is the ecologically relevant source of most nutrients required for the fly, then the animal could use the lack of internal AAs as a proxy for the concomitant lack of other nutrients, such as minerals, metals, and vitamins. The increase in yeast appetite triggered by AA deprivation would thus be sufficient to compensate for the lack of other nutrients. This would highlight that while synthetic diets are invaluable tools for studying the impact of nutrients on physiology and behavior, the results obtained from such studies always need to be interpreted in the context of ecologically relevant food sources such as yeast. It is also possible, however, that foods other than yeast can serve as sources for specific nutrients. Further focused analyses of the effect of specific nutrient classes on behavior will be required to identify the full capacity of Drosophila to maintain nutrient homeostasis.

Signaling mechanisms and neuronal circuits underlying amino acid sensing
How do flies sense the internal deficiency of AAs and what are the circuit mechanisms allowing them to increase yeast or AA intake upon this nutritional restriction? The brain should be able to detect changes in AA concentrations given that the concentration of free eAAs in the head drops dramatically upon their removal from the diet. Changes in behavior could therefore either be informed by direct sensing of internal AA titers by the nervous system and/or by detecting a signal released by peripheral tissues as response to lack of AAs (Fig 7B). Two molecular mechanisms have been proposed as mediating neuronal AA sensing: the TOR and GCN2 pathways. In Drosophila, neuronal TOR signaling has been proposed to influence food selection [12], and in Drosophila and vertebrates, GCN2 has been proposed to direct behavioral nutrient homeostasis by mediating post-ingestive neuronal sensing of AAs [70–72]. The involvement of neuronal GCN2 in nutrient selection in vertebrates has, however, been recently challenged [73]. We therefore have only very rudimentary clues as to what could be the mechanisms mediating internal sensing of AA availability and subsequent changes in behavior. Our data that flies can behaviorally react to the absence of a genetically engineered neAA deficiency suggest that whatever the sensing mechanism, it has to be able to sense the absence of any AA. Intriguingly, at the molecular level, nutrient-sensing pathways such as the TOR pathway have been proposed to mainly react to specific AAs [46]. Our data suggest that nutrient-sensing pathways could have a much broader spectrum of action. Alternatively, the sensing of AA deficiencies could rely on different molecular mechanisms, which could for example detect the
decrease in translation induced by the lack of AA$s$. Such a decrease could either be sensed per se or could lead to a decrease (or increase) of specific translation products, which could serve as signals to alter behavior. Pinpointing the site and cellular substrate of AA sensing as well as the underlying molecular mechanisms remains a key challenge in the field of nutrient homeostasis.

At the circuit level, the lack of AA$s$ is likely to lead to a change in chemosensory processing that would lead to a change in nutrient preference. For proteins, such changes have been proposed in locusts [74]. In Drosophila, chemosensory neurons have been shown to be directly modulated by the internal energy state of the animal [13]. Furthermore, mating has been shown to modulate salt taste processing using the same circuit as yeast intake [14]. How internal AA states affect yeast and AA chemosensory processing, however, still remains to be elucidated. The main obstacle is the lack of information on the identity of the chemosensory neurons mediating yeast and AA feeding in Drosophila. Identifying these neurons and analyzing how yeast perception is modulated by internal state should allow us to better understand how the internal AA state directs feeding decisions.

Host–microbiota interactions in nutrient choice

The extent to which the microbiota affects specific nutrient appetites has not been previously explored. We show that when flies are AA challenged, commensal bacteria reduce their compensatory yeast appetite. The increase in sugar appetite observed in flies harboring commensals further decreases the ratio of protein to carbohydrate intake, an important determinant of life history traits in animals, including vertebrates [2]. Given that a reduction in yeast and AA intake leads to an increase in lifespan, our observation that commensal bacteria reduce the intake ratio of proteins to carbohydrates could account for the shorter lifespan of axenic flies [75]. It is interesting to note that the flies harboring commensals are able to increase their reproductive output (our study and [76]) despite their lower protein intake. Commensal bacteria could therefore have a highly beneficial impact on the fly, enabling it to simultaneously maximize lifespan and reproductive output.

The increase in sugar appetite observed in gnotobiotic flies can be simply explained by the bacteria utilizing the sugar in the food and therefore inducing a carbohydrate deficit in flies. The decrease in yeast appetite, however, is more difficult to explain. One simple possibility could have been that these bacteria act as nutrients, which has been proposed for yeast and other fungi [77]. Our findings that inactivated bacteria do not induce a change in feeding behavior, that bacteria are not able to suppress the yeast appetite induced by neAA$s$ when we perturb the function of Henna, and that only specific commensal bacteria change yeast appetite, all indicate that the microbiota acts in a very specific way to alter food choice. Furthermore, previous data that microbes can improve the uptake of AA$s$ [77,78] are not sufficient to explain the suppression of yeast appetite in flies pretreated with commensals, as we use holidic media completely devoid of eAA$s$. A key question is thus: what could be the mechanisms by which gut bacteria change yeast appetite and increase egg laying?

Mechanisms by which gut bacteria could control nutrient homeostasis in the host

Intracellular symbiotic bacteria are known to provide eAA$s$ in other insects [69], and gut bacteria have been shown to provide significant amounts of eAA$s$ in vertebrates, including humans [79]. A straightforward hypothesis would therefore be that these two bacteria are able to provide the fly with eAA$s$. However, we were not able to detect changes in the levels of free eAA$s$ in flies pretreated with the commensal mix. In spite of this, we are reluctant to
completely rule out that the microbiota acts on yeast appetite and egg laying by providing eAAs. It is possible, for example, that in an eAA-deprived situation, bacterially produced eAAs are immediately utilized without increasing the pool of free AAs. In such a model, bacterially derived eAAs would be fully allocated to sustain reproduction as well as alleviate the process which triggers changes in yeast appetite upon eAA deprivation. In such a situation, it is conceivable that one would not be able to measure an increase in free eAAs provided by the bacteria. Our data, however, suggest that commensal bacteria do not act by providing eAAs to the host. What could be alternative mechanisms by which they influence behavior and egg production? They could secrete metabolites that help the host to increase its ability to use its remaining AAs, thereby buffering the fly from the effects of dietary eAAs. Intriguingly, both yeast appetite and reproduction are thought to be regulated by the nutrient-sensitive TOR pathway [12,80–83], and commensals have been shown to be able to modulate this pathway [37]. It is therefore possible that these bacteria act directly on nutrient sensing pathways by releasing metabolites that mimic the availability of eAAs (Metabolite X in Fig 7B). Distinguishing between these hypotheses will require comprehensive metabolome analyses of flies in different bacterial and nutrient states as well as careful genetic and behavioral studies, both at the level of the host and the bacteria.

The metabolic repertoire of an organism is evolutionarily fixed in its genome. As such, it represents a static set which can mainly be modulated by transcriptional control. The observation that flies ingest more food containing commensal bacteria suggests that they might be able to direct their feeding behavior to replenish or maintain a specific microbiome composition. It is therefore attractive to speculate that the dynamic nature of the microbiome in flies paired with the ability to modulate the replenishment of gut microbes through feeding could allow them to extend and adapt their metabolic repertoire by exploiting that of the microbiome [84]. This ability could partially explain the success of *Drosophila* in adapting to a wide range of habitats.

