# Title: Differential carbohydrate utilization and organic acid production by honey bee symbionts

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- 5 <sup>1,2</sup>Fredrick J. Lee, <sup>1</sup>Kayla I. Miller, <sup>1</sup>James B. McKinlay, and <sup>1</sup>Irene L. G. Newton\*
- 6 1=Department of Biology, Indiana University, Bloomington, IN
- 7 2=Current address: Department of Biology, Tufts University, Medford, MA
- 8 \*corresponding author
- 9 1001 E 3<sup>rd</sup> St
- 10 Bloomington, IN 47405
- 11 812-855-3883
- 12 irnewton@indiana.edu
- 13

## 14 **Abstract:**

15 The honey bee worker gut is host to a community of bacteria that primarily 16 comprises 8-10 bacterial species. Collectively, these microbes break down and ferment 17 saccharides present in the host's diet. The model of metabolism for these gut symbionts 18 is rooted in previous analyses of genomes, metagenomes, and metatranscriptomes of 19 this environment. Importantly, there is a correlation between the composition of the gut 20 microbiome and weight gain in the honey bee, suggesting that bacterial production of 21 organic acids might contribute to the observed phenomenon. Here we identify potential 22 metabolic contributions of symbionts within the honey bee gut. We show significant 23 variation in the metabolic capabilities of these microbes, highlighting the fact that 24 although the microbiota appears simple and consistent based on 16S rRNA gene 25 profiling, strains are highly variable in their ability to use specific carbohydrates and 26 produce organic acids. Finally, we confirm that the honey bee core microbes, especially 27 a clade of y-proteobacteria (i.e. Gilliamella), are highly active in vivo, expressing key 28 enzymatic genes critical for utilizing plant-derived molecules and producing organic 29 acids. These results suggest that Gilliamella, and other core taxa, may contribute 30 significantly to weight gain in the honey bee, specifically through the production of 31 organic acids.

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#### 33 Introduction:

34 Insects are some of the most abundant and diverse species on earth, inhabiting 35 a plethora of unique environments, ranging from the arid sands of the hottest desert to 36 frigid temperatures of glaciers (1). In order to thrive in such unusual environments, many 37 species of insects have evolved specialized symbioses with microbes capable of 38 synthesizing essential nutrients (e.g., amino acids and vitamins) or degrading complex 39 macromolecules (2). The world's most important agricultural pollinator, the honey bee 40 (Apis mellifera), is no exception, as it relies on microbes to aid in the digestion of its 41 plant-derived diet of honey and bee bread (3). After consumption, food ingested by the 42 bee is metabolized by the microbes that inhabit the honey bee gut, which include a small 43 number of dominant taxa that constitute bacterial species considered to be "core" to the 44 honey bee gut microbiota, including the proteobacteria Snodograssella alvi 45 (Betaproteobacteria), Parasaccharibacter apium (Alphaproteobacteria), Frischella 46 perrara and Gilliamella apicola (Gammaproteobacteria), two Lactobacillus species 47 (Firmicutes), a Bifidobacterium species (Actinobacteria), and a rarer Bacteroidetes (4). 48 Three of these phyla, the Gammaproteobacteria, Firmicutes, and Bifidobacteria, 49 dominate predicted functional capacity of the honey bee gut microbiome, based on a 50 metatranscriptomic survey on the honey bee gut (5). The current proposed model of 51 bacterial community metabolism in the bee gut, based on annotated gene presence and 52 expression, suggests that these core microbes participate in the breakdown and 53 fermentation of host dietary macromolecules into an array of alcohols, gases, short chain 54 fatty acids (SCFAs), and other organic acids, some of which likely serve to sustain the 55 host (5-10). Complementing this work, some studies have demonstrated, in vitro, the 56 ability of some strains to ferment a variety of substrates (5, 9, 11-14) and produce 57 organic acids (i.e. lactate and acetate)(11, 14). In addition to these observations, a 58 separate study also recently demonstrated that gnotobiotic, mono-colonized bees

59 possessed guts containing significant depletion in plant derivatives (i.e. flavonoids,  $\omega$ -60 hydroxy acids, and phenolamides), likely originating from pollen consumed by the host, 61 and an enrichment in fermentative products in the form of organic acids and SCFAs, 62 compared to bees with a depleted microbiota (15). The production of these fatty acids 63 may contribute to honey bee health. Indeed, a recent study identified a correlation 64 between host weight gain and organic acids in the digestive tract of bees (3). Bees 65 supplemented with gut bacteria showed an increase in the variety and amount of organic 66 acids present within the host digestive system, as well as a significant increase in host 67 weight gain and the expression of host genes related to insect growth and nutrient 68 homeostasis, compared to controls. Collectively, these results link the honey bee 69 symbionts to the production of organic acids and then to host growth and nutrient 70 acquisition. It has been proposed that honey bee core bacteria likely convert 71 carbohydrates in the bee diet to a variety of organic acids, which can either act as a 72 signaling molecules eliciting a host response, or act as building blocks for host derived 73 anabolism.

