

AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Bacterial diversity shift determined by different diets in the gut of the spotted wing fly *Drosophila suzukii* is primarily reflected on acetic acid bacteria.

This is the author's manuscript

Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/1634704> since 2017-11-23T16:16:59Z

Terms of use:

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)

This is the author's final version of the contribution published as:

VACCHINI V., GONELLA E., CROTTI E., PROSDOCIMI E., MAZZETTO F., CHOUAIA B., CALLEGARI M., MAPELLI F., MANDRIOLI M., ALMA A., DAFFONCHIO D. – Bacterial diversity shift determined by different diets in the gut of the spotted wing fly *Drosophila suzukii* is primarily reflected on acetic acid bacteria. *Environmental Microbiology Reports* 9(2), 2017, 91–103.

The publisher's version is available at:

<http://onlinelibrary.wiley.com/doi/10.1111/1758-2229.12505/full>

When citing, please refer to the published version.



Bacterial diversity shift determined by different diets in the gut of the spotted wing fly *Drosophila suzukii* is primarily reflected on acetic acid bacteria

Journal:	<i>Environmental Microbiology and Environmental Microbiology Reports</i>
Manuscript ID	Draft
Journal:	Environmental Microbiology Reports
Manuscript Type:	EMIR - Brief report
Date Submitted by the Author:	n/a
Complete List of Authors:	Daffonchio, Daniele; King Abdullah University of Science and Technology, Biological and Environmental Sciences & Engineering Crotti, Elena; University of Milan, DEFENS
Keywords:	cultivation-dependent approach, fluorescent in situ hybridization (FISH), symbionts, green fluorescent protein, 16S rRNA gene pyrosequencing

SCHOLARONE™
Manuscripts

1 **Bacterial diversity shift determined by different diets in the gut of the spotted wing fly *Drosophila***
2 ***suzukii* is primarily reflected on acetic acid bacteria**

3

4 Violetta Vacchini^{1#}, Elena Gonella^{2#}, Elena Crotti^{1#}, Erica M. Prosdocimi^{1°}, Fabio Mazzetto², Bessem
5 Chouaia^{1§}, Matteo Callegari¹, Francesca Mapelli¹, Mauro Mandrioli³, Alberto Alma² and Daniele
6 Daffonchio^{1,4*}

7

8 ¹Dipartimento di Scienze per gli Alimenti, la Nutrizione e l'Ambiente (DeFENS), Università degli Studi
9 di Milano, Milano, Italy

10 ²Dipartimento di Scienze Agrarie, Forestali e Alimentari (DISAFA), Università degli Studi di Torino,
11 Grugliasco, Italy

12 ³Dipartimento di Scienze della Vita (DSV), Università degli Studi di Modena e Reggio Emilia, Modena,
13 Italy

14 ⁴Biological and Environmental Sciences and Engineering Division (BESE), King Abdullah University
15 of Science and Technology (KAUST), Thuwal 23955-6900, Kingdom of Saudi Arabia.

16 [#]These authors have equally contributed to the work

17 [°]Current address: Blizard Institute, Queen Mary University of London, United Kingdom

18 [§]Current address: Department of Entomology, 5142 Comstock Hall, Cornell University, Ithaca, NY,
19 14853, United States of America

20 ^{*}Corresponding Author: Daniele Daffonchio, BESE, Biological and Environmental Sciences and
21 Engineering Division, King Abdullah University of Science and Technology (KAUST), Building 2,
22 Level 3, Room 3236, Thuwal 23955-6900, Kingdom of Saudi Arabia; Tel: +966 (2) 8082884; Email:
23 daniele.daffonchio@kaust.edu.sa.

24

25 **Running title:** Acetic Acid Bacteria of *Drosophila suzukii*

26

27 Abstract

28 The pivotal role of diet in shaping gut microbiota has been evaluated in different animal models,
29 including insects. *Drosophila* flies harbour an inconstant microbiota among which acetic acid bacteria
30 (AAB) are important components. Here, we investigated the bacterial and AAB components of the
31 invasive pest *Drosophila suzukii* microbiota, by studying the same insect population separately grown
32 on fruit-based or non-fruit artificial diet. AAB were highly prevalent in the gut under both diets (90 and
33 92% infection rates with fruits and artificial diet, respectively). Fluorescent *in situ* hybridization and
34 recolonization experiments with green fluorescent protein (Gfp)-labelled strains showed AAB capability
35 to massively colonize insect gut. High-throughput sequencing on 16S rRNA gene indicated that the
36 bacterial microbiota of guts fed with the two diets clustered separately. By excluding AAB-related
37 OTUs from the analysis, insect bacterial communities did not cluster separately according to the diet,
38 suggesting that diet-based diversification of the community is primarily reflected on the AAB
39 component of the community. Diet influenced also AAB alpha-diversity, with separate OTU
40 distributions based on diets. High prevalence, localization and massive recolonization, together with
41 AAB clustering behaviour in relation to diet, suggest an AAB role in the *D. suzukii* gut response to diet
42 modification.

43

44 Keywords

45 16S rRNA gene pyrosequencing, cultivation-dependent approach, fluorescent *in situ* hybridization
46 (FISH), symbionts, green fluorescent protein

47

48 **INTRODUCTION**

49 **The insect** gut microbiota plays very critical and essential roles for the host biology, physiology and
50 immunity (Hamdi *et al.*, 2011). **Diet, together with other factors, such as environmental habitat, host**
51 **developmental stage and phylogeny, profoundly affect its diversity and structure, consequently**
52 **influencing insect functionality** (Colman *et al.*, 2012; Yun *et al.*, 2014).

53 In last years, increased attention has been focused on the study of the bacterial microbiota associated to
54 different species of drosophilid flies. *Drosophila* represents a powerful insect model for a vast array of
55 studies, **including the defence mechanism-based investigations and the exploration of host-commensal**
56 **interactions** (Erkosar *et al.*, 2013; Lee and Lee, 2014). With the aim to unravel host-microbiome
57 interactions beyond laboratory boundaries, researchers have been prompted to investigate the gut
58 microbiota diversity of different natural species of drosophilid flies (Chandler *et al.*, 2011; Wong *et al.*,
59 2013; Cox and Gilmore, 2007). **By using molecular techniques four bacterial families have been found**
60 **to be commonly associated to field-captured or laboratory-reared flies, namely Enterobacteriaceae,**
61 **Acetobacteraceae, Lactobacillaceae and Enterococcaceae** (Brummel *et al.*, 2004, Chandler *et al.*, 2011,
62 Corby-Harris *et al.*, 2007, Cox and Gilmore, 2007, Ren *et al.*, 2007, Ridley *et al.*, 2012, Ryu *et al.*, 2008,
63 Sharon *et al.*, 2010, Storelli *et al.*, 2011, Wong *et al.*, 2011; Wong *et al.*, 2013). **In particular,**
64 **Acetobacteraceae (acetic acid bacteria, AAB) are among the dominant taxa in laboratory-reared *D.***
65 ***melanogaster* (Ryu *et al.*, 2008; Wong *et al.*, 2011). Conversely, field-captured *Drosophila* flies show**
66 **an inconstant bacterial community, where AAB are, however, frequently associated (Wong *et al.*, 2013).**
67 AAB are a bacterial group widespread in sugar- and ethanol-rich matrices, such as flowers' nectar,
68 fruits, vegetables and fermented matrices, all niches shared by drosophilid flies and from which they can
69 pass to the *Drosophila* gut, a sugar- and ethanol-rich environment (Blum *et al.*, 2013; Cox and Gilmore,
70 2007 Crotti *et al.*, 2010). AAB establish a delicate interaction with the insect innate immune system,
71 being involved in the suppression of the growth of pathogenic bacteria in healthy individuals (Ryu *et al.*,
72 2008), but also the modulation of the insulin pathway and the enhancement of the larval developmental

73 rate, body size, intestinal stem cells activity and energy metabolism (Shin *et al.*, 2011). A beneficial role
74 of AAB has been also demonstrated for mosquito larval development (Chouaia *et al.*, 2012; Mitraka *et*
75 *al.*, 2013).

76 The spotted wing fly *Drosophila suzukii* Matsumura (Diptera: Drosophilidae), an endemic pest in
77 South-East Asia, has been accidentally introduced in USA, Canada and Europe (Cini *et al.*, 2012;
78 Hauser, 2011; Lee *et al.*, 2011). Unlike its relatives that attack rotten fruits, *D. suzukii* lays eggs on
79 healthy soft summer fruits where the larvae grow (Walsh *et al.*, 2011; Mitsui *et al.*, 2006). So far, little
80 information is available on the bacterial community associated to *D. suzukii* specimens collected in USA
81 (Chandler *et al.*, 2014), while just few other publications studied *Wolbachia* infection (Mazzetto *et al.*,
82 2015; Cattel *et al.*, 2016; Siozios *et al.*, 2013).