Our understanding of how the microbiota influences behavior remains extremely rudimentary. In vertebrates, this task is made especially daunting by the complexity of their microbiota. *Drosophila*, on the other hand, has proven to be an especially powerful model for understanding microbe–host interaction because of the ability to isolate a single bacterial species promoting physiological effects such as improved growth [85,86]. Especially in vertebrates, many effects of the microbiome on the host, however, are likely to rely on interactions among different microbial species. Our finding that *Ap* acts together with *Lactobacilli* to influence food choice provides a powerful system for not only understanding how microbes act on the host to influence brain function but also how microbes cooperate to shape complex host traits. Microbes could act together by exchanging metabolites to act on the host. Alternatively, one bacterium could support the growth and survival of the other in nutritionally challenging situations, allowing it to exert its behavioral effect. The identification of these two bacterial species as mediators of food choice paired with the powerful genetic toolkit available in *Drosophila* provides a unique opportunity to identify the mechanisms by which microbes interact to shape the behavior of the host.

**The importance of nutritional–microbial interactions in influencing host behavior across phyla**

Our findings highlight a new function of the microbiota in modulating nutrient-specific appetites. Given that in *Drosophila*, AA state not only controls food intake but also more complex behavioral features, such as risk taking [16], the microbiota could influence behavior beyond feeding. Furthermore, because AAs and nutrient sensing play a pivotal role in controlling physiology, neurodevelopmental disorders [87,88], and behavior across metazoans, such
mechanisms could be conserved across phyla. Nutrition could therefore provide a framework for understanding how the microbiome influences behavior, disease, and physiology across phyla. Our findings highlight the power of the Drosophila model for dissecting complex nutritional–microbial–behavioral interactions and suggest the intriguing possibility that commensal bacteria influence behavior and brain function in invertebrates and vertebrates by tapping into the nutrient-sensing abilities of the nervous system.

Materials and methods

Methods and protocols for Drosophila rearing, media preparations, and microbial manipulations are available as a collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

Drosophila stocks and genetics

Unless stated otherwise, all experiments were performed with mated w^{1118} female flies. Ubiquitous (tubulin-Gal4 [89]), pan-neuronal (elav-Gal4 [90]), tracheal (btl-Gal4 [91]), or fat body (Cg-Gal4 [92], BL #7011) expression of RNAi delivering transgenes against Henna (CG7399) was achieved by crossing Gal4-carrying female flies with three independent UAS-Henna-RNAi stocks (HennaIR1: VDRC #35240; HennaIR2: NIG-RNAi #7399R-3; HennaIR3: BL #29540). The full genotypes of experimental flies are listed in S1 Table.

Drosophila rearing, media, and dietary treatments

Flies were reared on yeast-based medium (YBM) (per liter of water: 8 g agar [NZYTech, PT], 80 g barley malt syrup [Próvida, PT], 22 g sugar beet syrup [Gráfachter, DE], 80 g corn flour [Próvida, PT], 10 g soya flour [A. Centazi, PT], 18 g instant yeast [Saf-instant, Lesaffre], 8 ml propionic acid [Argos], and 12 ml nipagin [Tegospet, Dutscher, UK] [15% in 96% ethanol] supplemented with instant yeast granules on the surface [Saf-instant, Lesaffre]). To ensure a homogenous density of offspring among experiments, fly cultures were always set with 5 females and 4 males per vial and left to lay eggs for 7 d. Flies were reared in YBM until adulthood. Holidic media (HM) were prepared as described previously [7] using the HUNTaa formulation without food preservatives, with the exception of the HM used for pretreating axenic and gnotobiotic flies, for which we used an HM with an improved AA composition [93]. The different HM used in this study are described in S2 Table and S3 Table. In all experiments where we refer to all neAAs removal, L-glutamate was still present in the diet in order to prevent any possible adverse effects in neuronal function. Sucrose medium consisted of Kleenex tissue soaked with 5 ml of a 100 mM sucrose (Sigma-Aldrich, #84097) solution. For all experiments using the HM, the following dietary treatment protocol was used in order to ensure a well-fed state and minimize the microbial load in the flies [61] (S8 Fig): groups of 1–5-d-old flies (16 females and 5 males) were collected into fresh YBM-filled vials and transferred to fresh YBM after 48 h. Following a period of 24 h, flies were transferred to different HM for 72 h and immediately tested in the indicated assay. Flies treated using this protocol had a low titer of commensal bacteria (S3 Fig). For yeast dilution experiments presented in S5 Fig, flies were kept for 72 h prior to the behavioral assay on media containing 200 mM sucrose (Sigma-Aldrich, #84097), 2% agar (Difco, # 214530), and variable instant yeast concentrations: 5%, 2.5%, 1%, and 0% (Saf-instant, Lesaffre). After preparation, all yeast-based media were autoclaved before pouring into culture vials. Fly rearing, maintenance, and behavioral testing were performed at 25˚C in climate-controlled chambers at 70% relative humidity in a 12-h light–dark cycle (Aralab, FitoClima 60000EH). Polypropylene fly vials (VWR, #734–2261) were used.
These protocols are available in the following collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

**Axenic *Drosophila* generation and rearing**

The protocol to generate axenic *w^{1118}* fly cultures by sterilizing embryos was adapted from [94]: embryos were put for 2 min in 2.5% active chlorine (50% bleach) followed by 2 min in 70% ethanol and 2 min in autoclaved distilled water. The embryos were then transferred onto sterile food (autoclaved before pouring into culture vials) containing antibiotics (final concentrations: 416.7 μg/ml tetracycline [high dose], 41.67 μg/ml chloramphenicol, 41.67 μg/ml ampicillin, and 8.333 μg/ml erythromycin). In order to compensate for the developmental delay observed in axenic larvae [37], the yeast content of the medium was increased to 41.67 g per liter. Axenic *w^{1118}* flies were regularly transferred into vials containing freshly prepared, antibiotic-supplemented, high-yeast food (S8 Fig). The absence of bacteria was assessed by grinding flies in sterile 1x PBS and spreading the suspension on LB, MRS, or Mannitol plates. LB and MRS plates were incubated at 37˚C and Mannitol plates at 30˚C, respectively, before assessing the presence of bacterial colonies. The antibiotic treatment did not lead to any apparent malaise in the treated flies. Furthermore, to ensure that the antibiotics exposure would not directly affect the experimental animals, these were raised in sterile food without antibiotics (S8 Fig). Importantly, the results obtained using the gnotobiotic flies fully recapitulate the results obtained with the conventionally reared “low bacteria titer” flies.

All experiments in Figs 5A–5D, 6 and S7 were performed using axenic or gnotobiotic flies. These protocols are available in the following collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

**Bacterial species and cultures**

The following bacterial species and strains (kindly provided by François Leulier, IGFL, France, and Won-Jae Lee, SNU, South Korea) were used in this study: *Lactobacillus plantarum*<sup>WJL</sup> [62], *Lactobacillus brevis*<sup>EW</sup> [62], *Acetobacter pomorum* [62], *Commensalibacter intestini*<sup>A911T</sup> [62], and *Enterococcus faecalis* [63]. *Lactobacilli* were cultured in 10 ml of liquid MRS medium (Sigma-Aldrich, #69966) in 14 ml culture tubes (Thermo Fisher Scientific, #150268) at 37˚C for 24 h without agitation. *C. intestini*<sup>A911T</sup> and *A. pomorum* were cultured in a liquid mannitol medium (3 g/l Bacto peptone [Difco, #0118–17], 5 g/l yeast extract [Difco, #212750], 25 g/l D-mannitol [Sigma-Aldrich, #M1902]) at 30˚C for 48 h under 170 rpm agitation. *C. intestini*<sup>A911T</sup> was cultured in 20 ml of medium in 50-ml tubes (Falcon), and *A. pomorum* was cultured in 200 ml of medium in 500-ml flasks. *E. faecalis* was cultured in 200 ml of liquid LB medium (Sigma-Aldrich, #L3022) in 500-ml flasks at 37˚C for 24 h under 220 rpm agitation. Liquid cultures were set with colonies grown in fresh solid media (15 g/l agar [Difco, # 214530]).