74 In light of these recent findings, we sought to determine the metabolic 75 contribution of the Gammaproteobacteria, Firmicutes, and Bifidobacteria, within the bee 76 gut, as these phyla dominate the metatranscriptomic profile of the digestive tract of the 77 honey bee. Although there are well-defined species within these groupings, there is still 78 a large diversity of honey bee strains found within individual bees and colonies (16). The 79 strain diversity within each of the honey bee core clades is reflected in differences in 80 genomic content (7, 16), suggesting that significant functional diversity may exist within 81 each of the core 8-10 clades associated with the honey bee. In this study, we address 82 fundamental questions regarding metabolic characteristics of the honey bee gut 83 microbiota, such as: 1) What is the pattern of anaerobic utilization of environmentally 84 relevant carbon sources across dominant symbionts, and what are the direct byproducts

85 produced, per taxa, in vitro? 2) Are there functional differences in the patterns of 86 utilization between isolates of the same species? 3) What contribution does an individual 87 taxon make to the utilization and the subsequent fermentation of environmentally 88 relevant substrate throughout the digestive tract of the honey bee? Through utilizing a 89 combination of molecular, biochemical, and physiological approaches, both in vitro and 90 in vivo, we perform an in-depth metabolic analysis on 17 isolates from the honey bee 91 gut. Our results suggest that overall, bee-associated microbes are primed to metabolize 92 certain components of the honey bee diet, but that significant variation exists within 93 clades, suggesting that each member contributes differentially to honey bee metabolism. 94 95 Materials and Methods: 96 Bee Sampling and Bacterial Culture 97 Worker honey bees were collected from colonies located at an apiary in 98 Bloomington, IN and transported to lab in sterile vials on ice. Hindgut dissections were 99 performed using aseptic technique and sterile equipment. Whole digestive tracts of 2-3 100 bees were homogenized in 1X phosphate buffered saline (PBS) with a sterile disposable 101 pestle and a dilution series was plated onto Brain-Heart Infusion (BHI), de Man, Rogosa 102 and Sharpe (MRS), and Luria-Bertani (LB) agar (BD Difco™). All agar plates were 103 incubated under anaerobic conditions and supplemented with CO<sub>2</sub> using the GasPak EZ

- 104 system (BD, New Jersey) at  $37^{\circ}$ C for 4 5 days. Isolated colonies were subcultured on
- agar before genotyping (below).

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107 16S rRNA genotyping of bacterial isolates

108 For each isolate, genomic DNA was extracted using the QIAGEN DNeasy Blood &

109 Tissue kit with one modification: the inclusion of a bead-beating step before lysates were

110 loaded onto the Qiagen column. DNA was quantified using a spectrophotometer (BioTek

111 instruments) and used in PCR using Phusion® High Fidelity PCR mix with HF buffer 112 (New England BioLabs INC.), with 27F (5'-AGA-GTT-TGA-TCC-TGG-CTC-AG-3') and 113 1492R (5'-ACG-GCT-ACC-TTG-TTA-CGA-CTT-3') primers. A reaction consisted of 0.4 114 mM primers, 1x Phusion mix, PCR grade water, and 1 µL of sample DNA, under the 115 following conditions: 5 min at 98°C; 35 cycles-10 sec at 98°C, 30 sec at 55°C, 30 sec 116 72°C; Extension-10 min at 72°C. PCR products were visualized using gel 117 electrophoresis and all isolates with PCR products of the correct size were used for 118 sequencing (Beckman Coulter Genomics). Resulting ab1 files were concatenated and 119 exported to Fasta. To align, trim, and classify sequences, Mothur v.1.33.3 software was 120 used with the following commands (align.seqs, trim.seqs, and classify.seqs), using a 121 custom reference database (Greengeen + honey bee specific dataset) (Newton and 122 Roeselers, 2011) for classification. To demonstrate the evolutionary relationship 123 between isolates, a 16S rRNA gene phylogeny was constructed, using the concatenated 124 16S rRNA gene sequence used for classification. The tree was rooted with Aquifex 125 aeolicus (Accession: AJ309733.1) and to serve as representatives of the four dominant 126 bacterial families within the honey bee (Gilliamella apicola - Accession: NR 118433.1 127 (Gilliamella), Lactobacillus sp. wkB8 - Accession: NZ CP009531 - Locus Tag: 128 LACWKB8 RS00480 (Firm-5), Uncultured Lactobacillus sp. - Accession: HM113352.1 129 (Firm-4), and Bifidobacterium asteroides strain Hma3 - Accession: EF187236.1 (Bifido) 130 were used. Sequences were uploaded to SINA 1.2.11 aligner (ARB SILVA), and aligned 131 using the default settings, after which the FASTA formatted alignment was converted to 132 MEGA format (MEGA v6.06) and a maximum likelihood tree (GTR + G + I) was 133 constructed using 1000 bootstrap replicates. Divergence estimates were performed in 134 BLASTClust (Alva et al, 2016).

