Host genetic determinants of microbiota-dependent nutrition revealed by genome-wide analysis of *Drosophila melanogaster*

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Animals bear communities of gut microorganisms with substantial effects on animal nutrition, but the host genetic basis of these effects is unknown. Here we use *Drosophila* to demonstrate substantial among-genotype variation in the effects of eliminating the gut microbiota on five host nutritional indices (weight, protein, lipid, glucose and glycogen contents); this includes variation in both the magnitude and direction of microbiota-dependent effects. Genome-wide association studies to identify the genetic basis of the microbiota-dependent variation reveal polymorphisms in largely non-overlapping sets of genes associated with variation in the nutritional traits, including strong representation of conserved genes functioning in signalling. Key genes identified by the GWA study are validated by loss-of-function mutations that altered microbiota-dependent nutritional effects. We conclude that the microbiota interacts with the animal at multiple points in the signalling and regulatory networks that determine animal nutrition. These interactions with the microbiota are probably conserved across animals, including humans.
Animal phenotypes are strongly influenced by microorganisms that colonize their surfaces (for example, the skin, gut and reproductive tract) and sometimes internal organs. Some microbial effects can be attributed to specific microbial functions, for example, synthesis of specific nutrients or protective toxins. Other microbial effects on the host, including promotion of intestinal homeostasis, immunity and metabolic function, involve complex networks of interactions between the animal host and microbiota. These complex interactions have been interpreted as evidence that animal regulatory networks are structured to function in the context of the resident microbiota, with the implication that host health and vigour can be prejudiced by mismatch between host function and the composition or activities of the microbiota, a condition known as dysbiosis.

The purpose of this study was to quantify how the effect of the microbiota on host phenotype varies with host genotype, and to elucidate the genetic bases of these microbiota-dependent host traits. This issue has not been addressed directly for any system, even though it has important implications for our understanding of the genetic basis of human diseases linked to microbiota and can potentially make significant contributions to the development of personalized microbial therapies. More generally, understanding how the microbiota-dependent phenotype maps onto the host genotype will enrich our understanding of the evolution and function of interactions between animals and their resident microbiota.

Our research was conducted on the fruit fly Drosophila melanogaster and its gut microbiota, which is ideally suited for the study of microbiota-dependent effects for three reasons. First, experimental analysis is facilitated by robust methods to eliminate the gut microbiota by egg dechorionation, yielding axenic flies. This treatment does not affect the complement of the intracellular bacterium Wolbachia, which is present in many Drosophila lines and is vertically transmitted via the egg cytoplasm. Second, axenic individuals of Drosophila studied to date commonly display readily quantified nutritional traits, including elevated levels of indices of triglyceride, glycogen or free glucose, and these changes have been linked to altered function of the nutrient-sensing insulin/insulin-like peptide signalling (IIS) and target of rapamycin (TOR) signalling pathways that couple organisal growth to nutrient supply. Finally, the super-gene resource for Drosophila can be harnessed to interrogate the genetic architecture of microbiota-dependent effects. In particular, the Drosophila Genetic Reference Panel (DGRP) of inbred isofemale lines with sequenced genomes enable genotype–phenotype mapping by genome-wide association (GWA) and candidate lines with sequenced genomes enable genotype–phenotype mapping by genome-wide association (GWA) and candidate lines with sequenced genomes enable genotype–phenotype mapping by genome-wide association (GWA) and candidate lines with sequenced genomes enable genotype–phenotype mapping by genome-wide association (GWA). To standardize the microbiota in the test DGRP lines, this study was conducted on flies generated from dechorionated eggs and exposed to isolates of five bacterial species that were isolated from Drosophila guts, are found ubiquitously in association with laboratory-cultured and wild-caught Drosophila and, in combination, have been shown to restore the nutritional phenotype of Drosophila bearing its unmanipulated microbiota.

This study focused on the nutritional effects of the Drosophila microbiota. Using the DGRP, we demonstrated substantial among-line variation in nutritional response to elimination of the microbiota and identified host genetic variants (single-nucleotide polymorphisms, SNPs) associated with the microbiota-dependent nutritional traits. Many of the genes identified have fundamental roles in cell signalling and control of gene expression that are conserved in many animals, including humans. Our results show the central role of host genotype in shaping the impact of microbiota on animal nutrition.

