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Transcriptomic responses to diet quality and viral infection in Apis mellifera

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Abstract

Background: Parts of Europe and the United States have witnessed dramatic losses in commercially managed honey bees over the past decade to what is considered an unsustainable extent. The large-scale loss of bees has considerable implications for the agricultural economy because bees are one of the leading pollinators of numerous crops. Bee declines have been associated with several interactive factors. Recent studies suggest nutritional and pathogen stress can interactively contribute to bee physiological declines, but the molecular mechanisms underlying interactive effects remain unknown. In this study, we provide insight into this question by using RNA-sequencing to examine how monofloral diets and Israeli acute paralysis virus inoculation influence gene expression patterns in bees.

Results: We found a considerable nutritional response, with almost 2,000 transcripts changing with diet quality. The majority of these genes were over-represented for nutrient signaling (insulin resistance) and immune response (Notch signaling and JaK-STAT pathways). In our experimental conditions, the transcriptomic response to viral infection was fairly limited. We only found 43 transcripts to be differentially expressed, some with known immune functions (argonaute-2), transcriptional regulation, and muscle contraction. We created contrasts to explore whether protective mechanisms of good diet were due to direct effects on immune function (resistance) or indirect effects on energy availability (tolerance). A similar number of resistance and tolerance candidate differentially expressed genes were found, suggesting both processes may play significant roles in dietary buffering from pathogen infection.

Conclusions: Through transcriptional contrasts and functional enrichment analysis, we contribute to our understanding of the mechanisms underlying feedbacks between nutrition and disease in bees. We also show that comparing results derived from combined analyses across multiple RNA-seq studies may allow researchers to identify transcriptomic patterns in bees that are concurrently less artificial and less noisy. This work underlines the merits of using data visualization techniques and multiple datasets to interpret RNA-sequencing studies.

Keywords: Honey bee; RNA-sequencing; Israeli acute paralysis virus; Monofloral pollen; Visualization

1 Background

- 2 Commercially managed honey bees have undergone unusually large declines in the
- united States and parts of Europe over the past decade [1, 2, 3], with annual

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mortality rates exceeding what beekeepers consider sustainable [4, 5]. More than 70 percent of major global food crops (including fruits, vegetables, and nuts) at least benefit from pollination, and yearly insect pollination services are valued worldwide at \$175 billion [6]. As honey bees are largely considered to be the leading pollinator of numerous crops, their marked loss has considerable implications for agricultural sustainability [7].

Honey bee declines have been associated with several factors, including pesticide use, parasites, pathogens, habitat loss, and poor nutrition [8, 9]. Researchers generally agree that these stressors do not act in isolation; instead, they appear to influence the large-scale loss of honey bees in an interactive fashion as the environment changes [10]. Nutrition and viral infection are two broad factors that pose heightened dangers to honey bee health in response to recent environmental changes.

Pollen is a main source of nutrition (including proteins, amino acids, lipids, sterols, starch, vitamins, and minerals) in honey bees [11, 12]. At the individual level, pollen supplies most of the nutrients necessary for physiological development [13] and is believed to have considerable impact on longevity [14]. At the colony level, pollen 20 enables young workers to produce jelly, which then nourishes larvae, drones, older workers, and the queen [15, 16]. Various environmental changes (including urbanization and monoculture crop production) have significantly altered the nutritional profile available to honey bees. In particular, honey bees are confronted with a 24 less diverse selection of pollen, which is of concern because mixed-pollen (polyfloral) diets are generally considered healthier than single-pollen (monofloral) diets 26 [17, 18, 19]. Indeed, reported colony mortality rates are higher in developed land areas compared to undeveloped land areas [20], and beekeepers rank poor nutrition as one of the main reasons for colony losses [21]. Understanding how low diversity Rutter et al. Page 3 of 35

diets affect honey bee health will be crucial to resolve problems that may arise as agriculture continues to intensify throughout the world [22, 23].

Viral infection was a comparatively minor problem in honey bees until the last
century when the ectoparasitic varroa mite (*Varroa destructor*) spread worldwide
[24]. This mite feeds on honey bee hemolymph [25], transmits multiple viruses,
and supports replication of some viruses [26, 27, 28, 29]. More than 20 honey bee
viruses have been identified [30]. One of these viruses that has been linked to honey
bee decline is Israeli acute paralysis virus (IAPV), a positive-sense RNA virus of
the family Dicistroviridae [31]. IAPV infection causes shivering wings, decreased
locomotion, muscle spasms, paralysis, and high premature death percentages in
caged infected adult honey bees [32]. IAPV has demonstrated higher infectious
capacities than other honey bee viruses under certain conditions [33] and is more

Although there is growing interest in how viruses and diet quality affect the health and sustainability of honey bees, as well as a recognition that such factors might operate interactively, there are only a small number of experimental studies thus far directed toward elucidating the interactive effects of these two factors in honey bees [35, 36, 37, 38, 39]. We recently used laboratory cages and nucleus hive experiments to investigate the health effects of these two factors, and our results show the importance of the combined effects of both diet quality and virus infection. Specifically, ingestion by honey bees of high quality pollen is able to mitigate virus-induced mortality to the level of diverse, polyfloral pollen [40].

Following up on these findings, we now aim to understand the corresponding underlying mechanisms by which high quality diets protect bees from virus-induced mortality. For example, it is not known whether the protective effect of good diet is due to direct, specific effects on immune function (resistance), or if it is due to indirect effects of good nutrition on vigor (tolerance) [41]. Transcriptomics is

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one means to better understand the mechanistic underpinnings of dietary and viral
effects on honey bee health. Transcriptomic analysis can help us identify 1) the
genomic scale of transcriptomic response to diet and virus infection, 2) whether
these factors interact in an additive or synergistic way on transcriptome function,
and 3) the types of pathways affected by diet quality and viral infection. This
information, heretofore lacking in the literature, can help us better understand how
good nutrition may be able to serve as a "buffer" against other stressors [42].

There are only a small number of published experiments examining gene expression patterns related to diet effects [43] and virus infection effects [44] in honey bees, but there have also been several such studies in model organisms. For example, diet effects on transcriptomics have been studied in the insect model Drosophila. One recent transcriptomic study in *Drosophila melanogaster* reported an overexpression of genes related to immunity, metabolism, and hemocyanin in a high-fat diet and overexpression of genes related to cell cycle activity, DNA binding and transcription, and CHK kinase-like protein activity in a high-sugar diet [45]. This same study also discovered an upregulation of genes related to peptide and carbohydrate processing in both high-fat and high-sugar diets, a finding the authors attributed to a general increase in caloric intake. Another recent study investigated the transcriptomic effects of diets high in protein relative to sugar, diets high in sugar relative to protein, and diets with equal amounts of protein and sugar [46]. Drosophila mojavensis and Drosophila arizonae showed substantial differential expression between the dietary conditions: genes involved in carbohydrate and lipid metabolism were upregulated 78 in response to high sugar low protein diets and genes involved in juvenile hormone (JH) and ecdysone were upregulated in reponse to low sugar high protein diets. 80 Interestingly, prior studies have suggested that JH regulates body size by controlling ecdysone production, which modifies insulin signaling [47]. While similar transcriptomic studies have been limited in honey bees, one study found that pollen Rutter et al. Page 5 of 35

nutrition upregulates genes involved in macromolecule metabolism, longevity, and the insulin/TOR pathway required for physiological development [43].

There have been numerous studies on the transcriptomic effects of virus infection 86 in model organisms like fruit flies and mosquitoes that can provide a useful framework for interpreting virus responses in honey bees. These studies have shown that RNA silencing is a major antiviral strategy, but other pathways play substantial roles in many virus-host systems like the Toll pathway, IMD pathway, JAK/STAT pathway, Toll-7-autophagy pathway, and transcriptional pausing [48, 49]. It is important to note that general immune responses to viral infection in insects might be an indirect result of cellular damage [49]. In fact, every virus-host interaction has its own particularities derived from the diverse methods of replication and infection cycle evolved by different viruses. An intricate set of pro- and anti-virus host factors such as ribosomal proteins and autophagy pathways are involved, but the response depends on the virus species, as has been elucidated in Drosophila [48, 49]. Model insect studies can inform studies of honey bee virus response, using functional inference of as-of-yet uncharacterized honey bee genes based on orthology to Drosophila and other model organisms. Recent transcriptomic studies in honey 100 bees have improved our understanding of putative molecular mechanisms of honey 101 bee-virus interactions. 102

