

1 **Title:** Comprehensive characterization of internal and cuticle surface microbiota of laboratory-
2 reared F₁ *Anopheles albimanus* originating from different sites

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22 **Abstract**

23 **Background:** Research on mosquito-microbe interactions may lead to new tools for mosquito
24 and mosquito-borne disease control. To date, such research has largely utilized laboratory-
25 reared mosquitoes that typically lack the microbial diversity of wild populations. A logical
26 progression in this area involves working under controlled settings using field-collected
27 mosquitoes or, in most cases, their progeny. Thus, an understanding of how laboratory
28 colonization affects the assemblage of mosquito microbiota would aid in advancing mosquito
29 microbiome studies and their applications beyond laboratory settings.

30 **Methods:** Using high throughput 16S rRNA amplicon sequencing, we characterized the internal
31 and cuticle surface microbiota of F₁ progeny of wild-caught adult *Anopheles albimanus* from four
32 locations in Guatemala. A total of 132 late instar larvae and 135 2-5day old, non-blood-fed virgin
33 adult females that were reared under identical laboratory conditions, were pooled (3
34 individuals/pool) and analyzed.

35 **Results:** Results showed geographical heterogeneity in both F₁ larval internal ($p=0.001$; pseudo-
36 $F = 9.53$) and cuticle surface ($p=0.001$; pseudo- $F = 8.51$) microbiota, and only F₁ adult cuticle
37 surface ($p=0.001$; pseudo- $F = 4.5$) microbiota, with a more homogenous adult internal microbiota
38 ($p=0.12$; pseudo- $F = 1.6$) across collection sites. Overall, ASVs assigned to *Leucobacter*,
39 *Thorsellia*, *Chryseobacterium* and uncharacterized *Enterobacteriaceae*, dominated F₁ larval
40 internal microbiota, while *Acidovorax*, *Paucibacter*, and uncharacterized *Comamonadaceae*,
41 dominated the larval cuticle surface. F₁ adults comprised a less diverse microbiota compared to
42 larvae, with ASVs assigned to the genus *Asaia* dominating both internal and cuticle surface
43 microbiota, and constituting at least 70% of taxa in each microbial niche.

44 **Conclusions:** These results suggest that location-specific heterogeneity in field mosquito
45 microbiota can be transferred to F₁ progeny under normal laboratory conditions, but this may not
46 last beyond the F₁ larval stage without adjustments to maintain field-derived microbiota. Our
47 findings provide the first comprehensive characterization of laboratory-colonized F₁ *An.*
48 *albimanus* progeny from field-derived mothers. This provides a background for studying how
49 parentage and environmental conditions differentially or concomitantly affect mosquito
50 microbiome composition, and how this can be exploited in advancing mosquito microbiome
51 studies and their applications beyond laboratory settings.

52 **Keywords:** Mosquito microbiota, *Anopheles albimanus*, laboratory colonization, mosquito
53 microbiome, next generation sequencing, 16S rRNA gene amplicon sequencing

54 **Background**

55 Mosquitoes contain microbes that inhabit various tissues, such as the alimentary canal,
56 reproductive organs, and cuticle surface [1]. These microbes are thought to be principally
57 obtained from the mosquito habitat during larval development, and from food sources at the
58 adult stage [1]. In addition to acquisition from larval habitats and/or adult food sources,
59 transovarial bacterial transmission from adult females to their eggs, and transstadial
60 transmission across different immature stages, and into the adult stage, have been
61 demonstrated [2, 3]. As a key component of the mosquito microbiota, environmentally-acquired
62 microbes have been shown to affect mosquito life history traits such as the rate of pupation and
63 adult body size [4]. The mosquito microbiota has also been shown to affect the following aspects
64 of mosquito biology: immunity to human pathogens [5], reproduction [6], insecticide resistance
65 [7, 8], and ultimately vector competence—the mosquito’s ability to acquire, maintain and transmit
66 pathogens [5]. These effects of the microbiota on mosquito biology are being leveraged to
67 develop novel approaches for mosquito-borne disease control [9].