**Results**

**Genetic variation in microbiota-dependent nutritional traits.**

Gnotobiotic Drosophila associated with a defined five-species microbiota and axenic (microbe-free) Drosophila differed significantly in all of the five nutritional traits tested (Fig. 1). The global mean values of dry weight per fly were depressed in axenic flies and, after controlling for weight, axenic flies also had reduced glycogen and protein contents but elevated glucose and triglyceride (TAG) contents (Fig. 1a and Supplementary Table 1). Nevertheless, the among-line variation in the response to elimination of the microbiota was appreciable for all traits, and genotype accounted for 31%–73% of the total variance (Fig. 1b) and within-line variation was low (Supplementary Fig. 1). For every trait, the variation included lines that differed in response direction, that is, every trait was elevated in axenic flies of some lines, but decreased in other lines (Fig. 1c). Among the lines tested, 51% bore the intracellular bacterium Wolbachia, which was not eliminated by the egg dechorionation procedure used to generate axenic flies. The Wolbachia-positive lines had significantly elevated glycogen content relative to Wolbachia-free lines in both axenic and gnotobiotic flies (Fig. 1), perhaps due to sensitization to insulin signalling.

For each trait, the mean values were significantly correlated across the axenic and gnotobiotic flies (weight \( r = 0.8; \) protein/weight \( r = 0.61; \) glucose/weight \( r = 0.46; \) glycogen/weight \( r = 0.53; \) TAG/weight \( r = 0.77; \) all \( P < 0.001)\), reinforcing the evidence above for a genetic component to these traits. We also tested for correlations among the various phenotypic traits (Supplementary Table 2). No traits were significantly correlated in axenic flies, irrespective of Wolbachia status; this result indicates that generalized among-genotype variation in sensitivity to elimination of the microbiota is not a major driver of the microbiota-dependent effects. Among gnotobiotic flies, Wolbachia status determined which traits were correlated: TAG was positively correlated with glucose and glycogen contents in Wolbachia-positive gnotobiotic flies, but this effect was absent in Wolbachia-free flies (Supplementary Table 2). The data did not reveal specific lines whose nutritional status was globally hypersensitive to elimination of the microbiota, evidenced by the absence of positive correlations among response indices for each trait (Supplementary Fig. 2).

Collectively, these results demonstrate that Drosophila nutrition has a strong genetic component, that trait values are influenced by the microbiota and that there is genetic variation in the microbiota-dependence of Drosophila nutritional traits. Furthermore, the variation in the response of the various traits to elimination of the microbiota does not appear to be coupled.

**Genotype–phenotype associations.** To determine the association between SNPs, microbiota and the nutritional traits, the interactive effect of microbiota and SNP was tested for biallelic SNP markers in the Drosophila genome. We determined SNP significance by post-hoc corrections for GWA. The top ranking 0.01% of SNPs for each trait (Supplementary Fig. 3) comprised 184–226 SNPs, including SNPs associated with 77–144 genes, with a total of 436 genes mapped across the full data set
(Supplementary Table 3). Linkage disequilibrium between these SNPs was generally low, apart from an island of linked SNPs on chromosome 21 associated with variation in glycogen stores (Supplementary Fig. 4). The mapped SNPs were found more commonly in genic regions than expected, based on genome background (number of SNPs < 5,000 bp from known genes: mapped SNPs = 91%, genome background = 57%, proportions test \( P < 0.0001 \)). Significant gene ontology assignments (false discovery rate \( \leq 0.05 \)) and the loci responsible for assignments are presented in Supplementary Table 4.