Key factors or hallmarks of the antiviral conserved defense pathways mentioned above have been found in some virus-bee systems (e.g. RNA silencing, Toll, IMD, JAK/STAT pathway, autophagy, and endocytosis) [50]. In addition, a non-sequence-specific antiviral response mediated by unspecific dsRNA pathway was discovered in honey bees [51, 52]. As in Drosophila, particular antiviral factors in honey bees are linked to specific viruses, specific developmental stages, the analyzed tissue, the route of inoculation, and the time (post-inoculation) during which the study was performed. This was clearly demonstrated when comparing honey bee responses to

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two related iflaviruses with very different infection dynamics, sacbrood bee virus (SBV) vs. deformed wing virus (DWV) [53]. Authors observed differences in in-112 duction of defensin and hymenoptaecin immune-related genes, and suggested the 113 results reflect adaptations to the different routes of transmission [53]. In the case of 114 dicistroviruses, few works have studied the impact of IAPV infection at transcriptional level. Chen et al. 2014 analyzed responses to IAPV infection in larvae and 116 workers using microarrays [54]. Many of the DEGs found were involved in immune 117 response and energy-related metabolism, particularly in adults but not in brood. 118 The authors propose this observed difference could be connected to latent infections 119 in larvae (where host immunity is not perturbed) versus acute infections in adult-120 hood (induced by stressors faced during development) [54]. IAPV acute infection 121 also alters the DNA methylation pattern of numerous genes that do not overlap the 122 genes that are up- or down-regulated at the transcriptional level [44]. These works reiterate the conclusion that viruses trigger particular antiviral mechanisms by dif-124 ferent means and depending on several factors. The honey bee antiviral pathways induced by specific viruses were recently reviewed [50]; it is noteworthy that many honey bee factors discovered by transcriptomics need further characterization to 127 uncover their role in controlling (or promoting) viral infection in honey bees.

As far as we know, there are few to no studies investigating honey bee gene expres-129 sion patterns specifically related to monofloral diets, and few studies investigating 130 honey bee gene expression patterns related to the combined effects of diet in any 131 broad sense and viral inoculation in any broad sense [38]. In this study, we examine 132 how monofloral diets and viral inoculation influence gene expression patterns in 133 honey bees by focusing on four treatment groups (low quality diet without IAPV 134 exposure, high quality diet without IAPV exposure, low quality diet with IAPV 135 exposure, and high quality diet with IAPV exposure). For our diet factor, we exam-136 ined two monofloral pollen diets, rockrose (Cistus sp.) and chestnut (Castanea sp.). Rutter et al. Page 7 of 35

Rockrose pollen is generally considered less nutritious than chestnut pollen because it contains smaller amounts of protein, amino acids, antioxidants, calcium, and iron [40, 55]. We conduct RNA-sequencing analysis on a randomly selected subset of the honey bees we used in our previous study (as is further described in our methods section). We then examine pairwise combinations of treatment groups, the main effect of monofloral diet, the main effect of IAPV exposure, and the combined effect of the two factors on gene expression patterns.

We also compare the main effect of IAPV exposure in our dataset to that obtained 145 in a previous study conducted by Galbraith and colleagues [44]. While our study 146 examines honey bees from naturally mated colonies, the Galbraith study examined 147 honey bees from single-drone colonies. As a consequence, the honey bees in our 148 study will be on average 25% genetically identical, whereas honey bees from the Galbraith study will be on average 75% genetically identical [56]. We note that 150 the difference between these studies may be even greater than this as we used naturally mated honey bees from 15 different colonies. We should therefore expect that the Galbraith study may generate data with higher signal:to:noise ratios than 153 our data due to lower genetic variation between its replicates. At the same time, our honey bees will be more likely to display the health benefits gained from increased 155 genotypic variance within colonies, including decreased parasitic load [57], increased 156 tolerance to environmental changes [58], and increased colony performance [59, 60]. 157 Given that honey bees are naturally very polyandrous [61], our naturally mated 158 honey bees may also reflect more realistic environmental and genetic simulations. 159 Taken together, each study provides a different point of value: Our study likely 160 presents less artificial data while the Galbraith data likely presents less messy data. 161 We wish to explore how the gene expression effects of IAPV inoculation compare 162 between these two studies that used such different experimental designs. To achieve 163 this objective, we use visualization techniques to assess the signal:to:noise ratio Rutter et al. Page 8 of 35

between these two datasets, and differential gene expression (DEG) analyses to
determine any significantly overlapping genes of interest between these two datasets.

As RNA-sequencing data can be biased [62, 63, 64], this comparison allowed us
to characterize how repeatable and robust our RNA-sequencing results were in
comparison to previous studies. It also allowed us to shine light on how experimental
designs that control genetic variability to different extents might affect the resulting
gene expression data in honey bees. We suggest that in-depth data visualization
approaches can be useful for cross-study comparisons and validation of noisy RNAsequencing data in the future.

Results

175 Mortality and virus titers

We reanalyzed our previously published dataset with a subset that focuses on diet 176 quality and is more relevant to the current study. We show the data subset here to 177 inform the RNA-sequencing comparison because we reduced the number of treat-178 ments from the original published data (from eight to four) [40] as a means to focus 179 on diet quality effects. 180 As shown in Figure 1, mortality rates of honey bees 72 hours post-inoculation 181 significantly differed among the treatment groups (mixed model ANOVA across all 182 treatment groups, df = 3, 54; F = 10.03; p < 2.34e-05). The effect of virus treatment 183 (mixed model ANOVA, df = 1, 54; F = 24.73; p < 7.04e-06) and diet treatment 184 (mixed model ANOVA, df = 1, 54; F = 5.32; p < 2.49e-02) were significant, but 185 the interaction between the two factors (mixed model ANOVA, df = 1, 54; F =186 4.72e-02, p = 8.29e-01) was not significant. We compared mortality levels based 187 on pairwise comparisons: For a given diet, honey bees exposed to the virus showed 188 significantly higher mortality rate than honey bees not exposed to the virus. Bees fed 189 rockrose pollen had significantly elevated mortality with virus infection compared 190

to uninfected controls (Benjamini-Hochberg, p < 1.53e-03), and bees fed chestnut

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pollen similarly had significantly elevated mortality with virus infection compared to controls (Benjamini-Hochberg, p < 3.12e-03) (Figure 1).

As shown in Figure 2, IAPV titers of honey bees 72 hours post-inoculation significantly differed among the treatment groups (mixed model ANOVA across all treatment groups, df = 3, 33; F = 6.10; p < 2.03e-03). The effect of virus treatment (mixed model ANOVA, df = 1, 33; F = 15.04; p < 4.75e-04) was significant, but the 197 diet treatment (mixed model ANOVA, df = 1, 33; F = 2.55; p = 1.20e-01) and the 198 interaction between the two factors (mixed model ANOVA, df = 1, 33; F = 7.02e199 01, p = 4.08e-01) were not significant. We compared IAPV titers based on pairwise 200 comparisons: Bees fed rockrose pollen had significantly elevated IAPV titers with 201 virus infection compared to uninfected controls (Benjamini Hochberg, p < 7.56e-202 03). However, bees fed chestnut pollen did not have significantly elevated IAPV 203 titers with virus infection compared to uninfected controls (Benjamini Hochberg, p 204 = 6.29e-02). Overall, we interpreted these findings to mean that high-quality chest-205 nut pollen could partially "rescue" high virus titers resulting from the inoculation 206 treatment, whereas low-quality rockrose pollen could not (Figure 2).

208 Transcriptomic responses to virus infection and diet

We observed a substantially larger number of differentially expressed genes (DEGs) 209 in our diet main effect (n = 1,914) than in our virus main effect (n = 43) (Sup-210 plementary table 1 A and B, Additional file 1). In the diet factor, more DEGs 211 were upregulated in the more-nutritious chestnut group (n = 1,033) than in the 212 less-nutritious rockrose group (n = 881). In the virus factor, there were more virus-213 upregulated DEGs (n = 38) than control-upregulated DEGs (n = 5). While these 214 reported DEG counts are from the DESeq2 package, we saw similar trends for the 215 edgeR and limma package results (Supplementary table 1, Additional file 1 and 216 Additional file 18).

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GO analysis of the chestnut-upregulated DEGs revealed the following over-218 represented biological functions: Wnt signaling, hippo signaling, and dorso-ventral axis formation, as well as pathways related to circadian rhythm, mRNA surveillance, 220 insulin resistance, inositol phosphate metabolism, FoxO signaling, ECM-receptor in-221 teraction, phototransduction, Notch signaling, JaK-STAT signaling, MAPK signal-222 ing, and carbon metabolism (Supplementary table 2, Additional file 1). GO analysis 223 of the rockrose DEGs revealed pathways related to terpenoid backbone biosynthesis, 224 homologous recombination, SNARE interactions in vesicular transport, aminoacyl-225 tRNA biosynthesis, Fanconi anemia, and pyrimidine metabolism (Supplementary 226 table 3, Additional file 1). 227

With so few DEGs (n=43) in our virus main effect comparison, we focused on individual genes and their known functionalities rather than GO over-representation (Table 1). Of the 43 virus-related DEGs, only 10 had GO assignments within the DAVID database. These genes had putative roles in the recognition of pathogenrelated lipid products and the cleaving of transcripts from viruses, as well as involvement in ubiquitin and proteosome pathways, transcription pathways, apoptotic pathways, oxidoreductase processes, and several more functions (Table 1).