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136 Carbohydrate Utilization & Byproduct Production of Isolates

137 To prepare cell suspensions to assay substrate utilization and product formation. 138 all isolates classified as actinobacteria or bacilli were grown on MRS agar, while gamma-139 proteobacteria isolates were grown on BHI: these media correspond to the medium 140 under which the microbes were isolated. Broth cultures were grown anaerobically at 141 30°C for 6 days, after which cultures were pelleted via centrifugation and the supernatant 142 removed. The pellet was washed with salt solution (137.0 mM NaCl, 2.7 mM KCl, pH 143 6.0) to remove remaining media. Bacilli and actinobacteria, grown in MRS, were washed 144 twice to decrease the transfer of nutrients present in the spent MRS media. Isolates 145 were normalized for growth based on OD, across taxonomic classes, such that isolates 146 with more robust growth were diluted in a larger amount of salt solution. 147 To test the ability of isolates to utilize carbon substrates, we used the MT2 148 MicroPlate<sup>™</sup> assay. In the assay, each well in a 96-well plate contained an aliguot of a 149 TTC dye. To each well, 150 µL of cell suspension from each isolate was added, with or 150 without 15 µL of a filter-sterilized carbon source (either 1.1 M of D-(+)-glucose (SIGMA 151 G8270), D-(-)-fructose (ACROS Organics 57-48-7), D-sucrose (Fischer Scintific BP220-152 1), D-(+)-mannose (SIGMA M6020), L-rhamnose (SIGMA W373011), D-(+)-galactose 153 (SIGMA G0625), D-(+)-xylose (SIGMA X1500); or 13.1% D-(+)-cellobiose (SIGMA 154 22150), or 2.1% pectin (SIGMA – Pectin from apple – SIGMA 76282) solutions). 155 Negative controls received 15 µL of salt solution. To verify that substrate solutions were 156 void of microbes and that cross-contamination between wells was not occurring, an 157 additional negative control was filled with 15 µL of a sole carbon source with 150 µL of 158 salt solution. Each experimental condition and control was analyzed in triplicate within a 159 single plate. 96 well plates were covered with an optical adhesive film, and incubated 160 anaerobically at 30°C for 3 days. To measure reduction of TTC (and utilization of a 161 substrate), absorbance readings at 590 nm were taken, in triplicate, for each well. 162 Significant utilization was determined by performing a t-Test comparing 590 nm readings

163 between experimental (cell suspension + 15 µL substrate) and negative control (cell 164 suspension + 15 µL of salt solution) wells. All statistics were performed in IBM SPSS 165 Statistics Version 24 on datasets tested for a normal distribution. Biological triplicates for 166 each condition or control were then combined in a screw-top cryovial and stored at -80°C 167 until high performance liquid chromatography (HPLC) analysis (below). 168 To determine the soluble fermentation products, pooled samples were thawed, 169 vortexed, and then centrifuged at 13,000 RPM for 5 minutes to pellet bacterial debris. 170 After centrifugation, samples were filtered through a 0.45 µm HAWP membrane 171 (Millipore Corp., Bedford, Mass.) and acetate, lactate, formate, and succinate were 172 quantified by using a Shimadzu HPLC (Kyoto, Japan) as previously described (17). 173 Assaying the expression of enzymatic genes 174 For each dissection, the crop, midgut, and hindgut were aseptically incised from 175 the full digestive tract, and placed in a 2.0 mL Lysing matrix E tube with 500 uL of 176 Trizol® reagent. Samples were homogenized using a FastPrep®-24 instrument, twice, at 177 6.0 m/s for 40 seconds. RNA were extracted from samples using the standard protocol 178 provided by the manufacturer. After the final washing, RNA was resuspended in RNA 179 Storage Solution (Bioline) and stored at -80°C for downstream analysis. For each gut 180 chamber, each biological replicate consisted of three separate individuals. All analyses 181 were performed in triplicate with the same biological samples. 182 Primer sets specific for a particular enzymatic gene and group of honey bee 183 microbes (Gilliamella, Firm-5, and Bifido), were designed for gRT-PCR analyses 184 (Supplementary Table 1, Supplementary Figure 2). Selected primer sets fulfilled the 185 following requirements: 1) they amplified a product of the expected size, 2) the resulting

sequenced product had a top BLAST hit(s) to the intended bacterial taxa and the gene of
interest, or for the reference sequence used to design the primer set and 3) they showed

188 specificity for the bacterial group of interest in laboratory experiments (Supplemental

189 Table 1, Supplemental Figures 2). RNA extracted from individual gut chambers 190 (foregut, midgut, and hindgut) were used as a template for gRT-PCR reactions. Prior to 191 running reactions, all RNA samples were treated with RNase-free DNase I (Ambion) as 192 prescribed by the manufacture. RT-qPCR was performed using SensiMix ™ SYBER R 193 Hi-ROX One-Step kit and protocol. As a reference locus in these analyses, we used the 194 16S rRNA gene, using a universal bacterial primer set (S-\*-Univ-05150a0S-19/S-D-Bact-195 0787-b-A-20 (18)). Reactions were run on an Applied Biosystems StepOne RT-PCR 196 system: Initial denaturation-5 min at 98°C; 40 cycles -15 sec at 98°C, 1 min at 60°C, 30 197 sec 72°C; Extension-10 min at 72°C. A standard curve was generated for each primer 198 set separately, to account for primer efficiencies in our analyses. All statistics were 199 performed using IBM SPSS Statistics Version 24. 200 To quantify the expression of metabolic genes from honey bee specific clades of 201 bacteria, across gut chambers of the honey bee, Ct values acquired from the 202 quantification analyses of individual genes were initially normalized to the mean

203 amplification of the reference locus (16S rRNA gene) across gut sections. The

advantage of normalizing the data to 16S RNA gene is that it demonstrates the presence

205 of genes, relative to a proxy for the whole bacterial community present throughout the

206 digestive tract of the honey bee. Next, for each metabolic gene prescribed to a taxa of

207 interest, we standardized the data to the gut section with the lowest observed

amplification (i.e the crop) for calculating the relative expression (RNA). To determine

209 the relative ratio of expression of metabolic loci attributed to a particular taxa, across gut

sections, normalized datasets were standardized to the gut section and locus with the

211 lowest level of expression. To determine variation in the relative abundance, relative

212 expression, or relative ratio of expression of loci across gut sections, we performed an

- 213 ANOVA or Kruskal-Wallis test on the standardized data acquired for each individual
- 214 gene analyzed, which was dependent on results obtained from normality test of

individual gene datasets. Results with statistical significance ( $p \le 0.05$ ) were subjected to

216 pairwise comparisons using one-way ANOVA/post hoc or Mann-Whitney test.