83 Considering AAB abundance and importance in drosophilid flies, we aimed to assess the effect of two
84 different diets (i.e. based or not on fruit) on the diversity of bacterial and AAB microbiota of *D. suzukii*.
85 Specifically, we evaluated the possibility that AAB are involved in the gut microbiota diversification
86 when insects are exposed to two different alimentary regimes. For studying the effect of diets on the
87 bacterial microbiota diversity, we first confirmed the significance of AAB in the *D. suzukii* gut. We
88 determined their prevalence, the gut localization through fluorescent *in situ* hybridization (FISH) and
89 the ability to recolonize the insect gut by using green fluorescent protein (Gfp)-tagged derivatives of a
90 series of strains from a *D. suzukii* isolate collection. As a second step of the study we assessed the
91 changes of the bacterial microbiota structure and diversity by means of cultivation-independent
92 techniques.

93

94 RESULTS

95 **Prevalence of *Wolbachia* and AAB.** Since *Wolbachia* is a frequent symbiont of drosophilid flies, the
96 prevalence of this bacterium has been evaluated on adults obtained both from fruit and artificial diet
97 rearings. In flies reared on fruit *Wolbachia* showed an infection rate of 66% (33 out of 50 positive

98 specimens). *Wolbachia* prevalence was significantly lower (GLM, $p < 0.05$) in individuals maintained
99 on the artificial diet (infection rate of 28%, 14/50 positives). Conversely, AAB occurred in almost all of
100 the analysed individuals reared on both food sources, with 90 and 92% infection rates in flies
101 maintained on fruits and artificial diet, respectively (45 and 46 out of 50 individuals) with no significant
102 difference in infection incidence (GLM, $p=0.727$).

103

104 **AAB isolation.** Since the condition of fruit-based rearing is the closest to the diet of *D. suzukii* in field
105 conditions, we concentrated our efforts on individuals reared on this diet; however, specimens reared on
106 artificial diet have been also included in the analysis. The final collection included 234 isolates that were
107 de-replicated according to the ITS fingerprinting profiles. 16S rRNA gene sequencing of representatives
108 of each ITS profile identified the isolates as belonging to *Komagataeibacter*, *Gluconacetobacter*,
109 *Acetobacter* and *Gluconobacter* genera (Yamada *et al.*, 2012a; 2012b), while only 16.3% of the isolates
110 did not belong to Acetobacteraceae family (Tab. 1). Twenty-eight isolates have been affiliated to the
111 *Acetobacter* genus, including the species *A. cibinongensis*, *A. indonesiensis*, *A. orientalis*, *A.*
112 *orleanensis*, *A. peroxydans*, *A. persici* and *A. tropicalis*. *A. persici* and *A. indonesiensis* were the most
113 represented species. Eighteen *Gluconobacter* isolates have been affiliated to three species, *G.*
114 *kanchanaburiensis*, *G. kondonii* and *G. oxydans*. The unique isolate of *G. kondonii* in the collection has
115 been collected from an adult fly fed on fruits, while *G. kanchanaburiensis* isolates have been obtained
116 from specimens reared on artificial diet. Twelve isolates collected from adults fed on fruit showed high
117 sequence similarity with *G. oxydans*. One hundred and twenty-three isolates have been assigned to
118 *Gluconacetobacter* and *Komagataeibacter* genera. In particular, 118 *Komagataeibacter* isolates have
119 been obtained from fruit-fed *Drosophila*. Due to the phylogenetic proximity of the species of this genus,
120 discrimination at the species level was not possible with the actual 16S rRNA sequencing. *Ga.*
121 *liquefaciens* isolates (no. 4) have been obtained from three pupae and one larva using the TA1 medium.

122 Finally, the attribution to either *Gluconacetobacter* or *Komagataeibacter* genera could not be
123 discriminated according to the actual 16S rRNA sequence (Tab. 1).

124

125 **Localization of AAB in the *D. suzukii* gut and colonization by Gfp-labelled strains.** Fluorescent *in situ*
126 hybridization (FISH) on the insect dissected organs using the AAB-specific probe AAB455, gave
127 positive signals in the proventriculus and the gut (Fig. 1), whereas no fluorescence was detected in the
128 absence of probe. The proventriculus epithelium gave a strong signal, observable by merging the
129 interferential contrast (Fig. 1c) with the fluorescent (Fig. 1b) images. Magnification in fig. 1d allowed
130 the visualisation of fluorescent AAB microcolonies adhering to the peritrophic matrix.

131 *Gluconobacter* cells have been observed in the midgut (Fig. 1g) suggesting the distribution of this genus
132 in the inner side of the intestinal lumen. Fig. 1e-h show *Gluconobacter* distribution (Fig. 1g) in relation
133 to the dispersal of *Eubacteria* (Fig. 1f), indicating that it is surrounded by other bacteria, presumably
134 AAB (Fig. 1d). However, we could not ascertain such hypothesis because all the attempts to design
135 specific probes effective for *Acetobacter*, *Gluconacetobacter* and *Komagataeibacter* genera, failed.

136 Strains *G. oxydans* DSF1C.9A, *A. tropicalis* BYea.1.23 and *A. indonesiensis* BTa1.1.44 have been
137 successfully transformed with a plasmid carrying the Gfp cassette. Plasmid stability experiments
138 showed that *G. oxydans* DSF1C.9A retained the plasmid with a relatively high percentage (73.1%),
139 while this was not the case for strains BYea.1.23 and BTa1.1.44. Thus, colonization experiments of
140 adult flies have been performed under antibiotic (kanamycin) administration in the insect food. The Gfp-
141 labelled strains massively recolonized the fly foregut and midgut (Fig. 2); no auto-fluorescence has been
142 observed in control flies. *G. oxydans* DSF1C.9A successfully colonized the crop, the proventriculus and
143 the first part of the midgut (see the magnifications in Fig. 2b and 2c). The Gfp-labelled cells are clearly
144 restricted to the epithelium side of the proventriculus, embedded in the peritrophic matrix (Fig. 2c).
145 Likely, the midgut showed the same massive colonization pattern as the foregut (Fig. 2d-e). In this tract,
146 small hernias are also visible by interferential contrast (indicated by black arrowheads in Fig. 2e),

147 probably due to microscopic damages produced during the dissection. These hernias appeared full of a
148 gelatinous matrix that resulted Gfp-positive by CLSM, showing that Gfp-labelled cells are completely
149 sunk in the gel and suggesting that the bacterial cells are actually contained by the peritrophic matrix.
150 The black filaments around the organ are the Malpighian tubules, more evident in the CLSM picture
151 (Fig. 2d). Also *A. tropicalis* BYea.1.23(Gfp), and *A. indonesiensis* BTa1.1.44(Gfp) strains successfully
152 colonized the foregut and midgut (Fig. S1): since they showed an identical colonization pattern, only
153 strain BYea.1.23(Gfp) images are shown. The labelled bacteria were present in the whole tract and they
154 have been especially located close to the gut walls and within the peritrophic matrix (Fig. S1).

155

156 **Characterization of *D. suzukii* bacterial diversity by DNA-based analysis.** At first, to have a general
157 view of the bacterial community associated to *D. suzukii*, DNA extracted from 32 specimens has been
158 used, as template, in PCR-DGGE assays (targeting a fragment of the 16S rRNA gene, Tab. S1). In
159 particular, five larvae (n. 1-5), one pupa (n. 6) and ten adults (n. 7-16; Fig. S2a-b) reared on fruits have
160 been analysed, as well as four larvae (n. 29-32), four pupae (n. 25-28) and eight adults (n. 17-24) reared
161 on the artificial diet (Fig. S2c). Consistent with previous data reported for other drosophilid flies
162 (Chandler *et al.*, 2011; Wong *et al.*, 2013), *D. suzukii* specimens showed relatively simple bacterial
163 communities with the presence of few prevalent bacterial taxa. The lowest variability in the community
164 profiles has been observed among larvae reared on fruits and on the artificial diet: many PCR-DGGE
165 bands were conserved among the samples belonging to the same diet. Conversely, only few conserved
166 bands were detected among adults reared on fruits, which showed more complex profiles than larval
167 ones either reared on fruits or on the artificial diet (Fig. S2a-c). PCR-DGGE profiles allowed observing
168 the influence of diet on the insect bacterial community structure and composition (Fig. S2): the bacterial
169 community of adults reared on fruit diet was clearly more complex than the one of adults reared on
170 artificial diet. Moreover, PCR-DGGE sequencing results revealed high prevalence of AAB in insects
171 reared on both diet substrates (Tab. S2).