No interaction DEGs were observed between the diet and virus factors of the study, in any of the pipelines (DESeq2, edgeR, and limma).

The number of DEGs across the six treatment pairings between the diet and virus factor ranged from 0 to 955 (Supplementary table 8, Additional file 1). Again, diet level appeared to have greater influence on the number of DEGs than the virus level. Across every pair comparing the chestnut and rockrose levels, regardless of the virus level, the number of chestnut-upregulated DEGs was higher than the number of rockrose-upregulated DEGs (Supplementary table 8 C, D, E, F, Additional file 1). Virus-treated bees showed equal to or more upregulated genes relative to controls,

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under both diet treatments (Supplementary table 8 A and B, Additional file 1).

These trends were observed for all three pipelines used (DESeq2, edgeR, and limma).

Transcriptomic data visualization and comparison to a previous study

We wished to explore the signal:to:noise ratio between the Galbraith dataset and 247 our dataset. Note that the Galbraith dataset contained three samples for each 248 virus level, while our dataset contained twelve samples for each virus level. Ba-249 sic PCA plots were constructed with the DESeq2 analysis pipeline and showed 250 that the Galbraith dataset may separate the infected and uninfected honey bees 251 better than our dataset (Additional file 2). We also noted that the first replicate 252 of both treatment groups in the Galbraith data did not cluster as cleanly in the PCA plots. However, through this automatically-generated plot, we can only visualize information at the sample level. Wanting to learn more about the data at the gene level, we continued with new visualization techniques that are available online (https://lrutter.github.io/bigPint) and are in preparation for publication. 257

We used parallel coordinate lines superimposed onto side-by-side boxplots to visu-258 alize the DEGs associated with virus infection in the two studies. The background 250 side-by-side boxplot represents the distribution of all genes in the data, and each 260 parallel coordinate line represents one DEG. In a parallel coordinate line, connec-261 tions between samples with positive correlations should be flat, while connections 262 between samples with negative correlations should be crossed. We expect DEGs 263 to show more variability between treatments than between replicates. This means 264 the parallel coordinate lines should be flat between replicates but crossed between 265 treatments. However, overplotting problems would obscure our visualization if we 266 were to plot all DEGs onto the same side-by-side boxplot. As a result, we used 267 hierarchical clustering techniques to separate DEGs into common patterns as is described in the methods section.

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We see that the 1,019 DEGs from the Galbraith dataset form relatively cleanlooking visual displays, with consistent replicates and differences between treat-271 ments (Figure 3). We do see that the first replicate of the virus group (V.1) appears 272 somewhat inconsistent with the other virus replicates in Cluster 1, confirming that 273 the trend we saw in the PCA plot carried through into the DEG results. Cluster 274 reveals somewhat inconsistent replicates in the virus group, although most virus 275 standardized read counts (group V) remain consistently larger than most control 276 standardized read counts (group N). In contrast, we see that the 43 virus-related 277 DEGs from our dataset do not look as clean in their visual displays (Figure 4). The 278 replicates appear somewhat inconsistent in their estimated expression levels and 279 there is not always such a large (or even consistent) difference between treatment 280 groups. We see a similar finding when we also examine a larger subset of 1,914 281 diet-related DEGs from our study (Additional file 3).

We next used repLicate TREatment ("litre") plots, which we recently developed 283 and published in our bigPint software package. Litre plots allow users to visualize one DEG onto the Cartesian coordinates of one scatterplot matrix. In the litre plot, 285 each gene in the data is plotted once for every combination of replicates between treatment groups. For example, there are nine ways to pair a replicate from one 287 treatment group with a replicate from the other treatment group in the Galbraith 288 dataset (N.1 and V.1, N.1 and V.2, N.1 and V.3, N.2 and V.1, N.2. and V.2, N.2 280 and V.3, N.3 and V.1, N.3 and V.2, and N.3 and V.3). Hence, each gene in the 290 Galbraith dataset is plotted as nine points in the litre plot. With 11.825 genes in 291 the Galbriath data, 106,425 points would need to be plotted. Our dataset is even 292 more dramatic: There are 144 ways to pair a replicate from one treatment group 293 with a replicate from the other treatment group, and with 15,314 genes in our data, 294 we would need to plot 2,205,216 points. For either dataset, plotting all these points would reduce the speed of interactive functionality and cause overplotting problems.

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As a result, we use hexagon bins to summarize this massive information. Once the background of hexagons has been drawn to reveal the distribution of all between-treatment sample pair combinations for *all* genes, the user can superimpose all between-treatment sample pair combinations for one gene of interest.

Additional file 4 shows nine example litre plots for our dataset. The hexagon back-301 ground is the same for all nine litre plots because it simply shows the distribution of all between-treatment sample pair combinations for all genes in our dataset. In each litre plot, there are 144 magenta points superimposed that show all betweentreatment sample pair combinations for one DEG of interest. Additional file 5 and 6 similarly each show nine example litre plots for the Galbraith datset. We examined individual DEGs from the first cluster (Additional file 5) and second cluster (Additional file 6) of the Galbriath data because the first cluster had previously shown 308 less consistency in the first replicate of the treatment group (Figure 3). Notice that, 309 as previously explained, we now show each DEG as nine points for the Galbraith 310 dataset. We see that indeed the virus DEGs from our data (Additional file 4) show 311 less consistent replications and less differences between the treatment groups com-312 pared to the virus DEGs from the Galbraith data (Additional files 5 and 6). We also 313 observe that, in the Galbraith dataset, the DEG points in the first cluster show less 314 tight cluster patterns than the DEG points in the second cluster (Additional files 315 5 and 6), an observation we saw previously in the parallel coordinate plots (Figure 3). 317

Finally, we used scatterplot matrices from the bigPint software to further assess
the DEGs. A scatterplot matrix is another effective multivariate visualization tool
that plots read count distributions across all genes and samples. Specifically, it represents every gene in the dataset as a black point in each scatterplot. DEGs can
be superimposed as colored points to assess their patterns against the full dataset.
We expect DEGs to mostly fall along the x=y line in replicate scatterplots (denot-

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ing replicate consistency) but deviate from the x=y line in treatment scatterplots (denoting significant treatment changes). The x=y line is shown in red in our plots. We created standardized scatterplot matrices for each of the four clusters (from 326 Figure 3) of the Galbraith data (Additional files 7, 8, 9, and 10). We also created 327 standardized scatterplot matrices for our data. However, as our dataset contained 328 24 samples, we would need to include 276 scatterplots in our matrix, which would 329 be too numerous to allow for efficient visual assessment of the data. As a result, 330 we created four scatterplot matrices of our data, each with subsets of 6 samples 331 to be more comparable to the Galbraith data (Additional files 11, 12, 13, and 14). We can again confirm through these plots that the DEGs from the Galbraith data 333 appeared more as expected: They deviated more from the x=y line in the treatment scatterplots while staying close to the x=y line in replicate scatterplots. 335 Despite the virus-related DEGs (n = 1,019) from the Galbraith dataset displaying 336 the expected patterns more than those from our dataset (n = 43), there was significant overlap (p-value < 2.2e-16) in the DEGs between the two studies, with 26/38338 (68%) of virus-upregulated DEGs from our study also showing virus-upregulated response in the Galbraith study (Figure 6).

Tolerance versus resistance

Using the contrasts specified in Table 2, we discovered 122 "tolerance" candi-342 date DEGs and 125 "resistance" candidate DEGs. Within our 122 "tolerance" 343 gene ontologies, we found functions related to metabolism (such as carbohydrate 344 metabolism, fructose metabolism, and chitin metabolism). However, we also discov-345 ered gene ontologies related to RNA polymerase II transcription, immune response, 346 and regulation of response to reactive oxygen species (Figure 5A). Within our 125 347 "resistance" gene ontologies, we found functions related to metabolism (such as car-348 bohydrate metabolism, chitin metabolism, oligosaccharide biosynthesis, and general metabolism) (Figure 5B).