217

218 **Results**:

219 The honey bee core microbiota utilizes carbohydrates from the host diet

220 To determine the ability of gut isolates to utilize environmentally relevant 221 substrates, we isolated 17 honey bee associated bacteria, spanning three 222 transcriptionally dominant clades: Gilliamella, Lactobacillus Firm-5, and Bifidobacterium 223 (Figure 1, Supplemental Figure 1). Based on the construction of a 16S rRNA gene 224 phylogeny, including data from the 17 cultured isolates and 145 sequences previously 225 obtained from bee-associated microbiota, some of the microbial cultures used herein are 226 closely related to core honey bee bacteria while others fall outside of the core clades 227 (Supplemental Figure 1). When comparing the diversity within the clades (based on 16S) 228 rRNA genes), all isolates are unique at 98% and 99% divergence, and some are unique 229 at 97% divergence (Table 1), indicating that these microbes represent different bacterial 230 strains and species.

231 To test the ability of gut bacteria to utilize sole carbon sources relevant to the 232 honey bee diet, we used Biolog MT2<sup>™</sup> plates, containing an aliguot of an indicator 233 tetrazolium chloride dye (TTC), that when combined with a sole carbon source, along 234 with an aliguot of culture, detects substrate utilization via a concomitant color change 235 due to reduction of the dye. Each isolate was examined for the sole utilization of 9 236 carbon sources (glucose, fructose, sucrose, xylose, pectin, cellobiose, mannose, 237 rhamnose, and galactose) in pure culture under anaerobic conditions. Honey, which 238 provides the host with simple carbohydrates (e.g. glucose, fructose, and sucrose) and 239 trace amounts of secondary sugars (e.g. mannose, rhamnose, and galactose) (19-21). 240 Matured pollen (beebread), provides the host with essential nutrients (i.e. amino acids, 241 minerals, lipids), and is also rich in complex plant-derived substrates (e.g. cellobiose,

242 xylose, and pectin)(22, 23). Therefore, the compounds selected for analysis have

243 previously been identified, through biochemical assays, as molecules that constitute part

of the honey bee diet.

245 When examining the individual and collective data for the utilization of glucose. 246 fructose, and sucrose, nearly all isolates were able to utilize at least one or more of 247 these substrate(s), with the exception of 4 taxa: Staphylococcaceae EBHJ3 and 248 Lactobacillales MAH G07-3 within the Firmicutes and Enterobacteriaceae LB Tet D09 249 and LB Tet C11 (Figure 2, Supplemental Figure 2). At the phylum-level, at least 3 250 isolates within each phylum were capable of utilizing glucose, fructose, and sucrose, 251 with varying degrees of consistency (Figure 2A, Supplemental Figure 2A). Collectively, 252 when comparing isolates representative of taxa specific to the honey bee gut microbiota 253 (i.e. Gilliamella, Lactobacillus Firm-5, and Bifidobacterium), the general trend for 254 substrate utilization suggested that the honey bee bacteria were primed to utilize these 255 substrates (Figure 2). Interestingly, however, we did observe differences among strains 256 within clades in their ability to use different compounds. For example, although all 257 Bifidobacteria and Lactobacillus Firm-4 strains used glucose, fructose, and sucrose, the 258 Lactobacillus Firm-5 strain MAH H05-3 did not metabolize either monosaccharaide to a 259 significant extent. Similarly, Gilliamella strains differentially processed sucrose; although 260 both strains tested could utilize glucose and fructose, only Gamma-1 LB D06 could 261 utilize sucrose, although to a modest extent (Figure 2).

262

Acetate, lactate, succinate, and formate are the primary fermentation products produced
by honey bee gut microbiota

To determine the soluble fermentation products produced by each individual
isolate, post anaerobic metabolism, we analyzed the contents within each experimental

267 and control well of the MT2<sup>™</sup> plates described in the previous section using HPLC. For 268 this analysis, we focused on the products in pure cultures with glucose, fructose, and 269 sucrose. We chose to use simple sugars abundant in honey, rather than complex 270 polymeric carbon sources, because complex carbon sources are degraded to simple 271 sugars and funneled into the same catabolic pathways. With the exception of pectin, all 272 other saccharides analyzed for utilization are mono- or di- saccharides and are only a 273 few enzymatic reactions away from glycolysis or the pentose phosphate and Entner-274 Doudoroff pathways. While the ratio of fermentation products is expected to change 275 during growth on different substrates, it is less likely that different products would be 276 produced from different substrates. We used known standards (e.g., acetate, lactate, 277 formate, succinate) and identified peaks at the appropriate retention times within each 278 biological sample.