172 Thus, to sturdily investigate the diet influence on the insect bacterial community, 16S rRNA gene
173 pyrosequencing was performed on 14 specimens, including eight individuals reared on fruits and six on
174 the artificial diet and considering different developmental stages (five larvae, two pupae and seven
175 adults). Variability among the samples has been reported (Tab. S3; Fig. 3a). Using the Shannon Index to
176 measure α -diversity in each sample and plotting it on a rarefaction curve, we confirmed the saturation of
177 the bacterial diversity associated to the samples (Fig. S3). We obtained in total 178,856 reads after
178 quality evaluation and chimera removal. The different ecological estimators showed that, on average,
179 the bacterial communities associated with the specimens reared on fruits exhibited a greater diversity
180 than those from individuals reared on artificial diet (118 ± 42 and 78 ± 24 OTUs, respectively; Tab. S3).
181 As a matter of fact, the microbiota of *D. sukuzii* specimens reared on fruit showed on average a greater
182 richness (Chao1 = 137.4 ± 48.3), a higher diversity ($H' = 2.5 \pm 0.75$) and a higher evenness ($J = 0.52 \pm$
183 0.13), when compared to the microbiota of flies reared on artificial diet (Chao1 = 91.4 ± 31.1 ; $H' = 1.75$
184 ± 0.67 ; $J = 0.4 \pm 0.13$).

185 β -diversity has been evaluated through principal coordinates analysis (PCoA) on the similarity matrix
186 obtained by UniFrac. The two principal components explain 49.67% of the variation (Fig. 3b). PCoA
187 showed three clusters of samples ($p < 0.05$): the first one encompasses the two larvae and the sole pupa
188 reared on the artificial diet; the second one includes all the adults reared on the artificial diet, while the
189 third is constituted by all the specimens reared on fruits (Fig. 3b). Interestingly, the exclusion of AAB
190 OTUs from the analysis showed a loss of the clustering pattern observed before (Fig. 3c). Specifically,
191 the three abovementioned clusters were not significantly different one to each other ($p > 0.05$),
192 highlighting that AAB could be more responsive than other bacterial groups following diet
193 modification. Thus, we evaluated the distribution of AAB at OTU level among the specimens exploring
194 the 16S rRNA gene pyrosequencing dataset: a clustering tendency of the samples in relation to the
195 different diets has been further observed (Fig. 3d).

196 Looking to the bacterial community's composition, the results showed that the average percentage of
197 reads belonging to Acetobacteraceae family was 24.8% per specimen (18% in case of fruit-reared
198 insects and 33.9% for specimens fed with artificial diet; Fig. 3a). At genus level, 16S rRNA gene
199 pyrosequencing revealed that in *D. suzukii* specimens, reared on fruit and on the artificial diet,
200 Acetobacteraceae family was composed mainly by the genera *Acetobacter* and *Gluconobacter* (average
201 20% of 3.9% out of the total reads respectively, Fig. S4; Tab. S4).
202 Interestingly, reads affiliated to Rickettsiales, to which *Wolbachia* genus belongs, have been detected
203 only in flies reared on fruits, with an average of 27.5%, confirming results obtained by PCR-DGGE
204 (Fig. 3a; Fig. S2). *Wolbachia* was the only representative of Rickettsiales order in the dataset. Reads
205 clustering within Rhodospirillales order (the order to which Acetobacteraceae belongs) were present in
206 all the specimens with different abundance; in some cases it reached percentages of 85.2 and 85.4 out of
207 the total number of sequences per sample (MF1 and PP2, respectively). Members of other orders such as
208 Enterobacteriales, Xanthomonadales, Lactobacillales, Rhizobiales, Burkholderiales and
209 Sphingobacteriales constituted relevant fractions of the remaining bacterial communities (Fig. 3a).

210

211 DISCUSSION

212 Prevalence, FISH and 16S rRNA gene PCR-DGGE and pyrosequencing analyses confirmed that AAB
213 are invariably present in *D. suzukii* gut in our experimental conditions. In *D. melanogaster* and other
214 insects, AAB have been demonstrated as prevalent symbionts with important biological roles (Shin *et*
215 *al.*, 2011; Chouaia *et al.*, 2012; Mitraka *et al.*, 2013). For instance, *Acetobacter tropicalis*, a species that
216 we found in *D. suzukii*, was previously described in association with the olive fruit fly *Bactrocera oleae*
217 (Kounatidis *et al.*, 2009).

218 Localization and intimate association of AAB with *D. suzukii*, revealed by FISH (Fig. 1), support the
219 hypothesis that these bacteria may indeed influence the gut functionality. In the midgut, AAB
220 localization along with the peritrophic matrix suggests a bacterial interaction with the host gut

221 epithelium. Moreover, recolonization experiments with Gfp-labelled strains (i.e. *G. oxydans* DSF1C.9A,
222 *A. tropicalis* BYea.1.23 and *A. indonesiensis* BTa1.1.44) strongly supported the capability of AAB to
223 colonize the gut (Fig. 2 and Fig. S1). As indicated elsewhere (Favia *et al.*, 2007), recolonization
224 experiments have been performed under the antibiotic pressure of kanamycin, a required procedure
225 when Gfp cassette is encoded on a plasmid to avoid the loss of the plasmid itself. Certainly, the use of
226 antibiotic could have a negative side effect on the insect host and other gut symbionts. Further
227 investigations could help in verifying if the used concentration of antibiotic might have detrimental
228 effects for the host and/or the gut microbiota. However, such investigation was beyond the purpose of
229 the experiments that were designed to assess which gut portions were recolonized by the strains. For *A.*
230 *tropicalis* a very similar gut localization pattern to that of *D. suzukii* has been already observed in the
231 olive fruit fly *B. oleae* (Kounatidis *et al.*, 2009), where the bacterium was observed in contact with the
232 gut epithelium of the insect, entrapped in a polysaccharidic matrix. Similarly, in other insects, such as
233 the leafhopper *Scaphoideus titanus*, and *Anopheles* and *Aedes* mosquitoes, other AAB of the genus
234 *Asaia* massively colonize the epithelia of the gut and the reproductive organs (Crotti *et al.*, 2009;
235 Damiani *et al.*, 2010; Favia *et al.*, 2007; Gonella *et al.*, 2012). The AAB localization observed in the gut
236 of *D. suzukii* confirmed that guts of sugar-feeding insects are primary habitat for AAB, in which they
237 establish strict topological and presumably functional connections with the epithelial cells (Crotti *et al.*,
238 2010; Chouaia *et al.*, 2014).

239 *D. suzukii* microbiota diversity has been investigated at little extent and just one paper has been
240 published describing the insect bacterial community (Chandler *et al.*, 2014). By the use of a next
241 generation sequencing (NGS) technique, authors analyzed pools of specimens collected from cherries
242 sampled at different developmental stages, showing an high frequency of the gamma-Proteobacterium
243 *Tatumella*, while the two AAB *Gluconobacter* and *Acetobacter* genera were found at lower abundance
244 (Chandler *et al.*, 2014). Conversely, in our study, sequences related to *Tatumella* genus have not been
245 retrieved in any of the analysed samples, but a high prevalence of AAB have been found (average of

246 24.8%). Insects in Chandler and colleagues' work (2014) have been collected in USA, while our
247 populations derive from Italian field-collected individuals. Moreover, different variable regions on 16S
248 rRNA gene have been amplified in the two studies. Such environmental and methodological differences
249 may explain the differences between our and the Chandler *et al.* work (2014). However, further
250 investigations are needed to determine *Tatumella* prevalence in different *D. suzukii* populations,
251 considering with special attention insects collected in different locations, as already mentioned by
252 Chandler *et al.* (2014).