68 The use of next generation molecular biology tools has resulted in extensive characterization of
69 mosquito microbiota, with the initial focus on bacterial and archaeal components now expanding
70 to eukaryotic microbes [10, 11] and viruses [12, 13]. These advances in mosquito microbiota
71 research have led to field applications of mosquito symbionts for mosquito control. For *Aedes*
72 *aegypti*, the principal vector of dengue, Zika, chikungunya and yellow fever viruses, mosquito-
73 derived symbionts are now being used to suppress mosquito populations [14] and also being
74 considered to control the spread of pathogens [15]. However, studies exploring mosquito
75 symbionts for malaria control have largely remained at the laboratory stage [16, 17]. Similarly,
76 the microbiota of mosquito vectors in some geographical regions are well characterized and
77 studied compared to those from other regions. In malaria vectors for example, studies on the

78 microbiota have largely focused on Sub-Saharan African species—in particular, *Anopheles*
79 *gambiae*—and to a lesser extent on those from Southeast Asia [18]. In contrast, the microbiota
80 of Latin American malaria vectors have only recently been comprehensively characterized [7, 8,
81 19-21], with these studies describing associations between *An. albimanus* microbiota and
82 insecticide resistance [7, 8], and the factors that shape the composition of *An. darlingi*, *An.*
83 *albimanus*, *An. nunezovari*, *An. rangeli*, and *An. triannulatus* microbiota [19-21].

84 To exploit the mosquito microbiota for malaria and malaria vector control, research must
85 successfully advance from laboratory to field settings, a transition which can be fraught with
86 challenges. For example, some malaria vectors such as *An. darlingi*, *An. vestitipennis*, and *An.*
87 *gambiae* breed in sites that are small, temporary and often difficult to find and/or access [22-26],
88 making it hard to obtain sufficient immature field mosquitoes for experiments. Where larval
89 habitats are plentiful and easy to find and/or access, the subsequent rearing of field-collected
90 mosquitoes to obtain uniform characteristics can pose additional challenges [27, 28].

91 Additionally, some malaria vectors belong to species complexes whose members are
92 morphologically indistinguishable [29-32], constituting another layer of complexity that needs to
93 be considered in elucidating mosquito-microbe interactions in malaria vectors.

94 These challenges, which are common to research on mosquito ecology and control, are often
95 not reported or discussed in mosquito microbiome studies. Several failed attempts at collecting
96 and rearing sufficient immature mosquitoes from the field for our previous study on the role of
97 mosquito microbiota in insecticide resistance resulted in ultimately using either wild-caught
98 adults [7] or F₁ progeny derived from field-collected adult mosquitoes [8, 33]. While field-caught
99 adult mosquitoes or their F₁ progeny may offer insights into mosquito-microbe interactions in
100 field scenarios, obtaining adult field-collected mosquitoes with uniform and/or controlled
101 physiological characteristics is usually not feasible. Although geographically associated

102 heterogeneity in microbiota of field-collected mosquitoes has been previously described [34, 35],
103 there is limited information on the fate of field-acquired microbiota after laboratory colonization of
104 field-collected mosquitoes. So far, it has been observed that upon eclosion, newly-emerged
105 laboratory-reared adult mosquitoes show a reduction in bacterial diversity in contrast to earlier
106 developmental stages [36]. In addition, a recent study on the fate of field-acquired microbiota in
107 laboratory-colonized *An. gambiae s.l.* showed a reduction in bacterial diversity of the F₅ progeny
108 (the first point of measurement) that were reared in dechlorinated tap water in contrast to F₀ [37].
109 Another study of *Ae. aegypti* from different geographical locations showed no associations
110 between geographical location and microbiota composition after several generations of
111 laboratory colonization [38]. At what point the microbiota of laboratory-colonized mosquitoes
112 become homogenous and whether the microbiota of F₁ laboratory progeny represent their
113 parental origin—and thus could be used for symbiont-based translational studies—remains
114 largely undescribed.

115 Here, we present a comprehensive characterization of the microbiota of laboratory-reared F₁
116 progeny from field-caught adult *An. albimanus*. These represent unanticipated findings from a
117 larger study that was aimed at characterizing the effects of insecticide exposure and resistance
118 on *An. albimanus* microbiota [8]. We expected little or no immediate loss of geographical
119 heterogeneity in microbial composition upon initial laboratory colonization based on evidence of
120 this type of heterogeneity in field-derived populations [34, 35]. However, our data showed
121 geographical heterogeneity in both F₁ larval internal and cuticle surface microbiota, and only F₁
122 adult cuticle surface microbiota, with a more homogenous adult internal microbiota. These
123 findings lay the foundations for studying how parentage and environmental conditions
124 differentially or concomitantly affect mosquito microbiome composition, and how this can be

125 exploited in advancing mosquito microbiome studies and their applications beyond laboratory
126 settings.