To gain an overview of the biological functions potentially determining the host phenotypic responses to the microbiota, we interrogated known functions of genes associated with the 20 most significant SNPs found by each GWA (Supplementary Table 3). Associations between microbiota-dependent variation in dry weight and SNPs included a SNP in the transcription factor Lim3, a SNP in dpr10 (a predicted chemosensor enriched in expression in the brain) and two SNPs in the translational modifier boule (bol). Microbiota-dependent variation in protein content was associated with multiple SNPs in varicose, which has known roles in trachea development\(^ {28} \), and variation in both dry weight and protein are associated with SNPs in trol, a modulator of various signalling pathways, including fibroblast growth factor, hedgehog and Wnt\(^ {29,30} \). Microbiota-dependent variation in TAG was associated with SNPs in genes coding a number of central regulatory elements including the kinase happyhour (hpy; negative regulator of epidermal growth factor receptor signalling and a kinase of JNK and IIS/TOR signalling), and the genes coding the tyrosine protein phosphatase Ptp99A, highwire (hiw, a member of the BMP signaling pathway) and rugose (rg), homologue of the mammalian Neurobeachin, which regulates neurotransmitter receptor trafficking implicated in associative learning\(^ {31,32} \), as well as the transcription factors klumpfuss (klu) and scribbler (sbb). The candidate functional significance of IIS/TOR signalling indicated by the SNP in hpy is strengthened by lower-ranked associations of TAG with SNPs in two genes S6K (ranked 114) and melted (136) coding canonical IIS/TOR proteins downstream of hpy, as well as four further hpy SNPs (122, 124, 138 and 139). The most significant SNP associated with microbiota-dependent variation in glucose content was in slt
(slh), the ligand of roundabout signalling that controls intestinal stem cell fate. Other SNPs in this association included dunce (dcn), a cAMP-phosphodiesterase, which is involved in regulation of multiple behaviours, and the nicotinic acetylcholine receptor nAChR62. Glycogen showed particularly strong associations with the JAK-STAT activated transcription factor chimino, the epidermal growth factor receptor-responsive transcription factor seven up (svp), as well as the morphogen numb (an antagonist of Notch signalling) and CG5867, which contains a protein domain for juvenile hormone binding.

In total, SNPs linked to just 26 genes (6% of the 436 genes) were associated with more than one trait (Supplementary Table 5) and no gene mapped to more than two traits. The low incidence of genes associated with multiple traits is consistent with the responses of traits to removal of the microbiota (Supplementary Table 2).

Functional validation of GWA. The validation of associations between SNPs and microbiota-dependent phenotypic traits focused on genes linked to mapped SNPs. We reasoned that if a SNP correctly inferred the role of a gene, then the trait difference between corresponding flies with a null mutation and wild-type flies could be obtained in either axenic flies or gnotobiotic flies, but not in both treatments. Therefore, our definition of a validated effect of mutation is a statistically significant difference between the mutant and background in one but not both of the treatments. Therefore, our definition of a validated effect of mutation is a statistically significant difference between the mutant and background in one but not both of the axenic and gnotobiotic flies, with a between the mutant and background in one but not both of the treatments. Therefore, our definition of a validated effect of mutation is a statistically significant difference between the mutant and background in one but not both of the axenic and gnotobiotic flies, with a P-value \( \leq 0.025 \) (correcting for two comparisons) by a mixed-effects model. By this criterion, 9 (56%) of 16 predicted microbiota-dependent effects of the tested genes on TAG, glucose and glycogen were validated (Table 1 and Fig. 2). Some off-target effects were anticipated because mutations are predicted to cause greater disruption of genetic networks than natural polymorphisms. In total, four (27%) of 15 tests for off-target effects of genes validated for nutritional traits predicted in the GWAS study (GWAS) yielded significant effects (Table 1, Supplementary Table 6 and Supplementary Fig. 5). There were two classes of response to microbiota in validated mutants: the microbiota either suppressed or enhanced the effects of some mutations, such that the ratio of response (Fig. 2) was significantly greater than unity (for example, Dscam3, mthl1 and rg for TAG), or the microbiota promoted the effects of mutations on the host phenotypic trait (for example, glucose levels for CG32264, CG30288 and rg).

Taken together, these results provide experimental verification that GWAS is an effective route to identify genes with significant microbiota-dependent effects on nutritional traits of Drosophila.