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To visually explore gene expression patterns related to tolerance and resistance, we used hierarchical clustering to separate candidate DEGs into common patterns, and then visualized these clusters using parallel coordinate lines superimposed onto side-by-side boxplots. To reduce overplotting of parallel coordinate lines, we again used hierarchical clustering techniques to separate DEGs into common patterns. Perhaps unsurprisingly, we still see a substantial amount of noise (inconsistency 356 between replicates) in our resulting candidate DEGs (Additional files 15 and 16). 357 However, the broad patterns we expect to see still emerge: For example, based on 358 the contrasts we created to obtain the 'tolerance' candidate DEGs, we expect them 359 to display larger count values in the "NC" group compared to the "NR" group and 360 larger count values in the "VC" group compared to the "VR" group. Indeed, we see 361 this pattern in the associated parallel coordinate plots (Additional file 15). Likewise, 362 based on the contrasts we created to obtain the 'resistance' candidate DEGs, we still expect them to display larger count values in the "VC" group compared to the "VR" group, but we no longer expect to see a difference between the "NC" and "NR" groups. We do generally see these expected patterns in the associated parallel coordinate plots: While there are large outliers in the "NC" group, the "NR" replicates are no longer typically below a standardized count of zero (Additional file 16). The genes in Cluster 3 may follow the expected pattern the most distinctively 369 (Additional file 16). 370

Post hoc analysis

To better understand sources of transcriptomic noise, we explored whether pathogen response measurements (virus titers and mortality), which varied widely across samples, were correlated with observed patterns in gene expression.

The R-squared values between gene read counts and pathogen response measurements were generally low (R-squared < 0.1) across our dataset (Supplementary
table 9, Additional file 1). We further explored whether clusters of DEGs showed

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higher correlations with pathogen response measurements than non-DEGs (the latter serving as a control, where we do not expect a correlation). A Kruskal–Wallis test was used to determine if R-squared populations of DEG clusters significantly differed from those in the rest of the data. The p-values and Bonferroni correction values for each of the 36 tests (as described in the methods section) is provided in Supplementary table 9, Additional file 1. An overall trend emerges to suggest that DEGs may have significantly larger correlation with the pathogen response measurements compared to non-DEGs.

Discussion

Challenges to honey bee health are a growing concern, in particular the combined, 387 interactive effects of nutritional stress and pathogens [42]. In this study, we used RNA-sequencing to probe mechanisms underlying honey bee responses to two ef-389 fects, diet quality and infection with the prominent virus of concern, IAPV. In 390 general, we found a major nutritional transcriptomic response, with nearly 2,000 391 transcripts changing in response to diet quality (rockrose/poor diet versus chest-392 nut/good diet). The majority of these genes were upregulated in response to high 393 quality diet, and these genes were over-represented for functions (Supplementary table 2, Additional file 1) such as nutrient signaling metabolism (insulin resistance) 395 and immune response (Notch signaling and JaK-STAT pathways). These data suggest high quality nutrition may allow bees to alter their metabolism, favoring investment of energy into innate immune responses. While some insect systems have shown relatively low transcriptional responses 399 to dicistrovirus infection [65, 66], previous work on honey bees has revealed many 400 hundreds of DEGs [44]. Discrepancies between datasets may be due to noise and 401 complexity of the honey bee microbiome. The transcriptomic response to virus 402 infection in our experiment was fairly limited. We found only 43 transcripts to be differentially expressed, some with known immune functions (Table 1) such as Rutter et al. Page 17 of 35

argonaute-2 and a gene with similarity to MD-2 lipid recognition protein, as well as genes related to transcriptional regulation and muscle contraction. The small number of DEGs in this study may be partly explained by the large amount of noise in the data (Figure 4 and Additional files 2B, 4, 11, 12, 13, and 14).

Given the noisy nature of our data, and our desire to hone in on genes with real 409 expression differences, we compared our data to the Galbraith study [44], which 410 also examined bees response to IAPV infection. In contrast to our study, Galbraith et al. identified a large number of virus responsive transcripts, and generally had less noise in their data (Figure 3 and Additional files 2A, 5, 6, 7, 8, 9, and 10). To identify the most consistent virus-responsive genes from our study, we looked for 414 overlap in the DEGs associated with virus infection on both experiments. We found 415 a large, statistically significant (p-value < 2.2e-16) overlap, with 26/38 (68%) of 416 virus-responsive DEGs from our study also showing response to virus infection in 417 Galbraith et al. (Figure 6). This result gives us confidence that, although noisy, we 418 were able to uncover reliable, replicable gene expression responses to virus infection 419 with our data. 420

Data visualization is a useful method to identify noise and robustness in RNA-421 sequencing data [67]. In this study, we used extensive data visualization to improve 422 the interpretation of our RNA-sequencing results. For example, the DESeq2 pack-423 age comes with certain visualization options that are popular in RNA-sequencing 424 analysis. One of these visualization is the principal component analysis (PCA) plot, 425 which allows users to visualize the similarity between samples within a dataset. We 426 could determine from this plot that indeed the Galbraith data may show more simi-427 larity between its replicates and differences between its treatments compared to our 428 data (Additional file 2). However, the PCA plot only shows us information at the 429 sample level. We wanted to investigate how these differences in the signal:to:noise ratios of the datasets would affect the structure of any resulting DEGs. As a result, Rutter et al. Page 18 of 35

we also used three plotting techniques from the bigPint package: We investigated the 1,019 virus-related DEGs from the Galbraith dataset and the 43 virus-related 433 DEGs from our dataset using parallel coordinate lines, scatterplot matrices, and litre plots. To prevent overplotting issues in our graphics, we used a hierarchical 435 clustering technique for the parallel coordinate lines to separate the set of DEGs into smaller groups. We also needed to examine four subsets of samples from our 437 dataset to make effective use of the scatterplot matrices. After these tailorizations, 438 we determined that the same patterns we saw in the PCA plots regarding the entire 439 dataset extended down the pipeline analysis into the DEG calls: Even the DEGs 440 from the Galbraith dataset showed more similarity between their replicates and dif-441 ferences between their treatments compared to those from our data. However, the 442 365 DEGs from the Galbraith data in Cluster 1 of Figure 3 showed an inconsistent first replicate in the treatment group ("V.1"), which was something we observed in the PCA plot. This indicates that this feature also extended down the analysis pipeline into DEG calls. Despite the differences in signal between these two datasets, there was substantial overlap in the resulting DEGs. We believe these visualization applications can be useful for future researchers analyzing RNA-sequencing data to quickly and effectively ensure that the DEG calls look reliable or at least overlap with DEG calls from similar studies that look reliable. We also expect this type of 450 visualization exploration can be especially crucial when studying wild populations with high levels of genetic and environmental variation between replicates and/or 452 when using experiments that may lack rigid design control. 453

One of the goals of this study was to use our RNA-sequencing data to assess
whether transcriptomic responses to diet quality and virus infection provide insight
into whether high quality diet can buffer bees from pathogen stress via mechanisms
of "resistance" or "tolerance". Recent evidence has suggested that overall immunity is determined by more than just "resistance" (the reduction of pathogen fitness

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within the host by mechanisms of avoidance and control) [68]. Instead, overall immunity is related to "resistance" in conjunction with "tolerance" (the reduction of adverse effects and disease resulting from pathogens by mechanisms of healing) [41, 68]. Immune-mediated resistance and diet-driven tolerance mechanisms 462 are costly and may compete with each other [41, 69]. Data and models have sug-463 gested that selection can favor an optimum combination of both resistance and 464 tolerance [70, 71, 72, 73]. We attempted to address this topic through specific gene 465 expression contrasts (Table 2), accompanied by GO analysis of the associated gene 466 lists. We found an approximately equal number of resistance (n = 125) and toler-467 ance (n = 122) related candidate DEGs, suggesting both processes may be playing 468 significant roles in dietary buffering from pathogen induced mortality. Resistance 469 candidate DEGs had functions related to several forms of metabolism (chitin and 470 carbohydrate), regulation of transcription, and cell adhesion (Figure 5B). Tolerance candidate DEGs had functions related to carbohydrate metabolism and chitin metabolism; however, they also showed functions related to immune response, including RNA polymerase II transcription (Figure 5A). Previous studies have shown that transcriptional pausing of RNA polymerase II may be an innate immune re-475 sponse in D. melanogaster that allows for a more rapid response by increasing the accessibility of promoter regions of virally induced genes [74]. These possible 477 immunological defense mechanisms within our "tolerance" candidate DEGs and 478 metabolic processes within our "resistance" candidate DEGs may provide addi-479 tional evidence of feedbacks between diet and disease in honey bees [42]. 480