253 It is widely recognized the importance of diet in shaping the insect bacterial community (Montagna *et*
254 *al.*, 2015; Colman *et al.*, 2012; Yun *et al.*, 2014). Particularly, in *D. melanogaster* the establishment and
255 maintenance of the microbiota are determined by bacterial intake from external sources (Blum *et al.*,
256 2013). Differences in the diversity and dominance of bacterial species associated to several *Drosophila*
257 species are thus related to food source (Wong *et al.*, 2011). This has been substantiated by Chandler and
258 coworkers (2011) who observed that individuals of different *Drosophila* species reared on different food
259 sources enriched a similar microbiota when moved to the same medium. With the present study, we
260 confirmed that also in case of *D. suzukii* there are differences in the bacterial communities between
261 animals reared on fruits and on artificial diet (Fig. 3). Specifically, the fruit-based diet determined a
262 higher diversity in the bacterial community rather than the artificial diet, confirming what already
263 reported in literature about the reduction of the insect microbial community complexity in case of
264 artificial diet-fed animals in comparison to natural diet-fed ones (Lehman *et al.*, 2009). In our study, the
265 fruit-based diet can be considered similar to the natural one *D. suzukii* is exposed to in orchards. The
266 diet appeared as a more important factor than the life stage in discriminating the insect associated
267 microbiota, since discrimination at the life stage was possible only between juvenile stages and adults
268 reared on the artificial diet ($p < 0.05$; Fig. 3b). Chandler *et al.* (2011), analyzing clone libraries of the
269 bacterial community associated to different species of *Drosophila* flies, field-collected or reared in the
270 laboratory, found AAB in both types of individuals: sequences related to *Commensalibacter* and

271 *Acetobacter* have been retrieved, while the authors reported the nearly complete lack of *Gluconobacter*
272 sequences and the complete lack of *Gluconacetobacter* ones within their samples. In our 16S rRNA
273 gene-based survey of the *D. suzukii* microbiota, *Acetobacter* and *Gluconobacter* have been detected
274 while *Gluconacetobacter* and *Komagataeibacter* have not, although isolates of these two genera have
275 been obtained. The 16S rRNA sequence phylogenetic proximity of AAB genera and the small region,
276 targeting the bacterial 16S rRNA gene used in our PCR amplifications (about 500 bp), could have
277 masked the discrimination of *Gluconacetobacter* and *Komagataeibacter* sequences (Fig. S4). In this
278 perspective, the use of multiple primer pairs and the choice of longer regions (however taking into
279 account limitations of the current NGS techniques) could lead to a more representative view of the
280 structure of the host bacterial community. Another factor that might have introduced biases in the
281 microbiota analysis is the DNA extraction method. Even though in our work, DNA has been extracted
282 through one of the most widely used, cost-effective and efficient methods available for DNA extraction,
283 i.e. the using sodium dodecyl sulfate-proteinase K-CTAB treatment, the parallel use of alternative
284 methods on the same set of samples might help to better evaluate the reliability of the obtained data.
285 Our results indicated that AAB may play a role in structuring the gut community. In the AAB OTUs
286 distribution in relation to the specimens, a clustering pattern based on the food source was recognized
287 (Fig. 3d), further strengthening the results of the clustering already observed in fig. 3b. Such findings
288 indicate that AAB are primarily involved in the response to the diet, and suggest that they may be
289 directly or indirectly involved in the bacterial community shift following a different diet exposition. We
290 have evaluated the impact of the diet on the bacterial community, without considering the AAB
291 contribution: by excluding AAB OTUs from the analyzed dataset, we found the loss of the previously
292 observed clustering pattern ($p > 0.05$; compare Figs. 3b and 3c). Taken together, these data highlight not
293 only the differentiation of the AAB community in response to the diet type, but also indicate that AAB
294 are crucial in determining samples' grouping along with diet variation. It is also noteworthy that the
295 insects reared on the artificial diet originated from the same field population of the fruit-fed insects.

296 Another variable that could be associated with the distinction of the samples between fruit-fed and
297 artificial diet-fed animals is the presence of *Wolbachia*, but we concluded that it cannot be considered as
298 a driver of the bacterial community modification in this case. Although *Wolbachia* was detected by
299 PCR-DGGE and 16S rRNA barcoding just in fruit-fed samples, the complementary PCR analysis
300 performed for determining *Wolbachia* in the two diet groups, demonstrated its presence in the artificial
301 diet-fed animals. *Wolbachia* is generally considered as intracellular reproductive manipulator, described
302 in many insect species, including different *Drosophila* spp. (Werren *et al.*, 2008; McGraw and O'Neill,
303 2004). The different incidence in samples reared on fruits respect to the artificial diet could be explained
304 by the presence of inhibitory compounds in the artificial diet, hindering or somehow temporarily
305 influencing *Wolbachia* growth. Lack of *Wolbachia* by high throughput sequencing in flies reared on
306 artificial diet could be the result of the number of analyzed insects (n. = 6), since the *Wolbachia*
307 prevalence rate in our *D. suzukii* population has been verified to be 28%. On the other hand, the
308 *Wolbachia* strain associated to *D. suzukii* has been reported to be imperfectly maternally transmitted,
309 showing polymorphic infection (Hamm *et al.*, 2014). Moreover, the results could indicate a
310 diversification of infection rates linked to the diet source; indeed, prevalence analysis pointed out a
311 lower infection rate than previously reported in a similar population (Mazzetto *et al.*, 2015).

312 A competition phenomenon between *Asaia* and *Wolbachia* has been described to occur at the level of
313 mosquito gonads (Rossi *et al.*, 2015) and *Asaia* has been indicated as responsible for inhibiting
314 *Wolbachia* transmission in mosquitoes (Hughes *et al.*, 2014). In this study, we could not observe
315 competition phenomena between AAB and *Wolbachia*. However, no specific investigations have been
316 performed at gonad level. It should be underlined that so far competition has been described only for
317 *Asaia*, a symbiont that has never been described in *D. suzukii* or other *Drosophila* flies.

318 In conclusion, AAB's high prevalence in individuals fed on both diet types, their localization and ability
319 to massively recolonize the insect gut indicate that AAB are major components of the *D. suzukii*
320 microbiota and, similarly to *D. melanogaster*, they might play important roles in the physiology and

321 behaviour of the host. The AAB diversity shifts and their weight in determining the clustering behaviour
322 of the bacterial microbiota in relation to diet might indicate their crucial role in determining the
323 microbiota response to diet in *D. suzukii* gut.

324

325 EXPERIMENTAL PROCEDURES

326 **Insects.** Field-captured larvae of *D. suzukii* emerging from blueberries, raspberries and blackberries in
327 orchards of the Cuneo province, (Piedmont, North-West Italy) in summer 2013 have been reared for at
328 least eight generations in laboratory condition both on fruits (strawberries, blueberries, grapes and kiwi
329 fruits) and on a sugar-based artificial diet (composed with 71 g of corn flour, 10 g of soy flour, 5.6 g of
330 agar, 15 g of sucrose, 17 g of brewer's yeast, 4.7 ml of propionic acid, 2.5 g of vitamins mix for each Kg
331 of the preparation) at the Dipartimento di Scienze Agrarie, Forestali e Alimentari (DISAFA), University
332 of Torino. Insects have been kept in plastic cages (24 × 16 × 12 cm) in a growth chamber at 25 ± 1 °C,
333 65 ± 5% RH and 16L:8D photoperiod, until collected for analyses (Tab. S1). Bacterial community
334 evaluation was carried out on 2nd-3rd instar larvae, pupae, and 7-20 day-old adults.

335

336 **Prevalence of AAB and Wolbachia and AAB isolation.** Prevalences of *Wolbachia* and AAB have been
337 evaluated as described in Method S1. The strategy of isolation was to collect as many AAB isolated
338 colonies as possible according to diversity of colony morphology obtained from different sources (the
339 insect specimens) and different media. A bacterial collection has been obtained and identified as
340 indicated in Method S2. 16S rRNA sequences of representative isolates have been deposited in the ENA
341 database under the accession numbers LN884027-LN884133.

342

343 **Localization of *D. suzukii* AAB by fluorescent in situ hybridization (FISH) and colonization**
344 **experiments with Gfp labelled strains.** FISH has been carried out on tissues and organs dissected from
345 mass-reared *D. suzukii* adults in a sterile saline solution. The dissected organs have been fixed for two

346 minutes at 4°C in 4% paraformaldehyde and washed in Phosphate-Buffered Saline (PBS). All
347 hybridization experiment steps have been performed as previously described (Crotti *et al.*, 2009;
348 Gonella *et al.*, 2012), using fluorescent probes, specifically designed for the acetic acid bacterial group
349 (AAB455, sequence GCGGGTACCGTCATCATCGTCCCCGCT) and for *Gluconobacter* (Go15,
350 sequence AATGCGTCTCAAATGCAGTT and Go18, sequence GTCACGTATCAAATGCAGTTCCC).
351 The universal eubacterial probe, Eub338 (sequence GCTGCCTCCCGTAGGAGT), has been used to
352 detect the localization of the overall bacterial abundance and presence in the organs analysed (Gonella *et*
353 *al.*, 2012). Probes for AAB and Eubacteria have been labelled at the 5' end with the fluorochrome Texas
354 Red (TR; absorption and emission at 595 nm and 620 nm, respectively), whereas probes Go15 and
355 Go18 have been labelled with indodicarbocyanine (Cy5; absorption and emission at 650 nm and 670
356 nm, respectively). After hybridization, the samples have been mounted in anti-fading medium and then
357 observed in a laser scanning confocal microscope SP2- AOBS (Leica). Hybridization experiments in the
358 absence of probes have been performed as negative controls.