These protocols are available in the following collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

**Inoculation of HM with bacteria**

Prior to transferring the flies, each HM vial was inoculated with either single or different combinations of the following bacterial species: *L. plantarum*<sup>WJL</sup> (6.4 x 10⁴ CFU), *L. brevis*<sup>EW</sup> (5.31 x 10⁵ CFU), *C. intestini*<sup>A911T</sup> (9.04 x 10⁴ CFU), *A. pomorum* (9.5 x 10⁵ CFU), *E. faecalis* (1.11 x 10⁵ CFU), and *E. coli* (0.924 x 10⁶ CFU). To prepare this mixture, the necessary volume of liquid culture for each bacterial species was centrifuged three times at 3,000 rpm for 10 min and repeatedly resuspended in 1x PBS. To exclude an effect from residual components of bacterial
media, the equivalent volume of sterile bacterial media was centrifuged in parallel and used as a control. After the final centrifugation, both the control and the bacterial pellet were resuspended in sufficient 1x PBS to achieve an inoculation volume of 50 μl per vial. For the experiments with heat-inactivated bacteria, the bacterial suspension was incubated at 100°C for 10 min before inoculation in HM vials. The final suspensions were added to the surface of the HM and allowed to dry for approximately 1 h before the addition of flies. Note that even when not pretreated to be axenic, because of rearing protocol flies used in all experiments had a very low starting titer of internal microbes prior to inoculation (S3 Fig).

These protocols are available in the following collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

Calculation of internal bacterial load of flies
Flies were surface sterilized to remove any bacteria that could be found on the cuticle by washing them in 70% ethanol followed by two washes in sterile 1x PBS. Flies were grinded in 1x PBS (500 μl/18 flies) and diluted 180X. The suspension was then plated on LB, MRS, or Mannitol medium. LB and MRS plates were incubated at 37°C and mannitol plates at 30°C before counting the number of bacterial colonies.

These protocols are available in the following collection in protocols.io dx.doi.org/10.17504/protocols.io.hdtb26n.

Two-color food choice assay
Two-color feeding preference assays were performed as previously described [12]. Groups of 16 female and 5 male flies were briefly anesthetized using light CO₂ exposure and introduced into tight-fit-lid Petri dishes (Falcon, #351006). For the yeast choice assays, the flies were given the choice between nine spots of 10 μl sucrose solution mixed with red colorant (20 mM sucrose [Sigma-Aldrich, #84097]; 7.5 mg/ml agarose [Invitrogen, #16500]; 5 mg/ml Erythrosin B [Sigma-Aldrich, #198269]; 10% PBS) and nine spots of 10 μl yeast solution mixed with blue colorant (10% yeast [Saf-instant, Lesaffre]; 7.5 mg/ml agarose; 0.25 mg/ml Indigo carmine [Sigma-Aldrich, #131164]; 10% PBS) for 2 h. For the defined nutrient-choice assays, flies were given the choice between HM lacking AAs and containing 20 mM sucrose mixed with red colorant (option 1: sucrose) and HM lacking sucrose and containing the nutrients required for the experiment mixed with the blue colorant (option 2). In these experiments, the agar concentration in the HM was changed to 1.5%. After visual inspection of the abdomen under the stereo microscope (Zeiss, Stereo Discovery.V8), each female fly was scored as having eaten either sucrose (red abdomen), yeast (blue abdomen), or both (red and blue or purple abdomen) media. The yeast preference index (YPI) for the whole female population in the assay was calculated as follows: (n_blue yeast − n_red sucrose) / (n_red sucrose + n_blue yeast + n_both). Initially, dye-swap (red yeast versus blue sucrose choice) experiments were performed in parallel, and because the change of feeding preference was observed in both conditions, we opted to exclusively perform red sucrose versus blue yeast choice experiments. In all experiments, the observer was blind for both diet and genotype. All assays were performed between ZT6 and ZT9.

CAFE assay
CAFE assays were based on a protocol previously described [14,47] with some adaptations. On the assay day, flies were anesthetized with CO₂, sorted under a stereo microscope in groups of 18 females, and allowed to recover for 3 h at 25°C. The CAFE chamber consisted of a large plastic vial (50 x 100 mm) (Semadeni AG, #6128) with 6 5-μl glass capillaries (Hirschmann,
Capillaries were filled with 20 mM sucrose or 10% yeast extract (Sigma-Aldrich #1625) solutions and placed in an alternating circular fashion. Each group of flies was aspirated into a CAFE chamber, and during the 4 h of the assay, four experimental readings per capillary were scored ($t_0$, $t_0+1$ h, $t_0+3$ h, and $t_0+4$ h) to determine consumption. In order to correct for evaporation, each set of experimental chambers was accompanied by an empty chamber (no flies). Total sucrose or yeast extract consumption per time point was determined by subtracting the sum of the readings of the three capillaries of the respective solution in the empty chamber from the equivalent values in the experimental chamber. Consumption per fly was obtained by dividing sucrose or yeast extract total consumption by the number of living flies at the end of the assay.

**flyPAD assays**

FlyPAD assays were performed as described in [67]. For food choice experiments, single flies in different dietary conditions were tested in arenas that contained two kinds of food patches: 10% yeast and 20 mM sucrose, each mixed with 1% agarose.

To measure the phagostimulatory power of bacteria (Fig 5E), we used a flyPAD setup that had never been exposed to yeast. All tested flies were deprived from amino acids using HM–AAs. For the flyPAD assays, one feeding well per arena was filled with HM without sucrose, either intact media (holidic AA medium) or media supplemented with the bacterial mixture (holidic AA medium with five bacteria) (in HM: *L. plantarum* \(^{\text{WL}}\) 1.02 x 10^7 CFU, *L. brevis* \(^{\text{EW}}\) 8.49 CFU, *A. pomorum* 1.52 x 10^3 CFU, *C. intestini* \(^{\text{A911T}}\) 1.45 x 10^2 CFU, and *E. faecalis* 1.77 x 10^2 CFU). These media were prepared by adding agarose (1%) as a gelling agent together with cholesterol after autoclaving. Media were prepared on the experimental day and maintained at 30°C in a heat block. Preparation of the control and bacterial mixture pellets were performed as described above and directly resuspended in the HM without sucrose to generate holidic AA medium and holidic AA medium with five bacteria, respectively. Each medium was loaded into a single feeding well of the arena.

For all experiments, flies were individually transferred to flyPAD arenas by mouth aspiration and allowed to feed for 1 h at 25°C, 70% relative humidity. The total number of sips per animal over this hour was calculated using previously described flyPAD algorithms [67]. Noneating flies (defined as having fewer than two activity bouts during the assay) were excluded from the analysis.

**Egg-laying assays**

Groups of 16 female and 5 male flies were briefly anesthetized using light CO\(_2\) exposure and transferred to apple juice agar plates (per liter, 250 ml apple juice, 19.5 g agar, 20 g sugar, and 10 ml nipagin [15% in ethanol]), where they were allowed to lay eggs for 24 h. Flies were then removed and counted and eggs were counted. Egg laying was calculated by dividing the number of eggs by the number of living females at the end of the assay.