There were several limitations in this study that could be improved upon in future studies. For instance, our comparison between the Galbraith data (single-drone colonies) and our data (naturally mated colonies) was limited by numerous extraneous variables between these studies. In addition to different molecular pipelines and bioinformatic preprocessing pipelines used between these studies, the Galbraith Rutter et al. Page 20 of 35

study focused on one-day old worker honey bees that were fed sugar and artificial pollen diet, whereas our study focused on adult worker honey bees that were fed bee-collected monofloral diets. Furthermore, the Galbraith data used eviscerated abdomens with attached fat bodies and only considered symptomatic honey bees 480 for their infected treatment group, whereas we used whole bodies and considered 490 both asymptomatic and symptomatic honey bees for our infected treatment group. 491 There are also differences in the hours post inoculation and possible differences in 492 the inoculation amount between the studies. Further differences between the stud-493 ies can be found in their corresponding published methods sections [40, 44]. Our 494 comparative visualization assessment between these two datasets was also some-495 what limited because the virus effect in the Galbraith study used three replicates for each level, whereas the virus effect in our study used twelve replicates for each 497 level that were actually further subdivided into six replicates for each diet level. Hence the apparent reduction in noise observed in the Galbraith data compared to our data in the PCA plots, parallel coordinate plots, scatterplot matrices, and litre plots may be an inadvertent product of the smaller number of replicates used and the lack of a secondary treatment group rather than solely the reduction in genetic variability through the single-drone colony design itself. With this in mind, while our current efforts may be a starting point, future studies can shed more light on signal:to:noise and differential expression differences between naturally mated colony designs and single-drone colony designs by controlling for extraneous factors 506 more strictly than what we were able to do in the current line of work. 507

In addition, this study used a whole body RNA-sequencing approach. In future related studies, it may be informative to use tissue-specific methods. Previous work has shown that even though IAPV replication occurs in all honey bee tissues, it localizes more in gut and nerve tissues and in the hypopharyngeal glands. Likewise, the highest IAPV titers have been observed in gut tissues [34]. Recent evidence has Rutter et al. Page 21 of 35

suggested that RNA-sequencing approaches toward composite structures in honey bees leads to false negatives, implying that genes strongly differentially expressed in particular structures may not reach significance within the composite structure 515 [75]. These studies have also found that within a composite extraction, structures 516 therein may contain opposite patterns of differential expression. We can provide 517 more detailed answers to our original transcriptomic questions if we were to repeat 518 this same experimental design only now at a more refined tissue level. Another 519 future direction related to this work would be to integrate multiple omics datasets 520 to investigate monofloral diet quality and IAPV infection in honey bees. Indeed, 521 previous studies in honey bees have found that multiple omics datasets do not 522 always align in a clear-cut manner, and hence may broaden our understanding of 523 the molecular mechanisms being explored [44]. 524

5 Conclusions

To the best of our knowledge, there are few to no studies investigating honey bee gene expression specifically related to monofloral diets, and few to no studies ex-527 amining honey bee gene expression related to the combined effects of diet in any 528 general sense and viral inoculation in any general sense. It also remains unknown 529 whether the protective effects of good diet in honey bees is due to direct effects on 530 immune function (resistance) or indirect effects of energy availability on vigor and 531 health (tolerance). We attempted to address these unresolved areas by conducting 532 a two-factor RNA-sequencing study that examined how monofloral diets and IAPV 533 inoculation influence gene expression patterns in honey bees. Overall, our data sug-534 gest complex transcriptomic responses to multiple stressors in honey bees. Diet has 535 the capacity for large and profound effects on gene expression and may set up the 536 potential for both resistance and tolerance to viral infection, adding to previous evidence of possible feedbacks between diet and disease in honey bees [42].

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Moreover, this study also demonstrated the benefits of using data visualizations and multiple datasets to address inherently messy biological data. For instance, by verifying the substantial overlap in our DEG lists to those obtained in another study that addressed a similar question using specimens with less genetic variability, we were able to place much higher confidence in the differential gene expression results from our otherwise noisy data. We also suggested that comparing results derived 544 from multiple studies varying in level of genetic and environmental variability may 545 allow researchers to identify transcriptomic patterns that are concurrently more 546 realistic and less noisy. Altogether, we hope our results underline the merits of using 547 data visualization techniques and multiple datasets to understand and interpret 548 RNA-sequencing datasets. 549

550 Methods

551 Mortality and virus titers

Details of the procedures we used to prepare virus inoculum, infect and feed caged
honey bees, and quantify IAPV can be reviewed in our previous work [40, 33]. A
linear mixed effects model was used to relate the mortality rates and IAPV titers to
the main and interaction effects of the diet and virus factors. The model was fitted
to the data by restricted maximum likelihood (REML) using the "lme" function
in the R package "nlme". A random (intercept) effect for experimental setup was
included in the model. Post-hoc pairwise comparisons of the four (diet and virus
combination) treatment groups were performed and Benjamini-Hochberg adjusted
p-values were calculated to limit familywise Type I error rates [76].

Design of two-factor experiment

For our nutrition factor, we examined two monofloral pollen diets, rockrose (*Cistus* sp.) and chestnut (*Castanea* sp.). Rockrose pollen is generally considered less nutritious than chestnut pollen due to its lower levels of protein, amino acids, antioxidants, calcium, and iron [40, 55]. For our virus factor, one level contained bees that

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were infected with IAPV and another level contained bees that were not infected with IAPV. This experimental design resulted in four treatment groups (rockrose pollen without IAPV exposure, chestnut pollen without IAPV exposure, rockrose pollen with IAPV exposure, and chestnut pollen with IAPV exposure) that allowed us to assess main effects and interactive effects between diet quality and IAPV 570 infection in honey bees. 571 There are several reasons why our design focused only on diet quality (monofloral 572 diets) as opposed to diet diversity (monofloral diets versus polyfloral diets). First, when assessing diet diversity, a sugar diet is often used as a control. However, such an experimental design does not reflect real-world conditions for honey bees as they rarely face a total lack of pollen [55]. Second, in studies that compared honey bee health using monofloral and polyfloral diets at the same time, if the polyfloral 577 diet and one of the high-quality monofloral diets both exhibited similarly beneficial

effects, then it was difficult for the authors to assess if the polyfloral diet was better

than most of the monofloral diets because of its diversity or because it contained as

a subset the high-quality monofloral diet [55]. Third, as was previously mentioned,

honey bees are now confronted with less diverse sources of pollen. As a result, there

is a need to better understand how monofloral diets affect honey bee health.

RNA extraction

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Fifteen cages per treatment were originally produced for monitoring of mortality.
From these, six live honey bees were randomly selected from each cage 36 hours
post inoculation and placed into tubes [33]. Tubes were kept on dry ice and then
transferred into a -80C freezer until processing. From the fifteen possible cages,
eight were randomly selected for RNA-sequencing. From these eight cages, two of
the honey bees per cage were randomly selected from the original six live honey
bees per cage. These two bees were combined to form a pooled sample representing
the cage. Whole body RNA from each pool was extracted using Qiagen RNeasy

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MiniKit followed by Qiagen DNase treatment. Samples were suspended in water to 200-400 ng/ μ l. All samples were then tested on a Bioanalyzer at the Iowa State University DNA Facility to ensure quality (RIN > 8).

596 Gene expression

Samples were sequenced starting on January 14, 2016 at the Iowa State University DNA Facility (Platform: Illumina HiSeq Sequencing 2500 in rapid run mode; Category: Single End 100 cycle sequencing). A standard Illumina mRNA library was prepared by the DNA facility. Reads were aligned to the BeeBase Version 3.2 genome 600 [77] from the Hymenoptera Genome Database [78] using the programs GMAP and 601 GSNAP [79]. There were four lanes of sequencing with 24 samples per lane. Each 602 sample was run twice. Approximately 75-90% of reads were mapped to the honey 603 bee genome. Each lane produced around 13 million single-end 100 basepair reads. 604 We tested all six pairwise combinations of treatments for DEGs (pairwise DEGs). We also tested the diet main effect (diet DEGs), virus main effect (virus DEGs), and interaction term for DEGs (interaction DEGs). We then also tested for virus main effect DEGs (virus DEGs) in public data derived from a previous study exploring the gene expression of IAPV virus infection in honey bees [44]. We tested each 609 DEG analysis using recommended parameters with DESeq2 [80], edgeR [67], and 610 LimmaVoom [81]. In all cases, we used a false discovery rate (FDR) threshold of 0.05 611 [82]. Fisher's exact test was used to determine significant overlaps between DEG 612 sets (whether from the same dataset but across different analysis pipelines or from 613 different datasets across the same analysis pipelines). The eulerr shiny application 614 was used to construct Venn diagram overlap images [83]. In the end, we focused on 615 the DEG results from DESeq2 [80] as this pipeline was also used in the Galbraith 616 study [44]. We used the independent filtering process built into the DESeq2 software 617 that mitigates multiple comparison corrections on genes with no power rather than 618 defining one filtering threshold.