359 *G. oxydans* strain DSF1C.9A, *A. tropicalis* BYea.1.23 and *A. indonesiensis* BTa1.1.44 have been
360 transformed through electroporation introducing the plasmid pHM2-Gfp (Favia *et al.*, 2007) as
361 described in **Method S3**. Plasmid stability has been verified for the transformants as reported in **Method**
362 **S4**. Recolonization experiments using *G. oxydans* DSF1C.9A(Gfp), *A. tropicalis* BYea.1.23(Gfp) and *A.*
363 *indonesiensis* BTa1.1.44(Gfp) have been performed as indicated in **Method S5**.

364

365 ***Characterization of the D. suzukii bacterial community through molecular ecology approaches.***

366 Immediately after collection larval, pupal and adult individuals of *D. suzukii* have been washed once
367 with ethanol 70% and twice with saline and immediately stored at -20°C in ethanol until molecular
368 analyses. Total DNA has been individually extracted from larvae, pupae and adults by sodium dodecyl
369 sulfate-proteinase K-cethyltrimethyl ammonium bromide (CTAB) treatment, as described in Raddadi *et*
370 *al.* (2011).

371 PCR-DGGE has been performed as described in Method S6. The obtained sequences have been
372 deposited in the EMBL database under the accession numbers LN884134-LN884176.

373 Genomic DNA previously extracted from designated individuals (codes: LF1, LF2, LF3, PF1, MF1,
374 FF2, FF3, MF4, LP1, LP3, PP2, FP1, FP3, and MP3, Tab. S1, Tab. S3) were used in 16S rRNA gene
375 pyrosequencing as described in Method S7. 16S rRNA gene sequences obtained from 16S rRNA gene
376 pyrosequencing analysis have been deposited in European Nucleotide Archive with accession numbers
377 PRJEB10109. The OTU table obtained from 16S rRNA gene pyrosequencing analysis has been filtered
378 and only OTU sequences of AAB have been kept. Statistical significance ($p < 0.05$) of sample
379 distribution in different clusters along Axis 1 of PCoA analysis has been examined by t-test using the
380 software GraphPad Prism version 5.03. Heatmap based on the distribution of AAB OTUs has been
381 prepared as described in Method S8.

382

383 FUNDING INFORMATION

384 King Abdullah University of Science and Technology supported the study through the baseline research
385 funds to D.D. This work was partially funded by Consorzio di Ricerca Sperimentazione e Divulgazione
386 per l'Ortofrutticoltura Piemontese, within the project "Programma di ricerca, sperimentazione e
387 dimostrazione agricola in frutticoltura e orticoltura – 2014 – Indagini sul nuovo dittero esotico
388 *Drosophila suzukii* responsabile di gravi danni alle drupacee". E.C. acknowledges personal support from
389 "Piano Sviluppo di Ateneo: Linea B-Dotazione annuale per attività istituzionale" in the project "Acetic
390 acid bacteria cell factories".

391

392 REFERENCES

393 Bextine, B., Lauzon, C., Potter, S., Lampe, D., and Miller, T.A. (2004) Establishment of a genetically
394 marked insect-derived symbiont in multiple host plants. *Curr Microbiol* 48: 327-331.

- 395 Blum, J.E., Fischer, C.N., Miles, J., and Handelsman, J. (2013) Frequent replenishment sustains the
396 beneficial microbiome of *Drosophila melanogaster*. mBio 4. e00860-13.
- 397 Brummel, T., Ching, A., Seroude, L., Simon, A.F., and Benzer, S. (2004) *Drosophila* lifespan
398 enhancement by exogenous bacteria. Proc Natl Acad Sci U S A 101:12974-9.
- 399 Cattel, J., Kaur, R., Gibert, P., Martinez, J., Fraimout, A., Jiggins, F., Andrieux, T., Siozios, S., Anfora,
400 G., Miller, W., Rota-Stabelli, O., Mouton, L. (2016) *Wolbachia* in European populations of the
401 invasive pest *Drosophila suzukii*: regional variation in infection frequencies. PLoS ONE 11:
402 e0147766.
- 403 Chandler, J.A., James, P.M., Jospin, G., and Lang, J.M. (2014) The bacterial communities of
404 *Drosophila suzukii* collected from undamaged cherries. PeerJ 2: e474.
- 405 Chandler, J.A., Lang, J.M., Bhatnagar, S., Eisen, J.A., and Kopp, A. (2011) Bacterial communities of
406 diverse *Drosophila* species: Ecological context of a host-microbe model system. PLoS Genet 7(9):
407 e1002272.
- 408 Chouaia, B., Gaiarsa, S., Crotti, E., Comandatore, F., Degli Esposti, M., Ricci, I., *et al.* (2014) Acetic
409 acid bacteria genomes reveal functional traits for adaptation to life in insect guts. Genome Biol Evol
410 6(4): 912–920.
- 411 Chouaia, B., Rossi, P., Epis, S., Mosca, M., Ricci, I., Damiani C., *et al.* (2012). Delayed larval
412 development in *Anopheles mosquitoes* deprived of *Asaia* bacterial symbionts. BMC Microbiol 12:
413 S2.
- 414 Cini, A., Ioriatti C., and Anfora G. (2012) A review of the investigation of *Drosophila suzukii* in Europe
415 and draft research agenda for integrated pest management. Bull Insectol 65: 149-160.
- 416 Colman, D.R., Toolson, E.C., Takacs-Vesbach, C.D. (2012) Do diet and taxonomy influence insect gut
417 bacterial communities? Mol Ecol 21:5124-37.

- 418 Corby-Harris, V., Habel, K.E., Ali, F.G., and Promislow, D.E.L. (2007) Alternative measures of
419 response to *Pseudomonas aeruginosa* infection in *Drosophila melanogaster*. J Evol Biol 20(2):526-
420 33.
- 421 Cox, C., and Gilmore, M. (2007) Native microbial colonization of *Drosophila melanogaster* and its use
422 as a model of *Enterococcus faecalis* pathogenesis. Infect Immun 75:1565-1576.
- 423 Crotti, E., Damiani, C., Pajoro, M., Gonella, E., Rizzi, A., Ricci, I., *et al.* (2009) *Asaia*, a versatile acetic
424 acid bacterial symbiont, capable of cross-colonizing insects of phylogenetically-distant genera and
425 orders. Environ Microbiol 11: 3252-3264.
- 426 Crotti, E., Rizzi, A., Chouaia, B., Ricci, I., Favia, G., Alma, A., *et al.* (2010) Acetic acid bacteria, newly
427 emerging symbionts of insects. Appl Environ Microbiol 76: 6963-6970.
- 428 Damiani, C., Ricci, I., Crotti, E., Rossi, P., Rizzi, A., Scuppa, P., *et al.* (2010) Mosquito-bacteria
429 symbiosis: the case of *Anopheles gambiae* and *Asaia*. Microb Ecol 60 (3): 644-654.
- 430 Eriksson, A., Anfora, G., Lucchi, A., Lanzo, F., Virant-Doberlet, M., and Mazzoni, V. (2012)
431 Exploitation of insect vibrational signals reveals a new method of pest management. PLoS One 7:
432 e32954.
- 433 Erkosar, B., Storelli, G., Defaye, A., and Leulier F. (2013) Host-intestinal microbiota mutualism:
434 "learning on the fly". Cell Host Microbe 13: 8-14.
- 435 Favia, G., Ricci, I., Damiani, C., Raddadi, N., Crotti, E., Marzorati, M., *et al.* (2007) Bacteria of the
436 genus *Asaia* stably associate with *Anopheles stephensi*, an Asian malarial mosquito vector. Proc Natl
437 Acad Sci USA 104: 9047–9051.
- 438 Gonella, E., Crotti, E., Rizzi, A., Mandrioli, M., Favia, G., Daffonchio, D., and Alma, A. (2012)
439 Horizontal transmission of the symbiotic bacterium *Asaia* sp. in the leafhopper *Scaphoideus titanus*
440 Ball (Hemiptera: Cicadellidae). BMC Microbiol 12(Suppl 1): S4.