279 HPLC analysis revealed a variety of organic acids produced by honey bee 280 associated microbes, under anaerobic conditions. When comparing the supernatant of 281 experimental and control wells of isolates, almost all isolates that were positive for 282 utilizing a substrate exhibited the presence of fermentative products (as derived from 283 retention times identical to known substrate standards, *i.e.* lactate, acetate, succinate, 284 and formate; Figure 2). We expected to observe ethanol, acetoin, propanoate, 2,3-285 butanediol, based on the metabolic pathways identified in the metatranscriptomic data 286 but did not identify other significant peaks beyond the four described above, even though 287 the instrument is capable of detecting these compounds.

When comparing the general pattern of products produced across taxa, bacteria classified within the *Bifidobacterium* clade unanimously produced lactate and acetate, regardless of substrate. However, production of succinate and formate by this same clade was variable both across the phylogeny and across substrates (Figure 2). The Firm-5 and *Gilliamella* isolates produced these same organic acids, but with an even

293 larger degree of variability within the clades (Figure 2). Altogether, the data for the 17 294 isolates supports the hypothesis that honey bee gut microbiota are primed to consume 295 and ferment the components of the honey bee diet, but significant variation exists within 296 strains such that no uniform profile could be ascribed to a clade or species group. For 297 example, even if isolates of the same species used a substrate, they did not produce the 298 same organic acids; formate was detected from some strains of Bifidobacterium under 299 glucose but not under fructose, while all produced formate when given sucrose (Figure 300 2).

301

#### 302 Microbes utilize more complex carbohydrates and sugars toxic to the honey bee

303 We extended our analysis to plant derivatives of pollen (cellobiose, xylose, and 304 pectin), as well as secondary sugars commonly found in honey (mannose, rhamnose, 305 and galactose) and known to be toxic to the host in high guantities. For the plant derived 306 carbohydrates, we observed the same trend noted for the utilization of simple sugars, 307 with at least 2 isolates within each major taxon capable of utilizing cellobiose, xylose, 308 and pectin. Interestingly, all honey bee specific bacteria tested could utilize pectin while 309 more distantly related bacteria (Staphilococcaceae, Lactobacillales, Enterobacteriaceae) 310 could not (Figure 3). However, we found variation in the ability of strains within honey 311 bee associated clades in their use of other complex carbohydrates; xylose was only 312 utilized by half of the *Bifidobacterium* strains tested and only 1 out of 4 *Lactobacillus* 313 strains. In contrast, we observed no difference between honey bee specific and more 314 distantly related bacteria in their ability to utilize sugars toxic to the bee (Figure 3). Again, 315 however, we observed variation in the ability of strains to utilize these toxic sugars with 316 only some core members able to use all of these toxic sugars and some core members 317 unable to use any (*Bifidobacteriales* LB C04 and *Lactobacillus* Firm-4 MaH G10-2, 318 Figure 3). Overall, these results suggesting the honey bee associated bacteria utilize

mannose, rhamnose, and galactose, secondary sugars common in the honey bee diet,but toxic to the host.

321

322 The honey bee gut compartments are associated with altered gene expression by the 323 microbiota

324 We determined that isolates from the honey bee gut microbiota ferment sugars 325 common to the honey bee diet into organic acids in vitro (Figure 2). Recent work by 326 Zheng et al. (2017) demonstrated *in vivo*, the presence of a variety of organic acids 327 across distinct gut chambers of the honey bee digestive tract. Based on the 328 observations described above, we hypothesized that symbionts of the honey bee gut are 329 likely contributing to presence of organic acids *in vivo*, and since each gut chamber 330 houses a specific compilation of symbionts (24, 25), perhaps functional differences exist 331 in the metabolic activity of symbionts across individual gut chambers. To address this 332 hypothesis, we quantified the expression of genes that encode for fermentative enzymes 333 throughout the three gut chambers of the honey bee digestive tract (i.e. crop, midgut, 334 and hindgut (anterior to posterior)). We designed taxon-specific primers for two genes: 335 acetate kinase (ackA) and L-lactate dehydrogenase (Idh), encoding for enzymes 336 responsible for the production of acetate and lactate, respectively. In addition, we also 337 designed primers for clade specific enzymes (that is, the metabolic function was only 338 identified in one of the three phyla involved in cleaving glycosidic bonds within plant-339 derived compounds (e.g. cellulase and cellodextrinase for *Bifidobacteria*, pectate lyase 340 for *Gilliamella*, and glucosidase for the *Lactobacillus* Firm-5 clade; Supplementary Table 341 1)). We used gRT-PCR on RNA extracted from each gut section to quantify the 342 expression of these loci. For the Gilliamella and Lactobacillus Firm-5 clades, the midgut 343 and hindgut chambers were found to be the most active gut sections (Gilliamella (df = 2;  $x^2 = 7.05$ ; p = 0.029), Firm-5 (df = 2; F = 3.95; p = 0.040); Figure 4A and C). In contrast, 344