Discussion
This study has demonstrated substantial effects of host genetics on the nutritional response of Drosophila to elimination of the gut microbiota. For every index tested, both the magnitude and direction of the response varied among the Drosophila lines. For example, previous reports of elevated TAG and glucose in axenic flies of the Drosophila line Canton S background are consistent with results obtained for most of the DGRP lines studied here, but a minority displayed the reverse response, and both positive and negative responses to elimination of the microbiota in validated mutants: the microbiota either suppressed or enhanced the effects of some mutations, such that the ratio of response (Fig. 2) was significantly greater than unity (for example, Dscam3, mthl1 and rg for TAG), or the microbiota promoted the effects of mutations on the host phenotypic trait (for example, glucose levels for CG32264, CG30288 and rg).
microbiota were well represented among the lines for protein content and glycogen content. These results are complemented by the mutant analysis and suggest that interactions between the microbiota and host processes that regulate nutrient allocation are not tightly constrained and are responsive to genetic variation in the host. Host genotypic effects on microbiota-dependent nutritional traits may be general among animals, including the human and mouse, for which some traits of the microbiota are already known to vary with host genotype.26–38

A key result of this study is that the effects of eliminating the microbiota on different nutritional indices are not correlated. This suggests that although nutrient allocation patterns are coordinately regulated by integrated signalling circuits, the genetic variation that determines the responsiveness of these networks to the microbiota may have effects specific to individual nutritional indices. Fully consistent with this finding, GWAs revealed that variation in each nutritional trait mapped predominantly to distinct sets of loci and functional processes. Although an experimental design with greater statistical power may have yielded evidence for more pleiotropic genes, these results suggest that many of the genes with microbiota-dependent effects on nutrition have low pleiotropy and raise the possibility that many of the genes, which influence multiple nutritional indices, may generally be microbiota unresponsive.

The functional validations using mutant Drosophila provided a critical test for the reliability of the GWAs in predicting genes with microbiota-dependent effects on nutritional traits. These tests confirmed that host nutrition is determined by an interaction of the microbiota and host genotype. The nature of this interaction appears to depend on the gene in question, as the microbiota suppressed nutritional effects of some mutations, but were required to detect nutritional effects of other mutations. The mutants commonly had the effects predicted by GWAs. However, our ability to make statements about trait-specific effects of genes mapped by GWA is tempered by nonspecific effects of mutants of some genes on certain phenotypic traits. Possible technical causes of discrepancies between mutants and GWAs are false negatives in the GWAs and artefacts of mutants’ genetic backgrounds. Possible biological causes include off-target effects of mutants, owing to potentially greater disruption of genetic networks by transgenic mutation than by natural polymorphism or the related issue of SNPs having tissue-specific effects on gene function that are not mirrored by mutations. We therefore interpret the results of our functional validations as consistent with the absence of a signal of pleiotropy from GWA mappings, with the caveat that 27% of possible effects were not predicted by GWAs.

Successful validation of mutants is particularly relevant in the context of two limitations of the genome-wide analyses: poor definition of the critical probability for GWAS due to the ineligibility of standard corrections for multiple statistical comparisons27,39 and inflated P-values identified in QQ plots (Supplementary Fig. 6), potentially indicative of mis-specification in GWA models (Supplementary Note 1). We found that the microbiota-dependent effects for each trait were more frequently abrogated in Drosophila bearing mutations in genes mapped by specific GWAS than in collective pools for the individual genes of interest, confirming that mapped genes are microbiota-responsive (Fig. 2 and Table 1). We conclude that the validation studies of mutants support the genome-wide mappings as a strategy to generate hypotheses about the genetics underlying variation in impacts of the microbiota on nutrition.

The GWAs identified genes functioning in IIS/TOR signalling, a network identified previously to interact with the gut microbiota.17,18 IIS/TOR signalling is a nexus of nutrition and growth (for example, see refs 40,41), but with extensive cross-talk with other pathways involved in regulation of growth. The identification of genes contributing to multiple signalling pathways other than IIS/TOR indicates that the effects of the microbiota on host phenotype cannot be described exclusively in terms of any single defining pathway; instead, our results are congruent with the view that the microbiota interact at multiple points in the signalling and regulatory networks that define animal function. These data provide evidence for the argument made previously, but largely without supporting data, that many animal–microbiota interactions are probably conserved across the animal kingdom, including in humans, because the associations with resident microbiota are ubiquitous among animals and important for animal health and vigour.