- 441 Goodhue, R.E., Bolda, M., Farnsworth, D., Williams, J.C., and Zalom F.G. (2011) Spotted wing
442 drosophila infestation of California strawberries and raspberries: economic analysis of potential
443 revenue losses and control costs. *Pest Manag Sci* 67: 1396-1402.
- 444 Hamdi, C., Balloi, A., Essanaa, J., Crotti, E., Gonella, E., Raddadi, N., Ricci, I., Boudabous, A., Borin,
445 S., Manino, A., Bandi, C., Alma, A., Daffonchio, D., Cherif, A., 2011. Gut microbiome dysbiosis and
446 honeybee health. *J Appl Entomol* 135: 524–533.
- 447 Hamm, C.A., Begun, D.J., Vo, A., Smith, C.C.R., Saelao, P., Shaver, A.O., Jaenike, J., and Turelli, M.
448 (2014) *Wolbachia* do not live by reproductive manipulation alone: infection polymorphism in
449 *Drosophila suzukii* and *D. subpulchrella*. *Mol Ecol* 23: 4871–4885.
- 450 Hauser, M. (2011) A historic account of the invasion of *Drosophila suzukii* (Matsumura) (Diptera:
451 Drosophilidae) in the continental United States, with remarks on their identification. *Pest Manag Sci*.
452 67: 1352–1357.
- 453 Hughes, G.L., Dodson, B.L., Johnson, R.M., Murdock, C.C., Tsujimoto, H., Suzuki, Y., Patt, A.A., Cui,
454 L., Nossa, C.W., Barry, R.M., Sakamoto, J.M., Hornett, E.A., Rasgon, J.L. (2014) Native
455 microbiome impedes vertical transmission of *Wolbachia* in *Anopheles* mosquitoes. *Proc Natl Acad*
456 *Sci U S A* 111:12498-503.
- 457 Kounatidis, I., Crotti, E., Sapountzis, P., Sacchi, L., Rizzi, A., Chouaia, B., *et al.* (2009) *Acetobacter*
458 *tropicalis* is a major symbiont of the olive fruit fly (*Bactrocera oleae*). *Appl Environ Microbiol* 75:
459 3281-328.
- 460 Lee, J.C., Bruck, D.J., Deves, A.J., Ioriatti, C., Vogt, H., and Baufeld, P. (2011) In Focus: Spotted wing
461 drosophila, *Drosophila suzukii*, across perspectives. *Pest Manag Sci* 67: 1349-1351.
- 462 Lee, K.-A., and Lee, W.-J. (2014) *Drosophila* as a model for intestinal dysbiosis and chronic
463 inflammatory diseases. *Dev Comp Immunol* 42(1):102-10.

- 464 Lehman, R.M., Lundgren, J.G., and Petzke, L.M. (2009) Bacterial communities associated with the
465 digestive tract of the predatory ground beetle, *Poecilus chalcites*, and their modification by laboratory
466 rearing and antibiotic treatment. *Microb Ecol* 57:349-358.
- 467 Mazzetto, F., Gonella, E., and Alma, A. (2015) *Wolbachia* infection affects female fecundity in
468 *Drosophila suzukii*. *Bull Insectol* 68: 153-157.
- 469 McGraw, E.A., O'Neill S. (2004) *Wolbachia pipientis*: intracellular infection and pathogenesis in
470 *Drosophila*. *Curr Opin Microbiol* 7:67-70.
- 471 Mitraka, E., Stathopoulos, S., and Siden-Kiamos, I. (2013) *Asaia* accelerates larval development of
472 *Anopheles gambiae*. *Pathog Glob Health* 107: 305-311.8
- 473 Mitsui, H., Takahashi, H.K., and Kimura, M.T. (2006) Spatial distributions and clutch sizes of
474 *Drosophila* species ovipositioning on cherry fruits of different stages. *Popul Ecol* 48: 233-237.
- 475 Montagna, M., Chouaia, B., Mazza, G., Prosdocimi, E.M., Crotti, E., Mereghetti, V., *et al.* (2015)
476 Effects of the diet on the microbiota of the red palm weevil (Coleoptera: Dryophthoridae). *PLoS One*
477 10: e0117439.
- 478 Raddadi, N., Gonella, E., Camerota, C., Pizzinat, A., Tedeschi, R., Crotti, E., *et al.* (2011) “*Candidatus*
479 *Liberibacter europaeus*” sp. nov. that is associated with and transmitted by the psyllid *Cacopsylla*
480 *pyri* apparently behaves as an endophyte rather than a pathogen. *Environ Microbiol* 13: 414–426.
- 481 Ren, C., Webster, P., Finkel, S.E., and Tower, J. (2007) Increased internal and external bacterial load
482 during *Drosophila* aging without life-span trade-off. *Cell Metab* 6:144–152.
- 483 Ridley, E.V., Wong, A.C.N., Westmiller, S., and Douglas, A.E. (2012) Impact of the resident microbiota
484 on the nutritional phenotype of *Drosophila melanogaster*. *PLoS One* 7: e36765.
- 485 Rossi, P., Ricci, I., Cappelli, A., Damiani, C., Ulissi, U., Mancini, M.V., Valzano, M., Capone, A., Epis,
486 S., Crotti, E., Chouaia, B., Scuppa, P., Joshi, D., Xi, Z., Mandrioli, M., Sacchi, L., O'Neill, S.L.,
487 Favia, G. (2015) Mutual exclusion of *Asaia* and *Wolbachia* in the reproductive organs of mosquito
488 vectors. *Parasit Vectors* 8:278.

- 489 Ryu, J.H., Kim, S.H., Lee, H.Y., Bai, J.Y., Nam, Y.D., Bae, J.W., *et al.* (2008) Innate immune
490 homeostasis by the homeobox gene *Caudal* and commensal-gut mutualism in *Drosophila*. *Science*
491 319: 777–782.
- 492 Siozios, S., Cestaro, A., Kaur, R., Pertot, I., Rota-Stabelli, O., and Anfora, G. (2013) Draft genome
493 sequence of the *Wolbachia* endosymbiont of *Drosophila suzukii*. *Genome Announc.* 1(1), pii.00032-
494 13.
- 495 Sharon, G., Segal, D., Ringo, J.M., Hefetz, A., Zilber-Rosenberg, I., and Rosenberg, E. (2010)
496 Commensal bacteria play a role in mating preference of *Drosophila melanogaster*. *Proc Natl Acad*
497 *Sci USA* 107: 20051-20056.
- 498 Shin, S.C., Kim, S.H., You, H., Kim, B., Kim, A.C., Lee, K.A., *et al.* (2011) *Drosophila* microbiome
499 modulates host developmental and metabolic homeostasis via insulin signaling. *Science* 334: 670-
500 674.
- 501 Storelli, G., Defaye, A., Erkosar, B., Hols, P., Royet, J., and Leulier, F. (2011) *Lactobacillus plantarum*
502 promotes *Drosophila* systemic growth by modulating hormonal signals through TOR_dependent
503 nutrient sensing. *Cell Metab* 14: 403-414.
- 504 Walsh, D.B., Bolda, M.P., Goodhue, R.E., Dreves, A.J., Lee, J.C., Bruck, D.J., Walton, V.M., O'Neals,
505 S.D., and Zalom, F.G. (2011) *Drosophila suzukii* (Diptera: Drosophilidae): Invasive pest of ripening
506 soft fruit expanding its geographic range and damage potential. *Int J Pest Manage* 1: 1-7.
- 507 Werren, J.H., Baldo, L., Clark, M.E. (2008) *Wolbachia*: master manipulators of invertebrate biology.
508 *Nat Rev Microbiol* 6:741-51.
- 509 Wong, A., Chaston, J., and Douglas, A. (2013) The inconstant gut microbiota of *Drosophila* species
510 revealed by 16S rRNA gene analysis. *ISME J* 7: 1922-1932.
- 511 Wong, A., Ng, P., and Douglas, A. (2011) Low diversity bacterial community in the gut of the fruitfly
512 *Drosophila melanogaster*. *Environ Microbiol* 13: 1889-1900.