**Total mRNA extraction, RT-PCR, and quantitative real-time PCR**

Flies used for mRNA extraction were snap frozen in dry ice and kept at –80°C until used. Behavioral assays were performed in parallel to confirm that sibling flies presented the expected feeding phenotype. mRNA was extracted from flies (three flies per condition) using the following procedure: flies were grinded and homogenized for 20 s (using pestles #Z359947, Sigma) in 100 μl of PureZOL (#732–6890, Bio-Rad). 250 μl of PureZOL was further added and mixed by pipetting and incubated at RT for 10 min. Finally, 350 μl of 100% ethanol was added, and the samples were mixed and transferred to a Zymo column (Direct-zol RNA MicroPrep #R2062,
Zymo research). The manufacturer’s instructions were followed to purify the mRNA (including DNAse treatment), and samples were eluted in 15 μl of distilled RNase/DNase-free water. The concentration of the total mRNA samples was determined by performing a spectrophotometer scan in the UV region. Total RNA (1 μg) was reverse transcribed (RT) using the iScript Reverse Transcription Supermix for RT-PCR kit (#170–8840 Bio-Rad), following the manufacturer’s instructions. The expression of Henna was determined using real-time PCR. Each cDNA sample was amplified using SsoFast EvaGreen Supermix on the CFX96 Real-Time System (Bio-Rad). Briefly, the reaction conditions consisted of 1 μl of 1:10 diluted cDNA, 1 μl (10 μM) of each primer, 10 μl of supermix, and 7 μl of water. The cycle program consisted of enzyme activation at 95˚C for 30 s, 39 cycles of denaturation at 95˚C for 2 s, and annealing and extension for 5 s. The primers used in this reaction are listed in S4 Table. This experiment was performed using three experimental replicas and two technical replicas per genotype. Appropriate non-template controls were included in each 96-well PCR reaction, and dissociation analysis was performed at the end of each run to confirm the specificity of the reaction. Absolute levels of RNA were calculated from a standard curve and normalized to the internal controls (Actin42A and RpL32). The relative quantitation of each mRNA was performed using the comparative Ct method. Data processing was performed using Bio-rad CFX Manager 3.1 (Bio-Rad).

Amino acid measurements in fly heads

500 females per condition were collected on the same day as behavioral assays and were snap frozen in dry ice. Flies were kept at –80˚C until head preparation for amino acids measurements. Fly heads were separated from other body parts by vortexing the Eppendorf tubes and posteriorly passing the debris through 710-mm and 425-mm sieves (Retsch GmbH). Fly heads were counted before homogenization to ensure that the same number was used for all conditions. Heads were homogenized in 200 μl of 2.5% TCA and centrifuged for 10 min at top speed at 4˚C. The supernatant was recovered and stored at 4˚C for analysis. Amino acid quantification was performed by HPLC at a clinical laboratory (Joaquim Chaves Laboratories, PT). Amino acids were detected using AccQ.Tag (Waters, #176001235).

Supporting information

S1 Fig. Deprivation of amino acids specifically increases amino acid appetite. (A) Feeding assay in which flies were given the choice between the holidic medium lacking AAs (sucrose option) and the holidic medium lacking sucrose and one of the different nutrient classes. Animals were either kept on full holidic medium or holidic medium lacking AAs. (B) Feeding assay in which flies were given the choice between two options: 1) the holidic medium lacking amino acids (sucrose) and 2) the holidic medium lacking sucrose, lacking sucrose and all AAs, lacking sucrose and neAAs, or lacking sucrose and eAAs. Circles represent yeast preference in single assays, with line representing the median and whiskers the interquartile range. n = 12–15. (A) Significance was tested using the Kruskal-Wallis test followed by Dunn’s multiple comparison test or (B) the Mann Whitney test. Not significant (ns) p > 0.05, *** p < 0.001. Underlying data used in this Figure are provided in S1 Data.

S2 Fig. Henna RNAi leads to specific gene knockdown, and the knockdown effect on food choice is Tyr and tissue specific. (A) Henna mRNA levels measured from whole flies and normalized to two internal controls (Actin 42A and RpL32). The columns represent the mean and the error bars the standard error of the mean. n = 6. (B) Feeding preference of Henna knockdown animals using three independent hairpins. Animals were kept on holidic medium.
lacking neAAs, holidic diet lacking neAAs with 1x Tyr added back, or holidic medium lacking neAAs with 1x Pro added back. n = 10. (C) Feeding preference of control and Henna (Hn) knockout flies in different tissues upon removal of either all AAs or all neAAs. Cg-Gal4 drives expression in fat body, elav-Gal4 in neurons and btl-Gal4 in trachea. n = 14–20. (B and C) Circles represent yeast preference in single assays, with line representing the median and whiskers the interquartile range. Significance was tested using the Kruskal-Wallis test followed by Dunn’s multiple comparison test. Not significant (ns) p > 0.05, ** p < 0.01, *** p < 0.001. Underlying data used in this Figure are provided in S1 Data. (TIF)

S3 Fig. Levels of internal microbes in non-axenic flies. The internal load of bacteria inside flies was calculated as CFU/fly after bacterial colony count on LB (A), Mannitol (B), or MRS (C) media which sustain the growth of different bacterial species as indicated in the title of each graph. The load of bacteria was assessed for flies kept on holidic medium without His and without (empty columns) or with (filled columns) pretreatment with the commensal bacteria mix. Flies used to generate data in Figs 1, 2, 3, 4, 5E, 7, S1, S2, S4, S5A and S6 were treated using this or very similar rearing protocols. The columns represent the mean and the error bars, the standard error of the mean of 3 replicates from 2 independent experiments. Filled black circles represent pretreatment with the bacteria mix. Open circles represent no pretreatment with bacteria. AA deprivation is indicated as –His. Significance was tested using the unpaired t-test. * p < 0.05, ** p < 0.01. Underlying data used in this Figure are provided in S1 Data. (TIF)

S4 Fig. Commensal bacteria can reduce the protein appetite induced by dietary removal of any eAA. Feeding preference of animals kept either on holidic medium, or holidic medium lacking one of the 10 eAAs with or without pretreatment with 5 bacteria commensal mix. Data on the different graphs were collected on two independent days. Circles represent yeast preference in single assays, with line representing the median and whiskers the interquartile range. Filled circles represent assays in which flies had been pretreated with the 5 bacteria mix. n = 18–20. Significance was tested using the Kruskal-Wallis test followed by Dunn’s multiple comparison test, except for testing the effect of commensals, for which the Mann Whitney test was used. * p < 0.05, ** p < 0.01, *** p < 0.001. Underlying data used in this Figure are provided in S1 Data. (TIF)

S5 Fig. A non-commensal bacterium does not reduce yeast preference and commensal bacteria also affect food choice on low-yeast diets. (A) Feeding preference of animals kept either on holidic medium, or holidic medium lacking His with or without pretreatment with the 5 commensal bacteria mix or E. coli. Circles represent yeast preference in single assays, with line representing the median and whiskers the interquartile range. (A and B) Filled circles represent assays in which flies had been pretreated with the bacteria mix. n = 20. Significance was tested using the One-way analysis of variance test followed by Bonferroni’s multiple comparison test in (A) and using the Mann Whitney test in (B). Not significant (ns) p > 0.05, * p < 0.05, ** p < 0.01, *** p < 0.001. Underlying data used in this Figure are provided in S1 Data. (TIF)

S6 Fig. Commensal bacteria reduce yeast preference in Henna knockout flies upon eAA deprivation. Feeding preference of control and Henna knockout animals kept on holidic...
medium, or holcid medium lacking His with or without pretreatment with 5 commensal bacteria mix. n = 20. Circles represent yeast preference in single assays, with line representing the median and whiskers the interquartile range. Filled circles represent assays in which flies had been pretreated with the bacteria mix. Significance was tested using the Mann Whitney test followed by Bonferroni correction. * p<0.05, ** p<0.01, *** p<0.001. Underlying data used in this Figure are provided in S1 Data.