Immediately relevant to these considerations are genes with no known function, accounting for 43% of mapped genes. The nutritional functions of many of these genes may be microbiota dependent and may have gone undetected by traditional genetic screens, which do not control for microbiota effects. Experimental analyses that compare axenic hosts with hosts that are colonized with different microbial partners could promote understanding of these currently un-annotated genes. The absence of canonical immunity genes or immunity-related gene ontology terms significantly associated with any microbiota-dependent trait is also of interest. Various immune effectors have been implicated in the regulation of interactions with the microbiota in Drosophila,42,43 and other animals,44–47, raising the possibility that genetic variation in immunity may shape the microbiota in a manner that affects traits other than the nutrition-related traits studied here, or this variation is not exposed in associations with a specified microbiota. Our GWA analyses necessarily did not account for variation in the hosts’ ability to associate with the microbiota. The replication of host genotypes and properties of the microbiota (for example, total abundance and relative abundance) required to distinguish the effects of these two factors on host phenotype would require thousands of Drosophila lines. A key objective of future studies should be to map genetic variation in the predisposition of Drosophila for certain microbial communities and to characterize any correspondence between such variation and nutrition.

The GWA mappings elucidated in this study offer a rich inventory of genes with candidate roles in microbiota-dependent traits. Although our experimental design precludes detailed analysis of the evolutionary forces shaping this variation, because the inbred DGRP lines bear unnaturally segregated allelic combinations that expose variants, which may not be subject to selection under field conditions, the host genes identified probably include targets of microbiota-dependent selection. Future research to identify the mode of action and evolutionary significance of genes with microbiota-dependent function will need to account for the complexity of the interactions: that the impacts of the microbiota on host traits are strongly influenced by host genotype and involve variation in multiple genes with largely independent effects on different phenotypic traits.

Methods

The flies. Drosophila from the DGRP19 and transposon (P-element insertion) mutant flies obtained from the Bloomington Drosophila Resource Center, IN, USA (Supplementary Table 7) were cultured at 25 °C on a 12 h:12 h light–dark cycle on yeast–glucose medium (11 water, 100 g brewer’s yeast (MP Biomedicals 903312), 100 g glucose (Sigma 158968), 1.2% agar (Apex 66–103), 0.84% propionic acid, 0.08% phosphoric acid). Axenic and gnotobiotic flies were generated by allowing flies from stock cultures to oviposit overnight on grape juice agar (1 litre water, 100 g yeast, 100 g glucose, 113 ml Welch’s concentrated grape juice). The eggs were picked from the agar with a paint brush and dechorionated by 5 min of washing in 0.6% hypochlorite (10% Clorox brand bleach), to remove all microorganisms associated with the egg surface (but not the cytoplasmic Wolbachia), followed by three rinses with sterile water. To produce axenic flies, the eggs were added to 7.5 ml sterile yeast–glucose food medium in 50 ml Falcon tubes (Fisher Scientific) using aseptic technique in a laminar flow cabinet. To produce gnotobiotic flies,
A second set of three pools of 5 CO2-anaesthetized flies were homogenized in 125 ml per ml) with a sterile pestle, incubated at 37°C overnight at 30°C, excluding any row of the data matrix with missing data. Phenotypic traits were analysed with mixed models using the lme function from the nlme library, excluding any row of the data matrix with missing data. Mixed models were used to control for genotype nested within experimental block as a random factor. Differences in glycerol content between the TAG-digested and -undigested was immediately flash frozen for subsequent protein quantification, the remaining was high congruence between our detection of Wolbachia status-matched background stock, by a linear mixed model (lme in R version 3.0.2) accounting for genotype (mutant versus background) as a fixed effect and (n = 3, 3 replicates per experiment) as a random effect. The effect was quantified using the BioRad DC kit (BioRad 500-0111). Glucose and glycogen were quantified with the Sigma Glucose assay kit (Sigma-Aldrich G2020-1KT): glucose homogenates were assayed directly from TE-buffered homogenates, whereas glycogen was digested to glucose in the presence of 20 μg amylloglucosidase (Sigma-A7420 at 1 U/ml) before submitting the previous glucose reading. TAG was quantified using the Sigma Triglyceride assay kit (TR100-1KT), in which glycerol content is determined before and after digestion with supplied Triglyceride Reagent. TAG content was determined as difference in glycerol content between the TAG-digested and -undigested measurements of glycerol.