- 513 Yamada, Y., Pattaraporn, Y., Vu, H.T.L., Muramatsu, Y., Ochaikul, D., and Nakagawa, Y. (2012b)
514 Subdivision of the genus *Gluconacetobacter* Yamada, Hoshino and Ishikawa 1998: the proposal of
515 *Komagatabacter* gen. nov., for strains accommodated to the *Gluconacetobacter xylinus* group in the
516 α -Proteobacteria. *Ann Microbiol* 62: 849–859.
- 517 Yamada, Y., Yukphan, P., Lan, Vu, H.T., Muramatsu, Y., Ochaikul, D., Tanasupawat, S., and
518 Nakagawa, Y. (2012a) Description of *Komagataeibacter* gen. nov., with proposals of new
519 combinations (Acetobacteraceae). *J Gen Appl Microbiol* 58: 397-404.
- 520 Yun, J.H., Roh, S.W., Whon, T.W., Jung, M.J., Kim, M.S., Park, D.S., Yoon, C., Nam, Y.D., Kim, Y.J.,
521 Choi, J.H., Kim, J.Y., Shin, N.R., Kim, S.H., Lee, W.J., Bae, J.W. (2014) Insect gut bacterial
522 diversity determined by environmental habitat, diet, developmental stage, and phylogeny of host.
523 *Appl Environ Microbiol* 80:5254-64.

524

525 **TABLE**

526 **Table 1.** Identification of cultivable bacteria associated to *D. suzukii*. All the isolates showed a
 527 percentage of identity >97% in relation to the indicated species.
 528

Isolates	No. isolates	LP	PP	AP fly	AF fly
<i>Acetobacter tropicalis</i>	1	0	0	0	1
<i>Acetobacter orleanensis/malorum/cerevisiae</i>	4	0	0	0	4
<i>Acetobacter peroxydans</i>	1	0	0	0	1
<i>Acetobacter indonesiensis</i>	10	0	1	1	8
<i>Acetobacter persici</i>	10	0	1	1	8
<i>Acetobacter orientalis</i>	1	0	0	0	1
<i>Acetobacter cibirongensis</i>	1	0	0	0	1
<i>Gluconacetobacter liquefaciens</i>	4	1	3	0	0
<i>Komagataeibacter</i> sp	118	0	0	0	118
<i>Gluconacetobacter/Komagataeibacter</i> sp.	1	0	0	0	1
<i>Gluconobacter kondonii</i>	1	0	0	0	1
<i>Gluconobacter oxydans</i>	12	0	0	0	12
<i>Gluconobacter kanchanaburiensis</i>	5	3	1	1	0
<i>Pseudomonas geniculata</i>	1	0	0	1	0
<i>Serratia</i> sp.	8	2	6	0	0
<i>Micrococcus</i> sp.	5	0	0	0	5
<i>Microbacterium foliorum</i>	2	0	0	0	2
<i>Streptococcus salivarius</i>	1	0	0	1	0
<i>Staphylococcus</i> sp.	12	0	0	0	12
<i>Paenibacillus</i> sp.	2	0	0	0	2
<i>Lactococcus lactis</i>	1	0	0	0	1
<i>Lactobacillus plantarum</i>	1	0	1	0	0
Total	202	6	13	5	178

529 LP: larvae fed with artificial diet; PP: pupae fed with artificial diet; AP: Adults fed with artificial diet; AF: Adults fed with
 530 fruit diet
 531

532

533 FIGURES

534 **Figure 1.** AAB localization in the gut of *D. suzukii*. (a-d) FISH of the insect gut after hybridization with
 535 the Texas red-labelled probe AAB455, matching AAB. (a) Superposition of the interferential contrast
 536 (c) and the FISH (b) pictures of the midgut close to the proventriculus that is indicated by white arrows
 537 [for a scheme of the morphology of the initial part of the midgut and the upstream region refer to panel
 538 (a) of Figure 3]. (d) Magnification of the image in (b). The massive presence of AAB adherent to the
 539 peritrophic matrix (the black line below the first layer of cells indicated by black arrows) is observed.
 540 (e-h) FISH of posterior midgut with the Texas red-labelled universal eubacterial probe Eub338 (f) and
 541 the Cy5-labelled probe specific for *Gluconobacter*, Go615 and Go618 (g). (e) Intestine portion pictured
 542 by interferential contrast. (h) Superposition of hybridization signals of Eubacteria (red) and
 543 *Gluconobacter* (blue). Bars = 50 μm .

544

545 **Figure 2.** Colonization of *D. suzukii* foregut and midgut by Gfp-labelled *G. oxydans* DSF1C.9A1
 546 documented by confocal laser scanning microscopy. (a) The scheme represents the first tract of the
 547 digestive system and shows the different gut portions highlighted in the next panels. (b-d) Digestive
 548 tract portions including the crop, the proventriculus and the first part of the midgut. (c, d) Magnified
 549 views of the crop (c) and the proventriculus (d) showed in (b). Masses of fluorescent cells are observed
 550 in the crop (arrows). When the fluorescent strain cells reach the proventriculus (d), they colonize the gut
 551 part close to peritrophic matrix. (e-f) Interferential contrast (f) and confocal laser scanning (e) pictures
 552 of the posterior midgut of *D. suzukii* massively colonized by the *G. oxydans* strain labelled with Gfp.
 553 Small hernias (arrowhead) are shown. In some cases, the gelatinous matrix in the hernias present
 554 fluorescent cells. Bars = 50 μm .

555

556 **Figure 3.** Bacterial diversity associated with *D. suzukii* by 16S rRNA gene pyrosequencing. (a) 16S
 557 RNA gene pyrosequencing describing bacterial communities, at order level, associated with *D. suzukii*.

558 Names, under histograms, refer to fly specimens; in columns, the relative abundances in percentages of
559 the identified orders are showed. Sequences that did not match with anything in the database are
560 indicated as “Unclassified sequences”; bacterial sequences that have not been assigned to any
561 taxonomical group are indicated as “Bacteria_unclassified”; bacterial orders under 3% representation
562 per sample have been grouped and indicated as “Class. Bac. Orders under 3%”. (b) Principal coordinate
563 analysis (PCoA) on the phylogenetic β -diversity matrix on *D. suzukii* samples, considering all the
564 bacterial OTUs. (c) Principal coordinate analysis (PCoA) on the phylogenetic β -diversity matrix on *D.*
565 *suzukii* samples, considering all the bacterial OTUs, except for the ones belonging to AAB group. Red
566 circle indicates fruit-fed individuals, while blue circles mark specimens fed on the artificial diet. (d)
567 Distribution of AAB in *D. suzukii* hosts. The relative abundance of AAB OTUs, determined at 97%
568 identity, is showed in the heatmap. Coloured scale represents OTUs abundance for each sample
569 (indicated on the vertical axis). In bold are indicated samples from fruit-rearing; the remaining samples
570 are related to artificial diet-fed animals. First letter of codes refers to the fly stage (M: male adult; F:
571 female adult; L: larva; P: pupa); second letter of codes refers to feeding system (F: fruit-based diet; P:
572 artificial diet); third letter of codes is related to subsequent number of samples.