345 while the activity of the *Bifidobacterium* clade increases across gut sections from 346 posterior to anterior, the trend is not significant ((df = 2; 2 = 4.36; p = 0.113); Figure 4B). 347 To obtain a better understanding of genes responsible for the trends seen in the 348 Gilliamella and Lactobacillus Firm-5 clades (which both resulted in statistically significant 349 contrasts across the digestive tract), we performed a one-way ANOVA/Tukey-HSD to 350 analyze the differences in variation of relative gene expression across gut sections. For 351 Gilliamella, the analysis of ackA and ldh expression indicated that expression levels 352 significantly increased across gut sections in the following order: crop<midgut<hindgut 353 (Figure 4D). Pectate lyase was detected at equal levels of expression in the midgut and 354 hindgut, but no amplification was detected in the crop (Figure 4D). In contrast to 355 Gilliamella, we saw no pattern to the expression of ldh or ackA by Lactobacillus Firm-5 356 across the gut sections (Figure 4E). Importantly, we were not able to detect  $\beta$ -357 glucosidase and cellulase gene expression from Firm-5 and *Bifidobacterium* clades, in 358 any gut section, suggesting that they are not active in the honey bee gut. 359 360 **Discussion:** 

361

362 The honey bee gut microbiota is a deceptively simple community – although 363 dominated by 8-10 species (26) and transcriptionally dominated by only 3 phyla (5), the 364 strain diversity found within the gut is high (16). Here we show that strains within three 365 transcriptionally dominant clades associated with the bee are functionally distinct and 366 that there is indeed, no strict metabolic rule that describes each clade. Although overall, 367 the microbes seem primed to utilize carbohydrates in the honey bee diet, expressing 368 enzymes capable of cleaving glycosidic bonds and fermenting these substrates, strains 369 within each clade of core microbes investigated here differ in their ability to use different 370 simple sugars, sugars toxic to the bee, or complex carbohydrates found in the bee diet

371 (Figures 2, 3). They also differ in their production of organic acids, the production of 372 which are suspected to contribute to honey bee metabolism. The results presented in 373 this study reveal that bacterial taxa associated with the honey bee gut are differentially 374 present and metabolically active throughout the three major gut chambers, with 375 particular enrichment in the hindgut. Of the three taxa assayed for gene expression, the 376 Gilliamella clade of bacteria was the most active, showing exceptional expression of its 377 acetate kinase gene in the mid- and hindgut, a result supported by previous 378 metatranscriptomic analyses.

379 The utilization results presented in this study are supported by previous 380 observations of gross pH changes when honey bee isolates are cultured (9, 11-14) and 381 previous measures of acetic and lactic acid production from a few pure cultures (27, 28). 382 Although our results are well supported by predicted models based on the genomic 383 evidence, we did not detect the production of specific fermentative waste products 384 predicted (e.g. ethanol, acetoin, propanoate, 2,3-butanediol)(5). The lack of detection of 385 these products in our analysis could be due to the following: 1) a complete lack of 386 production under the *in vitro* conditions provided 2) or production below the detectable 387 range of the UV and refractive index detectors used.

In addition to analyzing the utilization of the substrates mentioned above, we also
examined the ability of isolates to utilize pectin, which is a heteropolysaccharide,

390 commonly found in the primary cell wall of terrestrial plants, such as pollen grains. In our

analysis, we identified multiple isolates within the *Gilliamella, Lactobacillus* Firm-5, and

392 Bifidobacterium clades able to utilize pectin (Figure 3). Importantly all tested core

393 microbiome strains utilized pectin in our assays – it was the only consistently utilized

394 carbohydrate tested – and more distantly related organisms (e.g. Staphylococcaceae,

395 Lactobacillales, Enterobacteriaceae) were unable to use it. Previous work has shown

that *Gilliamella* isolates are capable of degrading pectin, possessing a pectate lyase

397 gene capable of cleaving de-esterified pectin (7). In addition to Gilliamella, we also 398 report utilization of pectin and/or its derivatives by isolates within the Firm-4/-5 and 399 *Bifidobacterium* clades, observations supported by a recent publication that observed 400 growth of several core gut isolates in minimal media supplemented with pollen extract 401 (15). These results suggest that multiple constitutes of the honey bee microbiota may be 402 capable breaking-down and utilizing plant material abundant in the host's diet, an idea 403 that has previously been hypothesized (10). Indeed, the genome of *Bifidobacterium* 404 asteroides has an annotations for glycosidases and a pectinesterase, enzymes capable 405 of degrading pollen (6). Contrary to Bifidobacterium, genomes from bacteria in the 406 Lactobacillus Firm-5 clade currently have no annotated pectinases or glycosidases (8), 407 but these and many of the genomes of honey bee specific bacteria possess a plethora of 408 hypothetical genes of unknown function that might aid in the degradation and 409 metabolism of pectin (9). Importantly, although the pectin used in our *in vitro* assays was 410 of high quality, it is not a completely uniform nor pure substance and therefore it is 411 possible that trace amounts of other compounds might have been present and 412 metabolized by the honey bee specific microbes. Further experiments need to be 413 conducted to confirm the pectinase activity of the isolates analyzed. It should also be 414 noted, that the conditions provided in the utilization assay likely do not support the 415 growth of all 17 isolates as the salt solution lacks several elements required for growth. 416 However, although the assay does not rely on growth, it is possible that the utilization 417 patterns observed for individual isolates could potentially alter under alternative 418 conditions conducive for growth.