S7 Fig. *A. pomorum* and *Lactobacilli* are sufficient to reduce the yeast appetite induced by dietary removal of isoleucine. Numbers of sips on yeast as measured using the flyPAD of axenic flies pre-fed complete holcid medium or holcid medium without Ile and pretreated with different bacterial mixes. Filled black circles represent complete holcid medium or presence of specific bacteria in the pretreatment mix. Open black circles represent absence of specific bacteria in the pretreatment mix. Ile deprivation is indicated as -Ile. Boxes represent upper and lower quartiles with median. n = 25–55. Significance was tested using the Kruskal-Wallis test followed by Dunn’s multiple comparison test. *** p<0.001. Underlying data used in this Figure are provided in S1 Data.

S8 Fig. Diagram depicting the chronology of the dietary and microbial manipulations of flies used in experiments.

S1 Table. Detailed genotypes of flies used in this study.

S2 Table. Recipe for generation of HM used in this study. Gray fields indicate manipulated nutrients.

S3 Table. Detailed composition of amino acid solutions for HM used in this study.

S4 Table. Nucleotide sequence of primers used for qRT-PCR.

S1 Data. Raw data used for generation of all figures.

Acknowledgments

We thank François Leulier and Won-Jae Lee for providing bacterial strains and kind advice on growing them and Irene Miguel-Aliaga and Stefan Luschnig for providing fly strains. Further stocks were obtained from the Bloomington Drosophila Stock Center (NIH P40OD018537), the Vienna Drosophila Resource Center (Vienna, Austria), and the National Institute of Genetics. We thank François Leulier, Samuel Walker, Dennis Goldschmidt, and members of the Behavior and Metabolism laboratory for helpful discussions and comments on the manuscript and Gil Costa for illustrations.

Author Contributions

Conceptualization: Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Carlos Ribeiro.
Formal analysis: Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Ana Patrícia Francisco, Gabriela Tondolo Fioreze, Carlos Ribeiro.

Funding acquisition: Carlos Ribeiro.

Investigation: Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Ana Patrícia Francisco, Gabriela Tondolo Fioreze, Margarida Anjos, Célia Baltazar, Ana Paula Elias.

Methodology: Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Ana Patrícia Francisco, Carlos Ribeiro.

Project administration: Carlos Ribeiro.

Resources: Pavel M. Itskov, Matthew D. W. Piper.

Supervision: Carlos Ribeiro.

Visualization: Carlos Ribeiro.

Writing – original draft: Carlos Ribeiro.

Writing – review & editing: Ricardo Leitão-Gonçalves, Zita Carvalho-Santos, Ana Patrícia Francisco, Gabriela Tondolo Fioreze, Pavel M. Itskov, Carlos Ribeiro.

References

1. Joint WHO/FAO Expert Consultation on Diet, Nutrition and the Prevention of Chronic Diseases (2002: Geneva, Switzerland), World Health Organization. Dept. of Nutrition for Health and Development. Diet, nutrition and the prevention of chronic diseases : report of a joint WHO/FAO expert consultation, Geneva, 28 January—1 February 2002. Geneva: World Health Organization; 2003. http://www.who.int/iris/handle/10665/42665

2. Simpson SJ, Raubenheimer D. The nature of nutrition. Princeton, Princeton University Press; 2012.

3. Simpson SJ, Le Couteur DG, Raubenheimer D. Putting the Balance Back in Diet. Cell. 2015; 161: 18–23. https://doi.org/10.1016/j.cell.2015.02.033 PMID: 25815981

4. Levine ME, Suarez JA, Brandhorst S, Balasubramanian P, Cheng C-W, Madi F, et al. Low Protein Intake is Associated with a Major Reduction in IGF-1, Cancer, and Overall Mortality in the 65 and Younger but Not Older Population. Cell Metab. 2014; 19: 407–417. https://doi.org/10.1016/j.cmet.2014.02.006 PMID: 24606898

5. Solon-Biet SM, McMahon AC, Ballard JWO, Ruohonen K, Wu LE, Cogger VC, et al. The Ratio of Macronutrients, Not Caloric Intake, Dictates Cardiometabolic Health, Aging, and Longevity in Ad Libitum-Fed Mice. Cell Metab. 2014; 19: 418–430. https://doi.org/10.1016/j.cmet.2014.02.009 PMID: 24606899

6. Lee KP, Simpson SJ, Clissold FJ, Brooks R, Ballard JWO, Taylor PW, et al. Lifespan and reproduction in Drosophila: new insights from nutritional geometry. Proc Natl Acad Sci. 2008; 105: 2498–2503. https://doi.org/10.1073/pnas.0710787105 PMID: 18268352

7. Piper MDW, Blanc E, Leitão-Gonçalves R, Yang M, He X, Linford NJ, et al. A holidic medium for Drosophila melanogaster. Nat Methods. 2013; 11: 100–105. https://doi.org/10.1038/nmeth.2731 PMID: 24240321

8. Piper MDW, Partridge L, Raubenheimer D, Simpson SJ. Dietary Restriction and Aging: A Unifying Perspective. Cell Metab. 2011; 14: 154–160. https://doi.org/10.1016/j.cmet.2011.06.013 PMID: 21803286

9. Grandison RC, Piper MDW, Partridge L. Amino-acid imbalance explains extension of lifespan by dietary restriction in Drosophila. Nature. 2009; 462: 1061–1064. https://doi.org/10.1038/nature08619 PMID: 19950092

10. Gosby AK, Conigrave AD, Lau NS, Iglesias MA, Hall RM, Jebb SA, et al. Testing Protein Leverage in Lean Humans: A Randomised Controlled Experimental Study. Morison C, editor. PLoS ONE. 2011; 6: e25929. https://doi.org/10.1371/journal.pone.0025929 PMID: 22022472

11. Griffioen-Roose S, Mars M, Siebelink E, Finlayson G, Tomes D, de Graaf C. Protein status elicits compensatory changes in food intake and food preferences. Am J Clin Nutr. 2012; 95: 32–38. https://doi.org/10.3945/ajcn.111.025053 PMID: 22158729

12. Ribeiro C, Dickson BJ. Sex Peptide Receptor and Neuronal TOR/S6K Signaling Modulate Nutrient Balancing in Drosophila. Curr Biol. 2010; 20: 1000–1005. https://doi.org/10.1016/j.cub.2010.03.061 PMID: 20471268
13. Itskov PM, Ribeiro C. The Dilemma of the Gourmet Fly: The Molecular and Neuronal Mechanisms of Feeding and Nutrient Decision Making in Drosophila. Front Neurosci. 2013; 7: 12. https://doi.org/10.3389/fnins.2013.00012 PMID: 23407678

14. Walker SJ, Corrales-Carvajal VM, Ribeiro C. Postmating Circuitry Modulates Salt Taste Processing to Increase Reproductive Output in Drosophila. Curr Biol. 2015; 25: 2621–2630. https://doi.org/10.1016/j.cub.2015.08.043 PMID: 26412135