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Comparison to prior studies on transcriptomic response to viral infection

We compare the main effect of IAPV exposure in our dataset to that obtained in a 621 previous study conducted by Galbraith and colleagues [44] who also addressed honey 622 bee transcriptomic responses to virus infection. We applied the same downstream 623 bioinformatics analyses between our count table and the count table provided in 624 the Galbraith study. When we applied our bioinformatics pipeline to the Galbraith 625 count table, we obtained different differential expression counts compared to the 626 results published in the Galbraith study. However, there was substantial overlap and we considered this justification to use the differential expression list we obtained in order to keep the downstream bioinformatics analyses as similar as possible between 629 the two datasets (Additional file 17). 630

We used honey bees from naturally mated colonies, whereas Galbraith et al. [44]
used honey bees from single-drone colonies. In light of this, we should expect the
Galbraith et al. dataset to contain lower genetic variation between its replicates
and higher signal:to:noise ratios than our dataset. We use visualization techniques
to assess the signal:to:noise ratio between these two datasets, and differential gene
expression (DEG) analyses to determine any significantly overlapping genes of interest between these two datasets.

638 Visualization

We used an array of visualization tools as part of our analysis. We used the PCA plot
[84] from the DESeq2 package, a well-known and established tool. Along with that,
we used lesser-known multivariate visualization tools from our work-in-progress R
package called bigPint. Specifically, we used parallel coordinate plots [85], scatterplot matrices [86], and litre plots (which we recently developed based on "replicate
line plots" [87]) to assess the variability between the replicates and the treatments
in our data. We also used these plotting techniques to assess for normalization
problems and other common problems in RNA-sequencing analysis pipelines [87].

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Furthermore, we used statistical graphics to better understand patterns in our

DEGs. However, in cases of large DEG lists, these visualization tools had overplotting problems (where multiple objects are drawn on top of one another, making
it impossible to detect individual values). To remedy this problem, we first standardized each DEG to have a mean of zero and standard deviation of unity [88, 89].

Then, we performed hierarchical clustering on the standardized DEGs using Ward's
linkage. This process divided large DEG lists into smaller clusters of similar patterns, which allowed us to more efficiently visualize the different types of patterns
within large DEG lists (see Figures 3 and 4 for examples).

656 Gene ontology

DEGs were uploaded as a background list to DAVID Bioinformatics Resources 6.7 [90, 91]. The overrepresented gene ontology (GO) terms of DEGs were determined using the BEEBASE_ID identifier option (honey bee gene model) in the DAVID software. To fine-tune the GO term list, only terms correlating to Biological Processes were considered. The refined GO term list was then imported into REVIGO [92], which uses semantic similarity measures to cluster long lists of GO terms.

Probing tolerance versus resistance

To investigate whether the protective effect of good diet is due to direct, specific effects on immune function (resistance), or if it is due to indirect effects of good nutrition on energy availability and vigor (tolerance), we created contrasts of interest 666 (Table 2). In particular, we assigned "resistance candidate DEGs" to be the ones 667 that were upregulated in the chestnut group within the virus infected bees but not 668 upregulated in the chestnut group within the non-infected bees. Our interpretation 669 of these genes is that they represent those that are only activated in infected bees 670 that are fed a high quality diet. We also assigned "tolerance candidate DEGs" to 671 be the ones that were upregulated in the chestnut group for both the virus infected bees and non-infected bees. Our interpretation of these genes is that they represent

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those that are constitutively activated in bees fed a high quality diet, regardless of whether they are experiencing infection or not. We then determined how many genes fell into these two categories and analyzed their GO terminologies.

677 Post hoc analysis

We found considerable noisiness in our data and saw, through gene-level visualizations, that our DEGs contained outliers and inconsistent replicates. Hence, we wanted to explore whether our DEG read counts correlated with pathogen response 680 metrics, including IAPV titers, sacbrood bee virus (SBV) titers, and mortality rates. 681 For this process, we considered virus main effect DEGs (Figure 4), "tolerance can-682 didate" DEGs (Additional file 15), and "resistance candidate" DEGs (Additional 683 file 16). For each DEG in each cluster, we calculated a coefficient of determination 684 (R-squared) value to estimate the correlation between its raw read counts and the 685 pathogen response metrics across its 24 samples. We then used the Kruskal-Wallis 686 test to determine if the distribution of the R-squared values in any of the DEG clus-687 ters significantly differed from those in the non-DEG genes (the rest of the data). 688 As there were four clusters for each of the nine combinations of DEG lists ("tol-689 erance" candidate DEGs, "resistance" candidate DEGs, and virus-related DEGs) 690 and pathogen response measurements (IAPV titer, SBV titer, and mortality rate), this process resulted in 36 statistical tests.

693 Ethics approval and consent to participate

694 All honey bees used in this work were sampled in the United States, and no ethical use approval is required for this 695 species in this country.

696 Consent for publication

697 Not applicable.

98 Availability of data and materials

 699 The datasets generated and/or analysed during the current study are available in the [NAME] repository,

700 [PERSISTENT WEB LINK TO DATASETS]. Include our data, Galbraith data, scripts to reproduce tables and

701 figures (on GitHub).

702 Competing interests

703 The authors declare that they have no competing interests.

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707 Author's contributions

708 LR performed the bioinformatic and statistical analyses, produced the figures and tables, and drafted the

manuscript. BB conceptualized the study and critically revised the manuscript. AD contributed to experimental

710 design, carried out the laboratory experiments, and processed samples for virus titers and RNA-seq

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916 Figures

Figure 1 Mortality rates for the four treatment groups, two virus groups, and two diet groups. Left to right: Mortality rates for the four treatment groups, two virus groups, and two diet groups. "N" represents non-inoculation, "V" represents viral inoculation, "C" represents chestnut pollen, and "R" represents rockrose pollen. The mortality rate data included 59 samples with 15 replicates per treatment group, except for the "NC" group having 14 replicates. ANOVA values and p-values for the statistical tests are listed in the text of the paper. The letters above the bars represent significant differences with a confidence level of 95%.

Figure 2 IAPV titers for the four treatment groups, two virus groups, and two diet groups. Left to right: IAPV titers for the four treatment groups, two virus groups, and two diet groups. "N" represents non-inoculation, "V" represents viral inoculation, "C" represents chestnut pollen, and "R" represents rockrose pollen. The IAPV titer data included 38 samples with 10 replicates per treatment group, except for the "NR" group having 8 replicates. ANOVA values and p-values for the statistical tests are listed in the text of the paper. The letters above the bars represent significant differences with a confidence level of 95%.

Figure 3 Parallel coordinate plots of the 1,019 DEGs after hierarchical clustering of size four between the virus-infected and control groups of the Galbraith data [44]. Parallel coordinate plots of the 1,019 DEGs after hierarchical clustering of size four between the virus-infected and control groups of the Galbraith study. "N" represents non-inoculation, "V" represents viral inoculation. Clusters 1, 2, and 4 seem to represent DEGs that were overexpressed in the virus inoculated group, and Cluster 3 seems to represent DEGs that were overexpressed in the non-inoculated control group. In general, the DEGs appeared as expected, but there is rather noticeable deviation of the first replicate from the virus-treated sample ("V.1") from the other virus-treated replicates in Cluster 1.

Figure 4 Parallel coordinate plots of the 43 DEGs after hierarchical clustering of size four between the virus-infected and control groups of our study. Parallel coordinate plots of the 43 DEGs after hierarchical clustering of size four between the virus-infected and control groups of our study. "N" represents non-infected control group, and "V" represents treatment of virus. The vertical red line indicates the distinction between treatment groups. We see from this plot that the DEG designations for this dataset do not appear as clean compared to what we saw in the Galbraith dataset in Figure 3.

Figure 5 Gene ontology analysis results for the 122 DEGs related to our "tolerance" hypothesis and for the 125 DEGs related to our "resistance" hypothesis. GO analysis results for the 122 DEGs related to our "tolerance" hypothesis (A) and for the 125 DEGs related to our "resistance" hypothesis (B). The color and size of the circles both represent the number of genes in that ontology. The x-axis and y-axis are organized by SimRel, a semantic similarity metric [93].

Figure 6 Venn diagrams comparing the virus-related DEG overlaps between our dataset and the Galbraith dataset. Venn diagrams comparing the virus-related DEG overlaps between the Galbraith study (labeled as "G") and our study (labeled as "R"). From left to right: Total virus-related DEGs (subplot A), virus-upregulated DEGs (subplot B), control-upregulated DEGs (subplot C). Both the total virus-related and virus-upregulated DEGs showed significant overlap between the studies (p-value < 2.2e-16) as per Fisher's Exact Test for Count Data. There was one gene that was virus-upregulated in the Galbraith study but control-upregulated in our study.