Recent important work has identified a link between the presence of canonical gut symbionts and weight gain in honey bees; organic acids, which are presumed to be bacterially derived, are thought to contribute to the observed influence symbionts to host health (28). The fact that these microbes live within the bee gut and are consistently

423 associated with honey bees has led to the assumption that microbes isolated from the 424 bee are mutualists in this context – that they benefit the bee in some way. However, 425 supplementation of the bee diet with one of these core members (Snodarasella) results 426 in an increased susceptibility to parasites (29) and another (Frischella) results in scab 427 formation in the intestine (7). The converse can also be true; non-core members of the 428 bee can provide a benefit as well. Lactobacillus kunkeii, an environmentally acquired 429 microbe, protects from pathogens (30) and although Gilliamella strains have been found 430 to use sugars that are toxic to the bee (31), here we show that both non-core bacteria 431 and core microbes can utilize these sugars.

432The difference in the enzymatic activity pattern between members of the honey433bee microbiota suggests they might occupy different niches in the honey bee food,

434 digestive tract, or built environment. The idea that niche partitioning might exist in the

435 gut of the honey bee is physically plausible, as the alimentary system in the honey bee is

436 subdivided into three distinct chambers. Indeed, our analysis of gene expression (qRT-

437 PCR) suggests that expression of different enzymes can differ across gut sections,

438 genes, and taxa. It should be noted, however, that we could not detect the expression of

439 genes responsible for the breakdown of complex plant-derived molecules (i.e.

440 glucosidase, cellodextrinase, and cellulase). These results indicate that these genes are

441 either completely inactive, or are expressed at levels below detection. In either scenario,

the data suggest that perhaps complex carbohydrates are either present at minimal

443 levels in honey bee gut, or that the taxa analyzed in this study thrive on other resources

444 abundant in the environment (i.e. glucose, fructose, sucrose, and/or other plant

445 derivatives) by expressing other enzymatic loci capable of contributing to catabolism.

446 Our work suggests a dominant profile for *Gillamella in vivo*. Of the three analyzed 447 taxa, the *Gilliamella* clade was the most active in our analysis, with the *Bifidobacterium* 448 and *Lactobacillus* Firm-5 clades being roughly equal; this result fits well with previously

449 published transcriptomic data, where the gamma-proteobacteria dominated the 450 transcriptional profile of the gut (5). When assaying the relative expression of certain 451 genes (i.e. ackA and ldh) for the Gilliamella clade (Figure 4A, D), we saw highest levels 452 of expression in the hindgut, suggesting that Gilliamella may function differently in each 453 gut chamber or may be particularly active in the hindgut. Additionally, within each gut 454 section, Gilliamella consistently expresses more ackA than ldh, which might suggest a 455 preference for this taxon to produce acetate, a byproduct produced in vitro by Gilliamella 456 isolates under nearly all tested conditions (Figure 2). It is intriguing that *Gilliamella* 457 dominates transcriptomic analyses of the honey bee and related species are found in 458 bumble bees (8). Gilliamella and other proteobacterial species in the honey bee require 459 direct social contact between bees beyond mere trophallaxis (i.e. fecal exposure) to 460 achieve transmission (32). The fact that the honey bee workers acquire this important 461 symbiont through their social interactions, and that Gilliamella dominates the functional 462 profile of the honey bee worker gut suggests it to be an important, if not critical, part of 463 the honey bee core microbiota.

464 Recent work elegantly demonstrated through metabolomics that honey bees 465 primarily colonized by individual bacterial strains of taxa core to the gut microbiota show 466 depletion in plant derived substrates and an enrichment in fermentation product present 467 in the digestive system, compared to bees with a depleted gut microbiota (33). These 468 results are promising, as they demonstrate substrate utilization by individual core taxa 469 and enrichment of metabolites within the gut in vivo. However, our work here suggests 470 that the choice of strain used in colonization experiments may substantially influence 471 results; not all Gilliamella apicola strains produce the same organic acids under the 472 same conditions. It is perhaps the case that the diversity of strains found within the 473 honey bee gut complement each other, providing a range of organic acids under most 474 conditions and diets. This hypothesis awaits testing. Nevertheless, it is obvious that

475	individ	lual taxa found in the core microbiome of bees might not accurately reflect the			
476	metab	olic dynamics occurring in the gut of a honey bee with a natural microbiota, where			
477	microbe-microbe and microbe-host interactions likely govern niche partitioning and				
478	microbiome function.				
479					
480	Ackno	owledgements:			
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484					
485 486	Conflicts of interest statement:				
487 488	The authors declare no conflict of interest.				
489					
490	Refer	ences:			
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### 1 Tables:

- 2 **Table 1.** Cluster analysis of the 16S rRNA gene sequences of 17 cultured honey bee gut isolates
- 3 used for metabolic analysis. A FASTA file with near full-length 16S rRNA sequences for each
- 4 isolate was uploaded to BLASTClust (Alva et al., 2016). Standard settings were used for the
- 5 analysis, using 97% or 98% identity thresholds. The total number of isolates analyzed for each
- 6 class of bacteria is provided, along with the number of phylotype clusters at 97% and 98% ID.