15. Vargas MA, Luo N, Yamaguchi A, Kapahi P. A Role for S6 Kinase and Serotonin in Postmating Dietary Switch and Balance of Nutrients in D. melanogaster. Curr Biol. 2010; 20: 1006–1011. https://doi.org/10.1016/j.cub.2010.04.009 PMID: 20471266

16. Corrales-Carvajal VM, Faisal AA, Ribeiro C. Internal states drive nutrient homeostasis by modulating exploration-exploitation trade-off. eLife. 2016; 5: e19920. https://doi.org/10.7554/eLife.19920 PMID: 27770569

17. US Department of Agriculture, Agricultural Research Service, Nutrient Data Laboratory. USDA National Nutrient Database for Standard Reference, Release 28. https://www.ars.usda.gov/northeast-area/beltsville-md/beltsville-human-nutrition-research-center/nutrient-data-laboratory/docs/usda-national-nutrient-database-for-standard-reference/

18. Bos M, Burnet B, Farrow R, Woods RA. Development of Drosophila on sterol mutants of the yeast Saccharomyces cerevisiae. Genet Res. 1976; 28: 163–176. PMID: 795728

19. Subramanian S, Blanton LV, Frese SA, Charbonneau M, Mills DA, Gordon JI. Cultivating Healthy Growth and Nutrition through the Gut Microbiota. Cell. 2015; 161: 327–336. https://doi.org/10.1016/j.cell.2015.03.013 PMID: 25815983

20. Kau AL, Ahern PP, Griffin NW, Goodman AL, Gordon JI. Human nutrition, the gut microbiome, and immune system: envisioning the future. Nature. 2011; 474: 327–336. https://doi.org/10.1038/nature10213 PMID: 20471266

21. Turnbaugh PJ, Ley RE, Mahowald MA, Magrini V, Mardis ER, Gordon JI. An obesity-associated gut microbiome with increased capacity for energy harvest. Nature. 2006; 444: 1027–131. https://doi.org/10.1038/nature05414 PMID: 17183312

22. Ley RE, Turnbaugh PJ, Klein S, Gordon JI. Microbial ecology: Human gut microbes associated with obesity. Nature. 2006; 444: 1022–1023. https://doi.org/10.1038/4441022a PMID: 17183309

23. Sharon G, Garg N, Debelius J, Knight R, Dorrestein PC, Mazmanian SK. Specialized Metabolites from the Microbiome in Health and Disease. Cell Metab. 2014; 20: 719–730. https://doi.org/10.1016/j.cmet.2014.10.016 PMID: 25440054

24. Clemente JC, Ursell LK, Parfrey LW, Knight R. The Impact of the Gut Microbiota on Human Health: An Integrative View. Cell. 2012; 148: 1258–1270. https://doi.org/10.1016/j.cell.2012.01.035 PMID: 22424233

25. Cryan JF, Dinan TG. Mind-altering microorganisms: the impact of the gut microbiota on brain and behaviour. Nat Rev Neurosci. 2012; 13: 701–712. https://doi.org/10.1038/nrn3346 PMID: 22968153

26. Mayer EA, Knight R, Mazmanian SK, Cryan JF, Tillisch K. Gut Microbes and the Brain: Paradigm Shift in Neuroscience. J Neurosci. 2014; 34: 15490–15496. https://doi.org/10.1523/JNEUROSCI.3299-14.2014 PMID: 25392516

27. Sampson TR, Mazmanian SK. Control of Brain Development, Function, and Behavior by the Microbiome. Cell Host Microbe. 2015; 17: 565–576. https://doi.org/10.1016/j.chom.2015.04.011 PMID: 25974299

28. Ezenwa VO, Gerardo NM, Inouye DW, Medina M, Xavier JB. Animal Behavior and the Microbiome. Science. 2012; 338: 198–199. https://doi.org/10.1126/science.1227412 PMID: 23060664

29. Wong AC-N, Holmes A, Ponton F, Lihoireau M, Wilson K, Raubenheimer D, et al. Behavioral Microbiomics: A Multi-Dimensional Approach to Microbial Influence on Behavior. Front Microbiol. 2015; 6: 1359. https://doi.org/10.3389/fmicb.2015.01359 PMID: 26640464

30. Vijay-Kumar M, Aitken JD, Carvalho FA, Cullender TC, Mwangi S, Srinivasan S, et al. Metabolic Syndrome and Altered Gut Microbiota in Mice Lacking Toll-Like Receptor 5. Science. 2010; 328: 228–231. https://doi.org/10.1126/science.1179721 PMID: 20203013

31. Heijtz RD, Wang S, Anuar F, Qian Y, Bjorkholm B, Samuelsson A, et al. Normal gut microbiota modulates brain development and behavior. Proc Natl Acad Sci. 2011; 108: 3047–3052. https://doi.org/10.1073/pnas.1010529108 PMID: 21282636

32. Clarke G, Grenham S, Scully P, Fitzgerald P, Moloney RD, Shanahan F, et al. The microbiome-gut-brain axis during early life regulates the hippocampal serotonergic system in a sex-dependent manner. Mol Psychiatry. 2013; 18: 666–673. https://doi.org/10.1038/mp.2012.77 PMID: 22688187
33. Neufeld KM, Kang N, Bienenstock J, Foster JA. Reduced anxiety-like behavior and central neurochemical change in germ-free mice. Neurogastroenterol Motil. 2011; 23: 255–e119. https://doi.org/10.1111/j.1365-2982.2010.01620.x PMID: 21054680

34. Hsiao EY, McBride SW, Hsien S, Sharon G, Hyde ER, McCue T, et al. Microbiota Modulate Behavioral and Physiological Abnormalities Associated with Neurodevelopmental Disorders. Cell. 2013; 155: 1451–1463. https://doi.org/10.1016/j.cell.2013.11.024 PMID: 24315484

35. Sharon G, Segal D, Ringo JM, Heftez A, Zilber-Rosenberg I, Rosenberg E. Commensal bacteria play a role in mating preference of Drosophila melanogaster. Proc Natl Acad Sci. 2010; 107: 20051–20056. https://doi.org/10.1073/pnas.1009906107 PMID: 21041648

36. Huang J-H, Douglas AE. Consumption of dietary sugar by gut bacteria determines Drosophila lipid content. Biol Lett. 2015; 11: 20150469. https://doi.org/10.1098/rsbl.2015.0469 PMID: 26912894

37. Storelli G, Defaye A, Erkosar B, Hols P, Royet J, Leulier F. Lactobacillus plantarum Promotes Drosophila Systemic Growth by Modulating Hormonal Signals through TOR-Dependent Nutrient Sensing. Cell Metab. 2011; 14: 403–414. https://doi.org/10.1016/j.cmet.2011.07.012 PMID: 21907145

38. Shin SC, Kim S-H, You H, Kim B, Kim AC, Lee K-A, et al. Drosophila Microbiome Modulates Host Developmental and Metabolic Homeostasis via Insulin Signaling. Science. 2011; 334: 670–674. https://doi.org/10.1126/science.1212782 PMID: 22053049

39. Schwarzer M, Makki K, Storelli G, Machuc a-Gayet I, Srutkova D, Hermanova P, et al. Lactobacillus plantarum strain maintains growth of infant mice during chronic undernutrition. Science. 2016; 351: 854–857. https://doi.org/10.1126/science.aad8588 PMID: 26912894

40. Smith MI, Yatsunenko T, Manary MJ, Trehan I, Mkakosya R, Cheng J, et al. Gut Microbiomes of Malawian Twin Pairs Discordant for Kwashiorkor. Science. 2013; 339: 548–554. https://doi.org/10.1126/science.1229000 PMID: 23636771

41. Tanimura T, Isono K, Takamura T, Shimada I. Genetic dimorphism in the taste sensitivity to trehalose in Drosophila melanogaster. J Comp Physiol. 1982; 147: 433–437.