Wolbachia status of the flies. DNA was isolated from adult flies by the salting out method for most DGRP lines used in this study and for all P-element insertion mutants and backgrounds. Flies of five males were homogenized in 180 μl lysis buffer (20 mM Tris-HCl, 2 mM EDTA, 1.2% Triton X-100, with 200 μg lysosome per ml) with a sterile pestle, incubated at 37°C for 60 min and vortexed with 100-μl 0.1-mm glass beads (Scientific Industries SH-606) for 5 min. Twenty microliters of 50 mM sodium phosphate buffer (pH 7.0) and 5 μl of 5% (w/v) SDS was added to the samples, which were mixed with 10 μl 20 mg proteinase K per ml (Qiagen) and incubated at 55°C for 60 min. One hundred microliters of 3 M sodium acetate (pH 5.2) was added to each sample before incubation at 25°C for 10 min. Samples were centrifuged for 5 min at 5,000 g and 4°C. Three hundred microliters of isopropanol (cooled to −20°C) were then added to the supernatant, mixed and then incubated at room temperature for 30 min. Following 30 min centrifugation at 5,000 g at room temperature, the pellets were rinsed in 500 μl 70% ethanol (cooled to 20°C), vortexed, centrifuged for 5 min at 5,000g at room temperature and resuspended in sterile Milli-Q water. Wolbachia Status was determined by quantitative real-time PCR using three primer sets, including Wsp18F and Wsp18R, Sigma-AAT TGA TGA TGA AAG AAC-3′, respectively, which were used for specific detection of Wolbachia in duplicate samples. PCR reactions (25 μl) contained 20.575 μl ddH2O, 2.5 μl buffer, 0.25 μl 20 μM Wsp18R primer, 0.25 μl 20 μM Wsp18F primer, 0.25 μl 20 μM Wsp691R primer, 0.3 μl 20 μM dNTPs, 1 μl 50 mM MgCl2, 0.125 μl Platinum Taq DNA polymerase (5 U/μl) and 0.5 μl template DNA (2–100 ng μl). Cycle parameters were as follows: 95°C for 5 min, 30 cycles of 60 s, 95°C, 6 s 54°C, 60 s 72°C and a final extension 60 s 95°C, 5 min at 72°C. There was high congruence between our detection of Wolbachia and previous descriptions of DGRP Wolbachia status.

Quantitative and GWA analyses. All data were analysed in R (v3.0.2). The phenotypic traits were analysed with mixed models using the line function from the nlme library, excluding any row of the data matrix with missing data. The interactive effect of microbiota and Wolbachia as binary factors was analysed with genotype nested within experimental block as a random factor. Response indices were calculated for each trait as the difference between mean axenic and gnotobiotic values for the trait in each line, divided by the difference in mean of axenic and gnotobiotic trait value across all lines. To calculate correlations between traits, nutritional traits were normalized to line mean dry weight within each microbiota condition, hence, weight was excluded from these analyses to avoid autocorrelative artefacts. Significance of pairwise Pearson’s correlations were corrected for ten multiple comparisons by Bonferroni correction.

To associate phenotypic traits with genetic variants, GWAs were conducted, using the data for all replicates. R scripts were constructed specifically for this analysis so that all covariates could be fully accounted for. We associated traits with any marker that had a minor allele frequency greater than 0.3 in the pool of tested lines, giving a different subset of markers for each trait tested (number SNPs tested per trait: dry weight 1,881,661 SNPs; protein 1,805,384 SNPs; TAG 1,702,250; dry weight 1,977,756 SNPs; glycogen 1,447,725 SNPs). Models and numbers of DGRP lines used in each GWA are presented in Supplementary Table 8, and degrees of freedom in Supplementary Table 1. QQ plots of test−values from these associations were inflated above the null expectation (Supplementary Fig. 5) and this inflation could not be removed by model respecification or additional covariates (Supplementary Note 1 and Supplementary Fig. 6). Inferences about function and expression of genes bearing significant SNPs were based on Flybase FR0123_06 (ref. 52).