Bacterial Class	Total Number of isolates	Number of Phylotypes (%ID)	
	isolates	97	98
y-Proteobacteria	5	5	5
Bacilli	6	5	6
Actinobacteria	6	1	6

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Figures:

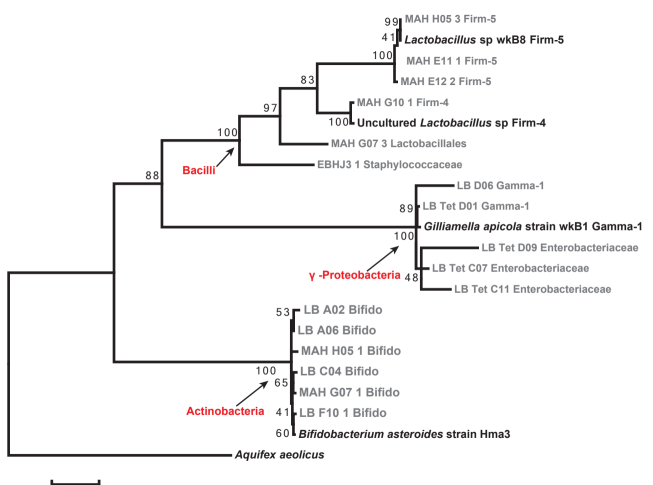
**Figure 1.** 16S rRNA gene phylogeny of gut isolates selected for metabolic analyses, in relationship to reference sequences of honey bee specific taxa. Sequences were aligned using SINA Aligner 1.2.11, the output file was used to construct a Maximum likelihood phylogeny, in MEGA 6.06 with GTR (G+I) and 1000 bootstraps. Bootstrap support is provided at nodes. Reference sequences indicated by black text. Cultured isolates indicated by grey text. Taxonomic class highlighted by arrow.

**Figure 2.** Phenotypic profile of substrate utilization by gut isolates. On the left of the figure are consensus phylogenetic trees derived from the 16S rRNA gene phylogeny. Utilization column is a heat map representation of OD590 measurements (background subtracted) from 1.4 (most purple) to zero (white). Asterisks indicate statistically significant results, compared to negative control, for glucose (A), fructose (B), or sucrose (C). Similarly, the detection of organic acids is denoted by boxes filled will the color that corresponds to a given compound detected: formate (blue), acetate (green), lactate (orange), and succinate (red), based on retention peaks on HPLC chromatograms.

**Figure 3.** Phenotypic profile of substrate utilization by honey bee isolates. On the left are cladograms derived from the 16S rRNA gene phylogeny. Isolates highlighted in grey indicate taxa considered core to the honey bee gut, while all other isolates are representative of non-core members. The utilization of other plant-derived sugars (i.e. cellobiose, xylose, and pectin) and dietary saccharides toxic to the host (mannose, rhamnose, and galactose) is a heat map representation of OD590 measurements

(background subtracted) from 1.4 (most purple) to zero (white). Asterisks indicate statistically significant results, compared to negative control.

**Figure 4.** Expression of metabolic genes derived from honey bee specific taxa. Reverse transcriptase quantitative PCR (RT-qPCR) expression shown for each enzymatic genes, relative to the average 16S rRNA gene expression, across chambers of the honey bee gut. Standardized to data from the crop, the overall relative ratio of expression of enzymes for *Gilliamella* (A), *Bifidobacterium* (B), and *Lactobacillus* clades(C) were determined across gut chambers. (D-F) The relative expression data of individual genes, for *Gilliamella* (D), *Bifidobacterium* (E), and *Lactobacillus* (F), across the gut sections. For each dataset, relative expression data were compared across gut sections, using ANOVA or Kruskal-Wallis test in SPSS. For statistically significant results ( $p \le 0.05$ ), pairwise comparisons were conducted using one-way ANOVA/post hoc or Mann-Whitney test in SPSS, significant difference ( $p \le 0.05$ ) between gut sections are denoted with letters.

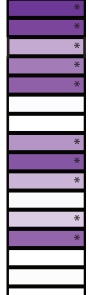


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bioRxiv preprint doi: https://doi.org/10.1101/294249; Hisilizationsted April 4, acta te copy Acetate for the Superinate ich Formate certified by peer review) is the author/funder, who has granted blocking a license to display t Bifidobacteriales LB C04 aCC-BY-NC-ND 40 International license

**Bifidobacteriales MAH G07-1** B

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**Bifidobacteriales LB C04 Bifidobacteriales MAH G07-1 Bifidobacteriales LB F10-1 Bifidobacteriales MAH H05-1 Bifidobacteriales LB A02 Bifidobacteriales LB A06** Staphylococcaceae EBHJ3-1 Lactobacillales MAH G07-3 Firm-4 MAH G10-2 Firm-5 MAH E12-2 Firm-5 MAH E11-1 Firm-5 MAH H05-3 Gamma-1 LB D06 Gamma-1 LB Tet D01 **Enterobacteriaceae LB Tet D09** Enterobacteriaceae LB Tet C11 Enterobacteriaceae LB Tet C07

**Bifidobacteriales LB C04 Bifidobacteriales MAH G07-1 Bifidobacteriales LB F10-1 Bifidobacteriales MAH H05-1 Bifidobacteriales LB A02 Bifidobacteriales LB A06** Staphylococcaceae EBHJ3-1 Lactobacillales MAH G07-3 Firm-4 MAH G10-2 Firm-5 MAH E12-2 Firm-5 MAH E11-1 Firm-5 MAH H05-3 Gamma-1 LB D06 Gamma-1 LB Tet D01 Enterobacteriaceae LB Tet D09 Enterobacteriaceae LB Tet C11 Enterobacteriaceae LB Tet C07

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<u>Plant-Derivatives</u>					
<u>Cellobiose</u>	<u>Xylose</u>	<u>Pectin</u>			
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# Sugars Toxic To The Host

Mannose Rhamnose Galactose				
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