42. Toshima N, Tanimura T. Taste preference for amino acids is dependent on internal nutritional state in Drosophila melanogaster. J Exp Biol. 2012; 215: 2827–2832. https://doi.org/10.1242/jeb.069146 PMID: 22837455

43. Bender DA. Amino Acid Metabolism. John Wiley & Sons; 2012.

44. Wu G, Wu Z, Dai Z, Yang Y, Wang W, Liu C, et al. Dietary requirements of “nutritionally non-essential amino acids” by animals and humans. Amino Acids. 2012; 44: 1107–1113. https://doi.org/10.1007/s00726-012-1444-2 PMID: 23247926

45. Simpson CL, Simpson SJ, Abisgold JD. The role of various amino acids in the protein compensatory response of locusta migratoria. Symposia Biologica Hungarica 1991; 39: 39–46.

46. Hara K, Yonezawa K, Weng Q-P, Kozlowski MT, Belham C, Avruch J. Amino Acid Sufficiency and mTOR Regulate p70 S6 Kinase and eIF-4E BP1 through a Common Effector Mechanism. J Biol Chem. 1998; 273: 14484–14494. PMID: 9603962

47. Ja WW, Carvalho GB, Mak EM, de la Rosa NN, Fang AY, Liong JC, et al. Prandiology of Drosophila and the CAFE assay. Proc Natl Acad Sci. 2007; 104: 8253–8256. https://doi.org/10.1073/pnas.0702726104 PMID: 17494737

48. Silva FJ, Bel Y, Botella LM, Cotton RGH, Ferré J. Immunological detection of phenylalanine hydroxylase protein in Drosophila melanogaster. Biochem J. 1992; 287: 85–89. PMID: 1417795

49. Fölling A. Über Ausscheidung von Phenylbrenztraubensäure in den Harn als Stoffwechselanomalie in Verbindung mit Imbezillität. Hoppe-Seyler’s Z Für Physiol Chem. 1934; 227: 169–181.

50. Mitchell JJ, Trakadis YJ, Scriver CR. Phenylalanine hydroxylase deficiency. Genet Med Off J Am Coll Med Genet. 2011; 13: 697–707.

51. de Groot MJ, Hoekema M, Blau N, Reijngoud DJ, van Sproons FJ. Pathogenesis of cognitive dysfunction in phenylketonuria: Review of hypotheses. Mol Genet Metab. 2010; 99, Supplement: S86–S89.

52. MacDonald A, Rocha JC, van Rijn M, Feillet F. Nutrition in phenylketonuria. Mol Genet Metab. 2011; 104: S10–S18. https://doi.org/10.1016/j.ygme.2011.08.023 PMID: 21944460

53. Vajro P, Strisciuglio P, Houssin D, Huault G, Laurent J, Alvarez F, et al. Correction of Phenylketonuria after Liver Transplantation in a Child with Cirrhosis. N Engl J Med. 1993; 329: 363–363.

54. Udenfriend S, Cooper JR. The enzymatic conversion of phenylalanine to tyrosine. J Biol Chem. 1952; 194: 503–511. PMID: 14927641

55. Kilby BA, Neville E. Amino-acid metabolism in locust tissues. J Exp Biol. 1957; 34: 276–289.

56. Arrese EL, Soulages JL. Insect Fat Body: Energy, Metabolism, and Regulation. Annu Rev Entomol. 2010; 55: 207–225. https://doi.org/10.1146/annurev-ento-112408-085356 PMID: 19725772
57. Neckameyer WS, Coleman CM, Eadie S, Goodwin SF. Compartmentalization of neuronal and peripheral serotonin synthesis in Drosophila melanogaster. Genes Brain Behav. 2007; 6: 756–769. https://doi.org/10.1111/j.1601-183X.2007.00307.X PMID: 17376153

58. Schmid MR, Anderl I, Vesala L, Vanha-aho L-M, Deng X-J, Rämet M, et al. Control of Drosophila Blood Cell Activation via Toll Signaling in the Fat Body. PLoS ONE. 2014; 9: e102568. https://doi.org/10.1371/journal.pone.0102568 PMID: 25102059

59. Hacquard S, Garrido-Oter R, González A, Spaepen S, Ackermann G, Lebeis S, et al. Microbiota and Host Nutrition across Plant and Animal Kingdoms. Cell Host Microbe. 2015; 17: 603–616. https://doi.org/10.1016/j.chom.2015.04.009 PMID: 25974302

60. Wong AC-N, Dobson AJ, Douglas AE. Gut microbiota dictates the metabolic response of Drosophila to diet. J Exp Biol. 2014; 217: 1894–1901. https://doi.org/10.1242/jeb.101725 PMID: 24577449

61. Blum JE, Fischer CN, Miles J, Handelsman J. Frequent Replenishment Sustains the Beneficial Microbiome of Drosophila melanogaster. mBio. 2013; 4: e00860–13. https://doi.org/10.1128/mBio.00860-13 PMID: 24194543

62. Ryu J-H, Kim S-H, Lee H-Y, Bai JY, Nam Y-D, Bae J-W, et al. Innate Immune Homeostasis by the Homeobox Gene Caudal and Commensal-Gut Mutualism in Drosophila. Science. 2008; 319: 777–782. https://doi.org/10.1126/science.1149357 PMID: 18218863

63. Cox CR, Gilmore MS. Native Microbial Colonization of Drosophila melanogaster and Its Use as a Model of Enterococcus faecalis Pathogenesis. Infect Immun. 2007; 75: 1565–1576. https://doi.org/10.1128/IAI.01496-06 PMID: 17220307

64. Hansen IA, Attardo GM, Park J-H, Peng Q, Raikhel AS. Target of rapamycin-mediated amino acid signaling in mosquito anautogeny. Proc Natl Acad Sci U S A. 2004; 101: 10626–10631. https://doi.org/10.1073/pnas.0403460101 PMID: 15229322

65. Drummond-Barbosa D, Spradling AC. Stem Cells and Their Progeny Respond to Nutritional Changes during Drosophila Oogenesis. Dev Biol. 2001; 231: 265–278. https://doi.org/10.1006/dbio.2000.0135 PMID: 11180967

66. Hosios AM, Hecht VC, Danai LV, Johnson MO, Rathmell JC, Steinhauer ML, et al. Amino Acids Rather than Glucose Account for the Majority of Cell Mass in Proliferating Mammalian Cells. Dev Cell. 2016; 36: 540–549. https://doi.org/10.1016/j.devcel.2016.02.012 PMID: 26954548

67. Itskov PM, Moreira J-M, Vinnik E, Lopes G, Safarik S, Dickinson MH, et al. Automated monitoring and quantitative analysis of feeding behaviour in Drosophila. Nat Commun. 2014; 5: 4560. https://doi.org/10.1038/ncomms5560 PMID: 25087594

68. Newell PD, Douglas AE. Interspecies Interactions Determine the Impact of the Gut Microbiota on Nutrient Allocation in Drosophila melanogaster. Appl Environ Microbiol. 2014; 80: 788–796. https://doi.org/10.1128/AEM.02742-13 PMID: 24242251