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917 **Tables**

| BeeBase ID | Gene Name | Known functions | Us | Galbraith |
|------------|----------------------------------|--|------------------|-----------|
| GB41545 | MD-2-related | Implicated in lipid recognition, | | - |
| | lipid-recognition | particularly in the recognition of | N | |
| | protein-like | pathogen related products | | |
| | | Interacts with small interfering RNAs | N s V V d n v er | V |
| GB50955 | Protein argonaute-2 | to form RNA-induced silencing | | |
| | | complexes which target and cleave | | |
| | | transcripts that are mostly from | | |
| | | viruses and transposons | | |
| | UBA-like | Found in diverse proteins involved | | V |
| GB48755 | domain-containing | in ubiquitin/proteasome | V | |
| | protein 2 | pathways | | |
| GB47407 | Histone H4 | Capable of affecting transcription, | | V |
| | | DNA repair, and DNA replication | V | |
| | | when post-transcriptionally modified | | |
| GB42313 | Leishmanolysin-like peptidase | Encodes a protein involved in cell | V | V |
| | | migration and invasion; implicated in | | |
| | | mitotic progression in D. melanogaster | | |
| | Rho guanine | Implicated in regulation of apoptopic processes, cell growth, signal transduction, and transcription | | |
| GB50813 | nucleotide | | | V |
| | exchange factor 11 | | | |
| | Thioredoxin | Converse of a general protein | | <u> </u> |
| GB54503 | domain-containing | Serves as a general protein disulphide oxidoreductase | | - |
| | protein | | | |
| GB53500 | Transcriptional | Regulator gene that codes for a | V | V |
| GD33300 | regulator Myc-B | transcription factor | | v |
| GB51305 | Tropomyosin-like | Related to protein involved in muscle | N | N |
| | | contraction | 1 1 | |
| GB50178 | Cilia and | Induces components required for wild-type motility and | | |
| | flagella-associated | | | V |
| | protein 61-like | stable assembly of motile cilia | | |

Table 1 Known functions of the mapped subset of 43 DEGs in the virus main effect of our study. Whether the gene was overrepresented in the virus or non-virus group is also indicated for both our study and the Galbraith study. Functionalities were extracted from Flybase, National Center for Biotechnology Information and The European Bioinformatics Institute databases.

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| DEGs | Interpretation | Results | |
|------|--------------------------------|--|--|
| 43 | Genes that change expression | | |
| | due to virus effect regardless | Table 1 | |
| | of diet status in bees | | |
| | Genes that change expression | Supplementary | |
| 941 | due to diet effect in | tables 4 and 5, | |
| | uninfected bees | Additional file 1 | |
| 376 | Genes that change expression | Supplementary | |
| | due to diet effect in | tables 6 and 7, | |
| | infected bees | Additional file 1 | |
| | "Tolerance" genes that turn | | |
| 122 | on by good diet regardless of | Figure 5A | |
| | virus infection status in bees | | |
| 125 | "Resistance" genes that turn | | |
| | on by good diet only in | Figure 5B | |
| | infected bees | | |
| | 43 941 376 | Genes that change expression due to virus effect regardless of diet status in bees Genes that change expression due to diet effect in uninfected bees Genes that change expression due to diet effect in infected bees "Tolerance" genes that turn on by good diet regardless of virus infection status in bees "Resistance" genes that turn on by good diet only in infected bees | |

Table 2 Contrasts in our study for assessing GO and pathways analysis.

Additional Files

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Additional file 1 — Supplementary tables.

Table 1: Number of DEGs across three analysis pipelines for (A) the diet main effect in our study, (B) the virus main effect in our study, and (C) the virus main effect in the Galbraith study. For the diet effects, "C" represents chestnut diet and "R" represents rockrose diet. For the virus effects, "N" represents control non-inoculated and "V" represents virus-inoculated. Table 2: Pathways related to the 1,033 DEGs that were upregulated in the chestnut treatment from the diet main effect. Table 3: Pathways related to the 881 DEGs that were upregulated in the rockrose treatment from the diet main effect. Table 4: GO analysis results for the 601 DEGs that were upregulated in the NC treatment from the NC versus NR treatment pair analysis. These DEGs represent genes that are upregulated when non-infected honey bees are given high quality chestnut pollen compared to being given low quality rockrose pollen. Table 5: GO analysis results for the 340 DEGs that were upregulated in the NR treatment from the NC versus NR treatment pair analysis. These DEGs represent genes that are upregulated when non-infected honey bees are given low quality rockrose pollen compared to being given high quality chestnut pollen. Table 6: GO analysis results for the 247 DEGs that were upregulated in the VC treatment from the VC versus VR treatment pair analysis. These DEGs represent genes that are upregulated when infected honey bees are given high quality chestnut pollen compared to being given low quality rockrose pollen. Table 7: GO analysis results for the 129 DEGs that were upregulated in the VR treatment from the VC versus VR treatment pair analysis. These DEGs represent genes that are upregulated when infected honey bees are given low quality rockrose pollen compared to being given high quality chestnut pollen. Table 8: Number of DEGs across three analysis pipelines for all six treatment pair combinations between the diet and virus factor. "C" represents chestnut diet, "R" represents rockrose diet, "V" represents virus-inoculated, and "N" represents control non-inoculated. Table 9: Kruskal-Wallis p-value and Bonferroni corrections for the 36 combinations of DEG lists, pathogen response metrics, and cluster number. (XLS).

940 Additional file 2 — PCA plots for the Galbraith dataset and for our dataset.

PCA plots for the Galbraith dataset (A) and for our dataset (B). "V" represents virus-inoculated, and "N"

942 represents control non-inoculated. The x-axis represents the principal component with the most variation and the

y-axis represents the principal component with the second-most variation (PNG).

944 Additional file 3 — Parallel coordinate lines of the diet-related DEGs of our dataset.

Parallel coordinate plots of the 1,914 DEGs after hierarchical clustering of size six between the chestnut and

946 rockrose groups of our study. Here "C" represents chestnut samples, and "R" represents rockrose samples. The

947 vertical red line indicates the distinction between treatment groups. We see from this plot that the DEG designations

948 for this dataset do not appear as clean compared to what we saw in the Galbraith dataset in Figure 3 (PNG).

 $\,$ Additional file 4 — Example litre plots from the virus-related DEGs of our dataset.

950 Example litre plots of the nine DEGs with the lowest FDR values from the 43 virus-related DEGs of our dataset.

"N" represents non-infected control samples and "V" represents virus-treated samples. Most of the magenta points

952 (representing the 144 combinations of samples between treatment groups for a given DEG) do not reflect the

expected pattern as clearly compared to what we saw in the litre plots of the Galbraith data. They are not as

 $_{954}$ clustered together (representing replicate inconsistency) and they sometimes cross the x=y line (representing lack of

955 difference between treatment groups). This finding reflects what we saw in the messy looking parallel coordinate

lines of Figure 4 (PNG).