Empirical validation of GWA. The validation experiments used flies with P-element insertion mutations in genes that included the 20 most significant SNPs and the five most significant non-synonymous SNPs in coding regions of genes associated with each trait. To exclude mutants with generally deleterious effects on fitness and condition, we excluded from study genes in which lethal mutations are known and balanced homozygous–lethal mutants. A total of 15 genes were tested in three separate experiments, by exactly the same procedures as in the GWAS, except that dry weights of five individual flies per replicate were determined. The effect of the microbiota on each mutant was analysed, relative to a Wolbachia-status-matched background stock, by a linear mixed model (lme in R version 3.0.2) accounting for genotype (mutant versus background) as a fixed effect and treatment and phenotyped for GWA at 6–8 days post eclosion. The microbiological status of axenic and gnotobiotic flies was checked by plating a sample of fly homogenate (generated as below) onto 1% mMR agar and incubating for 48 h at 30°C under aerobic conditions for Acetobacter and under CO2 for Lactobacillus. As low bacterial counts can be generated by contamination during fly sorting, our validation experiments were conducted on three replicate sets of five male flies in each of these three separate experiments, by exactly the same procedures as in the GWAS, except that dry weights of five individual flies per replicate were determined.

The effect of the microbiota on each mutant was analysed, relative to a Wolbachia-status-matched background stock, by a linear mixed model (lme in R version 3.0.2) accounting for genotype (mutant versus background) as a fixed effect and (n = 3, 3 replicates per experiment) as a random effect. The effect was calculated separately for the mutant/background pair raised under each of axenic or gnotobiotic conditions: a significant effect of genotype (P < 0.025, that is, correcting for two comparisons) under only one of axenic or gnotobiotic conditions indicated the mutant had a different response to the microbiota than the wild-type or background host. The ratio of the gnotobiotic:axenic index value in the mutant, relative to the ratio of the gnotobiotic:axenic index value of the background was calculated to display interactions among host genotype (mutant versus background) and treatment (gnotobiotic or axenic).

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Acknowledgements

We thank David Duneau, Soren Franzenburg, Brenda Gustin, Ginny Howick, Xiangfeng Jing, Susan Rottschaefer, Calum Russell, Robin Schwenke and Rob Unckless for their assistance in experiments. Nicolas Buchon provided advice in the design of mutant studies, and Julien Ayroles, Keegan Kelsey, Jason Mezey, Cornelie Schietz and Rob Unckless provided valuable analytical support. Financial support was provided by NIH grant R01GM095372 and The Sarkaria Institute for Insect Physiology and Toxicology.

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Conceived and designed experiments: A.E.D., J.M.C., A.J.D., B.P.L. and A.G.C. Conducted experiments: A.J.D., J.M.C., P.D.N., S.L.H., S.W., D.R.S., L.D., A.C.-N.W., B.P.L. Analyzed data: A.J.D., J.M.C. and A.E.D.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: The authors declare no competing financial interests.

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How to cite this article: Dobson, A. J. et al. Host genetic determinants of microbiota-dependent nutrition revealed by genome-wide analysis of *Drosophila melanogaster*. *Nat. Commun.* 6:6312 doi: 10.1038/ncomms7312 (2015).
Corrigendum: Host genetic determinants of microbiota-dependent nutrition revealed by genome-wide analysis of Drosophila melanogaster

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Nature Communications 6:6312 doi: 10.1038/ncomms7312 (2015); Published 18 Feb 2015; Updated 15 May 2015

In the top row of Supplementary Fig. 5b in this Article, the graphs for glycogen (left panel) and glucose (right panel) were inadvertently switched. The correct version of the figure appears below.