69. Douglas AE. Nutritional interactions in insect-microbial symbioses: aphids and their symbiotic bacteria Buchnera. Annu Rev Entomol. 1998; 43: 17–37. https://doi.org/10.1146/annurev.ento.43.1.17 PMID: 15012383

70. Bjordal M, Arquier N, Kniazeff J, Pin JP, Léopold P. Sensing of Amino Acids in a Dopaminergic Circuitry Promotes Rejection of an Incomplete Diet in Drosophila. Cell. 2014; 156: 510–521. https://doi.org/10.1016/j.cell.2013.12.024 PMID: 24485457

71. Hao S. Uncharged tRNA and Sensing of Amino Acid Deficiency in Mammalian Piriform Cortex. Science. 2005; 307: 1776–1777. https://doi.org/10.1126/science.1104882 PMID: 15774759

72. Maurin A-C, Jousse C, Averous J, Parry L, Bruhat A, Cherasse Y, et al. The Gcn2 kinase biases feeding behavior to maintain amino acid homeostasis in omnivores. Cell Metab. 2005; 1: 273–277. https://doi.org/10.1016/j.cmet.2005.03.004 PMID: 16054071

73. Leib DE, Knight ZA. Re-examination of Dietary Amino Acid Sensing Reveals a Gcn2-Independent Mechanism. Cell Rep. 2015; 13: 1081–1089. https://doi.org/10.1016/j.celrep.2015.09.055 PMID: 26526991

74. Simpson SJ, Simpson CL. Mechanisms Controlling Modulation by Haemolymph Amino Acids of Gustatory Responsiveness in the Locust. J Exp Biol. 1992; 168: 269–287.

75. Brummel T, Ching A, Seroude L, Simon AF, Benzer S. Drosophila lifespan enhancement by exogenous bacteria. Proc Natl Acad Sci U S A. 2004; 101: 12974–12979. https://doi.org/10.1073/pnas.0405207101 PMID: 15322271

76. Elgart M, Stern S, Salton O, Gnanis K, Heifetz Y, Soen Y. Impact of gut microbiota on the fly’s germ line. Nat Commun. 2016; 7: 11280. https://doi.org/10.1038/ncomms11280 PMID: 27080728

77. Yamada R, Deshpande SA, Bruce KD, Mak EM, Ja WW. Microbes Promote Amino Acid Harvest to Rescue Undernutrition in Drosophila. Cell Rep. 2015; 10: 865–872.
78. Erkosar B, Storelli G, Mitchell M, Bozonnset L, Bozonnset N, Leulier F. Pathogen Virulence Impedes Mutualist-Mediated Enhancement of Host Juvenile Growth via Inhibition of Protein Digestion. Cell Host Microbe. 2015; 18: 445–455. https://doi.org/10.1016/j.chom.2015.09.001 PMID: 26439865

79. Metges CC. Contribution of microbial amino acids to amino acid homeostasis of the host. J Nutr. 2000; 130: 1857S–1864S. PMID: 10867063

80. Attardo GM, Higgs S, Klingler KA, Vanlandingham DL, Raikhel AS. RNA interference-mediated knockdown of a GATA factor reveals a link to anautogeny in the mosquito Aedess aegypti. Proc Natl Acad Sci. 2003; 100: 13374–13379. https://doi.org/10.1073/pnas.2235649100 PMID: 14595016

81. Arsic D, Guerin PM. Nutrient content of diet affects the signaling activity of the insulin/target of rapamycin/p70 S6 kinase pathway in the African malaria mosquito Anopheles gambiae. J Insect Physiol. 2008; 54: 1226–1235. https://doi.org/10.1016/j.jinsphys.2008.06.003 PMID: 18634792

82. Ables ET, Laws KM, Drummond-Barbosa D. Control of adult stem cells in vivo by a dynamic physiological environment: diet-dependent systemic factors in Drosophila and beyond. Wiley Interdiscip Rev Dev Biol. 2012; 1: 657–674. https://doi.org/10.1002/wdev.48 PMID: 23799567

83. LaFever L, Feoktistov A, Hsu H-J, Drummond-Barbosa D. Specific roles of Target of rapamycin in the control of stem cells and their progeny in the Drosophila ovary. Development. 2010; 137: 2117–2126. https://doi.org/10.1242/dev.050351 PMID: 20504961

84. Theis KR, Dheilly NM, Klassen J-L, Brucker RM, Bosch TCG, et al. Getting the Hologenome Concept Right: an Eco-Evolutionary Framework for Hosts and Their Microbiomes. mSystems. 2016; 1: e00028–16.

85. Erkosar B, Storelli G, Defaye A, Leulier F. Host-Intestinal Microbiota Mutualism: “Learning on the Fly.” Cell Host Microbe. 2013; 13: 8–14. https://doi.org/10.1016/j.chom.2013.12.004 PMID: 23332152

86. Lee W-J, Brey PT. How Microbiomes Influence Metazoan Development: Insights from History and Drosophila Modeling of Gut-Microbe Interactions. Annu Rev Cell Dev Biol. 2013; 29: 571–592. https://doi.org/10.1146/annurev-cellbio-101512-122333 PMID: 23808845

87. Novarino G, El-Fishawy P, Kayseril H, Meguid NA, Scott EM, Schroth J, et al. Mutations in BCKD-kinase Lead to a Potentially Treatable Form of Autism with Epilepsy. Science. 2012; 338: 394–397. https://doi.org/10.1126/science.1224631 PMID: 22956686

88. Tärlungeanu DC, Deliu E, Dotter CP, Kara M, Janiesch PC, Scalise M, et al. Impaired Amino Acid Transport at the Blood Brain Barrier Is a Cause of Autism Spectrum Disorder. Cell. 2016; 167: 1481–1494.e18. https://doi.org/10.1016/j.cell.2016.11.013 PMID: 27912058

89. Lee T, Luo L. Mosaic Analysis with a Repressible Cell Marker for Studies of Gene Function in Neuronal Morphogenesis. Neuron. 1999; 22: 451–461. PMID: 10197526

90. Luo L, Liao YJ, Jan LY, Jan YN. Distinct morphogenetic functions of similar small GTPases: Drosophila Drac1 is involved in axonal outgrowth and myoblast fusion. Genes Dev. 1994; 8: 1787–1802. PMID: 795857

91. Förster D, Luschnig S. Src42A-dependent polarized cell shape changes mediate epithelial tube elongation in Drosophila. Nat Cell Biol. 2012; 14: 526–534. https://doi.org/10.1038/ncb2456 PMID: 22446736

92. Asha H, Nagy I, Kovacs G, Stetson D, Ando I, Dearolf CR. Analysis of Ras-Induced Overprolife ration in Drosophila Hemocytes. Genetics. 2003; 163: 203–215. PMID: 12586708

93. Piper MD, Soultoukis GA, Blanc E, Mesaros A, Herbert SL, Juridic P, et al. Matching Dietary Amino Acid Balance to the In Silico Translated Exome Optimizes Growth and Reproduction without Cost to Lifespan. Cell Metab. 2017; 25: 610–621. https://doi.org/10.1016/j.cmet.2017.02.005 PMID: 28273481

94. Wayland MT, Defaye A, Rocha J, Jayaram SA, Royet J, Miguel-Alia ga I, et al. Spotting the differences: Probing host/microbiota interactions with a dedicated software tool for the analysis of faecal outputs in Drosophila. J Insect Physiol. 2014; 69: 126–135. https://doi.org/10.1016/j.jinsphys.2014.05.023 PMID: 24907675