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Additional file 5 — Example litre plots of DEGs from Cluster 1 of the Galbraith dataset.
      Example litre plots of the nine DEGs with the lowest FDR values from the 365 DEGs in Cluster 1 (originally shown
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      in Figure 3) of the Galbraith dataset. "N" represents non-infected control samples and "V" represents virus-treated
      samples. Most of the light orange points (representing the nine combinations of samples between treatment groups
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      for a given DEG) deviate from the x=y line in a tight bundle as expected (PNG).
      Additional file 6 — Example litre plots of DEGs from Cluster 2 of the Galbraith dataset.
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      Example litre plots of the nine DEGs with the lowest FDR values from the 327 DEGs in Cluster 2 (originally shown
      in Figure 3) of the Galbraith dataset. "N" represents non-infected control samples and "V" represents virus-treated
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      samples. Most of the dark orange points (representing the nine combinations of samples between treatment groups
      for a given DEG) deviate from the x=y line in a compact clump as expected. However, they are not as tightly
      bunched together compared to what we saw in the example litre plots of Cluster 1 (shown in Additional file 5). As a
967
      result, what we see in these litre plots reflects what we saw in the parallel coordinate lines of Figure 3: The replicate
      consistency in the Cluster 1 DEGs is not as clean as that in the Cluster 2 DEGs, but is still relatively clean (PNG).
      Additional file 7 — Scatterplot matrix of DEGs from Cluster 1 of the Galbraith dataset.
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      The 365 DEGs from the first cluster of the Galbraith dataset (originally shown in Figure 3) superimposed as light
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      orange dots onto all genes as black dots in the form of a scatterplot matrix. The data has been standardized. "N"
      represents non-infected control samples and "V" represents virus-treated samples. We confirm that the DEGs
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      mostly follow the expected structure, with their placement deviating from the x=y line in the treatment
      scatterplots, but adhering to the x=y line in the replicate scatterplots. However, we do see that sample "V.1" may
      be somewhat inconsistent in these DEGs, as its presence in the replicate scatterplots shows DEGs deviating from
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      the x=y line more than expected and its presence in the treatment scatterplots shows DEGs adhering to the x=y
      line more than expected. This inconsistent sample was something we observed in Figure 3 (PNG).
      Additional file 8 — Scatterplot matrix of DEGs from Cluster 2 of the Galbraith dataset.
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      The 327 DEGs from the second cluster of the Galbraith dataset (originally shown in Figure 3) superimposed as dark
      orange dots onto all genes as black dots in the form of a scatterplot matrix. The data has been standardized. "N"
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      represents non-infected control samples and "V" represents virus-treated samples. We confirm that the DEGs
      mostly follow the expected structure, with their placement deviating from the x=y line in the treatment
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      scatterplots, but adhering to the x=y line in the replicate scatterplots (PNG).
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      Additional file 9 — Scatterplot matrix of DEGs from Cluster 3 of the Galbraith dataset.
      The 224 DEGs from the third cluster of the Galbraith dataset (originally shown in Figure 3) superimposed as
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      turquoise dots onto all genes as black dots in the form of a scatterplot matrix. The data has been standardized. "N"
      represents non-infected control samples and "V" represents virus-treated samples. We confirm that the DEGs
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      mostly follow the expected structure, with their placement deviating from the x=y line in the treatment
      scatterplots, but adhering to the x=y line in the replicate scatterplots (PNG).
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      Additional file 10 — Scatterplot matrix of DEGs from Cluster 4 of the Galbraith dataset.
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      The 103 DEGs from the fourth cluster of the Galbraith dataset (originally shown in Figure 3) superimposed as pink
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      dots onto all genes as black dots in the form of a scatterplot matrix. The data has been standardized. "N"
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      represents non-infected control samples and "V" represents virus-treated samples. We confirm that the DEGs
      mostly follow the expected structure, with their placement deviating from the x=y line in the treatment
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      scatterplots, but adhering to the x=y line in the replicate scatterplots. We also see that the second replicate from
      the virus-treated sample ("V.2") may be somewhat inconsistent in these DEGs, as its presence in the replicate
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      scatterplots results in the DEGs unexpectedly deviating from the x=y line and its presence in the treatment
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      scatterplots results in the DEGs unexpectedly adhering to the x=y line (PNG).
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      Additional file 11 — Scatterplot matrix of virus-related DEGs from our dataset, showing only replicates 1, 2, and 3.
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      The 43 virus-related DEGs from our dataset superimposed as magenta dots onto all genes in the form of a
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1002
      scatterplot matrix. Only replicates 1, 2, and 3 are shown from both treatment groups. The data has been
      standardized. "N" represents non-infected control samples and "V" represents virus-treated samples. We see that,
      compared to the scatterplot matrices from certain clusters of the Galbraith data, the 43 DEGs from this subset of
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      six samples from our data do not paint as clear of a picture, sometimes unexpectedly deviating from the x=y line in
      the replicate plots and sometimes unexpectedly adhering to the x=y line in the treatment plots (PNG).
      Additional file 12 — Scatterplot matrix of virus-related DEGs from our dataset, showing only replicates 4, 5, and 6.
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      The 43 virus-related DEGs from our dataset superimposed as magenta dots onto all genes in the form of a
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      scatterplot matrix. Only replicates 4, 5, and 6 are shown from both treatment groups. The data has been
      standardized. "N" represents non-infected control samples and "V" represents virus-treated samples. We see that,
      compared to the scatterplot matrices from certain clusters of the Galbraith data, the 43 DEGs from this subset of
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six samples from our data do not paint as clear of a picture, and most of them unexpectedly adhere to the x=y line

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in the treatment plots (PNG).

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Additional file 13 — Scatterplot matrix of virus-related DEGs from our dataset, showing only replicates 7, 8, and 9.

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The 43 virus-related DEGs from our dataset superimposed as magenta dots onto all genes in the form of a
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      scatterplot matrix. Only replicates 7, 8, and 9 are shown from both treatment groups. The data has been
      standardized. "N" represents non-infected control samples and "V" represents virus-treated samples. We see that,
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      compared to the scatterplot matrices from certain clusters of the Galbraith data, the 43 DEGs from this subset of
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      six samples from our data do not paint as clear of a picture, sometimes unexpectedly deviating from the x=y line in
      the replicate plots and sometimes unexpectedly adhering to the x=y line in the treatment plots (PNG).
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      Additional file 14 — Scatterplot matrix of virus-related DEGs from our dataset, showing only replicates 10, 11, and
      12
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      The 43 virus-related DEGs from our dataset superimposed onto all genes in the form of a scatterplot matrix. Only
      replicates 10, 11, and 12 are shown from both treatment groups. The data has been standardized. "N" represents
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      non-infected control samples and "V" represents virus-treated samples. We see that, compared to the scatterplot
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      matrices from certain clusters of the Galbraith data, the 43 DEGs from this subset of six samples from our data do
      not paint as clear of a picture, and most of them unexpectedly deviate from the x=v line in the virus-related
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      replicate plots (PNG).
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      Additional file 15 — Parallel coordinate plots of the "tolerance" candidate DEGs.
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      Parallel coordinate plots of the 122 DEGs after hierarchical clustering of size four between the "tolerance" candidate
      DEGs. Here "N" represents non-infected control group, "V" represents treatment of virus, "C" represents
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      high-quality chestnut diet, and "R" represents low-quality rockrose diet. The vertical red line indicates the
      distinction between treatment groups. We see there is considerable noise in the data (non-consistent replicate
      values), but that the general patterns of the DEGs follow what we expect based on our "tolerance" contrast (PNG).
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      Additional file 16 — Parallel coordinate plots of the "resistance" candidate DEGs.
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      Parallel coordinate plots of the 125 DEGs after hierarchical clustering of size four between the "resistance"
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      candidate DEGs. Here "N" represents non-infected control group, "V" represents treatment of virus, "C" represents
      high-quality chestnut diet, and "R" represents low-quality rockrose diet. The vertical red line indicates the distinction
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      between treatment groups. We see there is considerable noise in the data (non-consistent replicate values), but that
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      the general patterns of the DEGs follow what we expect based on our "resistance" contrasts (PNG).
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      Additional file 17 — Venn diagrams comparing the virus-related DEG overlaps in the Galbraith data using our
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      pipeline and the pipeline used by Galbraith et al.
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      Venn diagrams comparing the virus-related DEG overlaps of the Galbraith data from the DESeq2 bioinformatics
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      pipelines used in the Galbraith study (labeled as "G.O.") and the DESeq2 bioinformatics pipelines used in our study
      (labeled as "G.R"). While we were not able to fully replicate the DEG list published in the Galbraith study, our DEG
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      list maintained significant overlaps with their DEG list. From left to right: Total virus-related DEGs (subplot A),
      virus-upregulated DEGs (subplot B), control-upregulated DEGs (subplot C) (PNG).
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      Additional file 18 — Venn diagrams of main effect DEG overlaps across DESeq2, edgeR, and limma
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      Venn diagrams comparing DEG overlaps across DESeq2, edgeR, and limma for our diet main effect (top row), our
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      virus main effect (middle row), and the Galbraith virus main effect (bottom row). Within a given subplot, "D'
      represents DESeq2, "E" represents edgeR, and "L" represents limma. From left to right on top row: Total
      diet-related DEGs (subplot A), chestnut-upregulated DEGs (subplot B), rockrose-upregulated DEGs (subplot C).
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      From left to right on middle row: Total virus-related DEGs (subplot D), virus-upregulated DEGs (subplot E),
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      control-upregulated DEGs in our data (subplot F). From left to right on bottom row: Total virus-related DEGs
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      (subplot G), virus-upregulated DEGs (subplot H), control-upregulated DEGs in the Galbraith data (subplot I)
      (PNG). With the exception of the limma pipeline resulting in zero DEGs in our virus main effect analysis, we found
      significant overlaps between DEG lists across the different pipelines (DESeq2, edgeR, and limma). In general,
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      DESeq2 resulted in the largest number of DEGs and limma resulted in the least number of DEGs (PNG).
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      Additional file 19 — Analysis of correlation between DEG read counts and pathogen response metrics
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      Distribution of R-squared values for DEG cluster read counts and pathogen response metrics. Columns left to right:
      SBV titers, mortality rates, and IAPV titers. Rows top to bottom: Tolerance candidate DEGs, resistance candidate
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      DEGs, and virus-related DEGs. Each subplot includes five boxplots which represent the R-squared value distributions
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      for four DEG clusters and all remaining non-DEGs in the data. The top number above each boxplot represents the
      number of genes included. The first four boxplots also include a bottom number, which represents the
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1065
      Kruskal-Wallis p-value of the comparison of the R-squared distribution of the cluster and the R-squared distribution
      of the non-DEG data (PNG).
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```