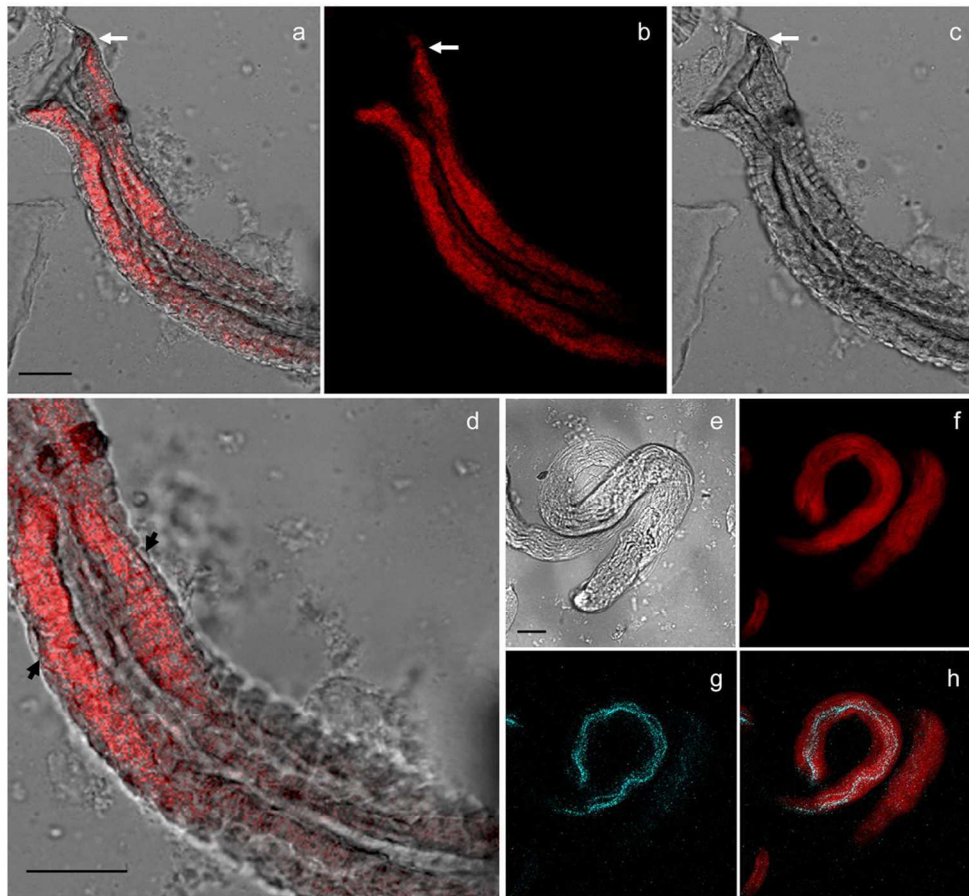


Figure 1. AAB localization in the gut of *D. sukii*. (a-d) FISH of the insect gut after hybridization with the Texas red-labelled probe AAB455, matching AAB. (a) Superposition of the interferential contrast (c) and the FISH (b) pictures of the midgut close to the proventriculus that is indicated by white arrows [for a scheme of the morphology of the initial part of the midgut and the upstream region refer to panel (a) of Figure 3]. (d) Magnification of the image in (b). The massive presence of AAB adherent to the peritrophic matrix (the black line below the first layer of cells indicated by black arrows) is observed. (e-h) FISH of posterior midgut with the Texas red-labelled universal eubacterial probe Eub338 (f) and the Cy5-labelled probe specific for *Gluconobacter*, Go615 and Go618 (g). (e) Intestine portion pictured by interferential contrast. (h) Superposition of hybridization signals of Eubacteria (red) and *Gluconobacter* (blue). Bars = 50 μm .

205x189mm (300 x 300 DPI)

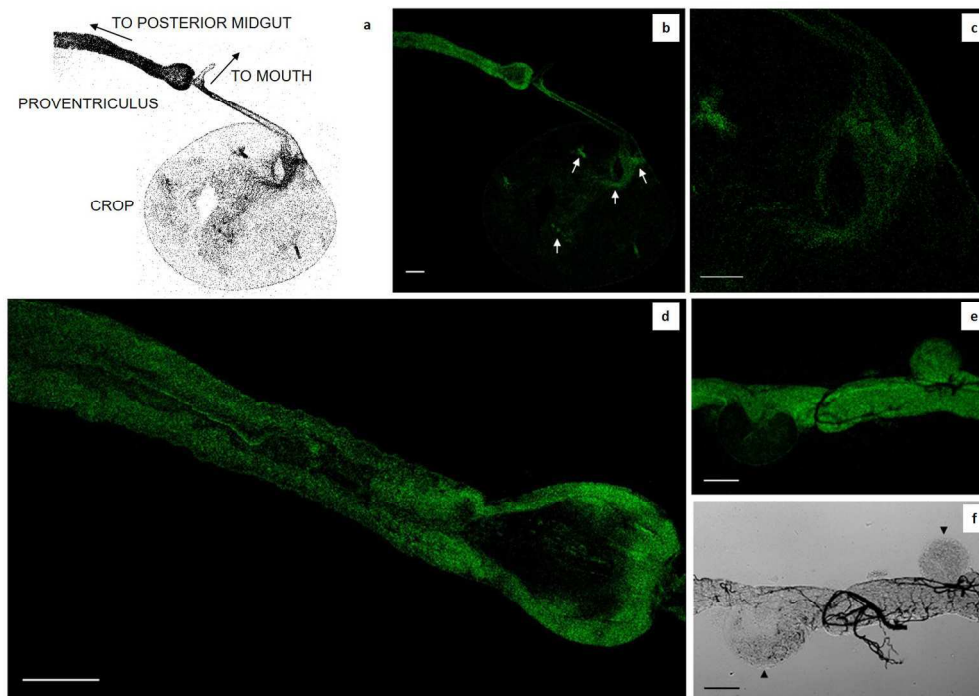


Figure 2. Colonization of *D. sukukii* foregut and midgut by Gfp-labelled *G. oxydans* DSF1C.9A1 documented by confocal laser scanning microscopy. (a) The scheme represents the first tract of the digestive system and shows the different gut portions highlighted in the next panels. (b-d) Digestive tract portions including the crop, the proventriculus and the first part of the midgut. (c, d) Magnified views of the crop (c) and the proventriculus (d) showed in (b). Masses of fluorescent cells are observed in the crop (arrows). When the fluorescent strain cells reach the proventriculus (d), they colonize the gut part close to peritrophic matrix. (e-f) Interferential contrast (f) and confocal laser scanning (e) pictures of the posterior midgut of *D. sukukii* massively colonized by the *G. oxydans* strain labelled with Gfp. Small hernias (arrowhead) are shown. In some cases, the gelatinous matrix in the hernias present fluorescent cells. Bars = 50 μ m.

277x194mm (300 x 300 DPI)

Only

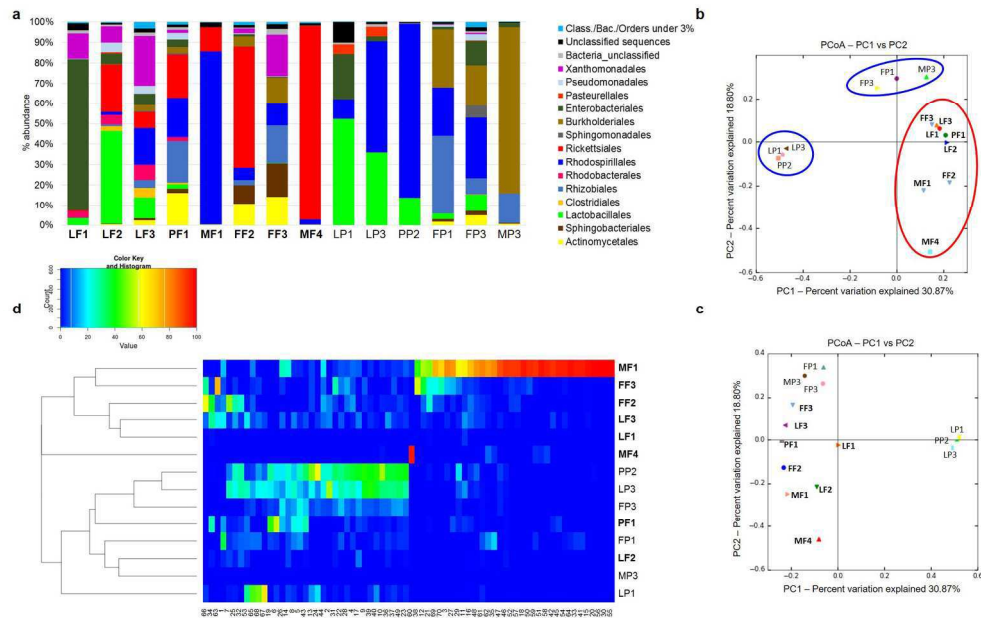


Figure 3. Bacterial diversity associated with *D. sukuzii* by 16S rRNA gene pyrosequencing. (a) 16S RNA gene pyrosequencing describing bacterial communities, at order level, associated with *D. sukuzii*. Names, under histograms, refer to fly specimens; in columns, the relative abundances in percentages of the identified orders are showed. Sequences that did not match with anything in the database are indicated as "Unclassified sequences"; bacterial sequences that have not been assigned to any taxonomical group are indicated as "Bacteria_unclassified"; bacterial orders under 3% representation per sample have been grouped and indicated as "Class. Bac. Orders under 3%". (b) Principal coordinate analysis (PCoA) on the phylogenetic β -diversity matrix on *D. sukuzii* samples, considering all the bacterial OTUs. (c) Principal coordinate analysis (PCoA) on the phylogenetic β -diversity matrix on *D. sukuzii* samples, considering all the bacterial OTUs, except for the ones belonging to AAB group. Red circle indicates fruit-fed individuals, while blue circles mark specimens fed on the artificial diet. (d) Distribution of AAB in *D. sukuzii* hosts. The relative abundance of AAB OTUs, determined at 97% identity, is showed in the heatmap. Coloured scale represents OTUs abundance for each sample (indicated on the vertical axis). In bold are indicated samples from fruit-rearing; the remaining samples are related to artificial diet-fed animals. First letter of codes refers to the fly stage (M: male adult; F: female adult; L: larva; P: pupa); second letter of codes refers to feeding system (F: fruit-based diet; P: artificial diet); third letter of codes is related to subsequent number of samples.

372x231mm (150 x 150 DPI)