127 **Results**

128 Using high throughput deep sequencing of the universal bacterial and archaeal 16S rRNA gene,
129 we characterized the microbiota of the internal and cuticle surface microbial niches of laboratory
130 reared *An. albimanus* F₁ larvae (n=132) and adult (n=135) progeny originating from mothers that
131 were collected from four locations in south central Guatemala (Fig 1). Mosquitoes were
132 processed as pools (3 individuals per pool), resulting in a total of 44 larval and 45 adult pools. F₁
133 larvae were from El Tererro, Las Cruces 3 and Las Cruces 4, while F₁ adults were from Las
134 Cruces 1, Las Cruces 3 and Las Cruces 4 (Table 1). Following quality control of the resulting
135 sequencing reads, amplicon sequence variants (ASVs) were identified and used in downstream
136 analysis. Since adult mosquito microbiota is distinct from that of immature stages [39, 40], ASVs
137 from larvae and adults were analyzed separately, as were ASVs from each microbial niche
138 (internal or cuticle surface).

139

140 **Internal and cuticle surface microbiota of laboratory colonized F₁ *An. albimanus* larvae** 141 **differed by geographic origin**

142 Non-pairwise Bray-Curtis distance comparison showed significant differences in internal
143 ($p=0.001$) and cuticle surface ($p=0.001$) microbiota between F₁ larvae from different collection
144 sites. Thus, irrespective of microbial niche, the microbial community structure (composition and
145 relative abundance of ASVs) of F₁ laboratory-colonized larvae differed by collection location.
146 Pairwise PERMANOVA comparison of Bray-Curtis distances further showed significant
147 differences in microbial community structure in larval internal ($q<0.01$) and cuticle surface
148 microbiota ($q<0.01$) between every pair of collection site (Table 2a). This location-driven
149 heterogeneity in microbial community structure was further demonstrated by principal coordinate

150 analysis (PCoA), where F_1 larval internal and cuticle surface microbiota clustered distinctly by
151 collection site (Fig. 2).

152 Non-pairwise Shannon diversity comparisons showed significant differences in internal
153 ($p=0.009$) but not cuticle surface ($p=0.09$) microbiota of F_1 laboratory-colonized larvae from
154 different collection sites, indicating that there was inter-sample variation in the diversity of
155 internal but not cuticle surface microbiota of larvae when all collection sites were taken into
156 consideration. A pairwise Kruskal-Wallis comparison of Shannon diversity indices showed that
157 the inter-sample variation in diversity of larval internal microbiota held true when every pair of
158 collection sites was considered except between Las Cruces 3 and 4 (Table 2b and Suppl. 1).
159 Larvae originating from Las Cruces 3 had the highest internal microbiota diversity, followed by
160 Las Cruces 4 and El Terrero (Suppl. 1)

161

162 **Cuticle surface, but not internal, microbiota of laboratory colonized F_1 adult *An.***
163 ***albimanus* differed by geographic location**

164 Non-pairwise Bray-Curtis diversity comparisons showed significant differences in cuticle surface
165 ($p=0.001$), but not internal microbiota ($p=0.12$) between adult F_1 mosquitoes from different
166 collection sites, suggesting a loss of location-driven heterogeneity in microbial community
167 structure in internal but not cuticle surface microbial niche of laboratory-colonized F_1 adults.
168 Pairwise PERMANOVA comparisons of Bray-Curtis distances also showed significant
169 differences in microbial community structure of F_1 adult cuticle surface microbiota ($q<0.01$)
170 between every pair of collection sites (Table 2a). These results were corroborated by PCoA
171 which showed that F_1 adult cuticle surface microbiota, but not internal microbiota, clustered
172 distinctly by collection site (Fig. 2).

173 Non-pairwise Shannon diversity comparisons showed no differences in the internal ($p=0.42$) or
174 cuticle surface ($p=0.4$) microbiota of F_1 adults from different collection sites, indicating that there
175 was little or no inter-sample variation in diversity of F_1 adult microbiota when all collection sites
176 were taken into consideration. Pairwise Kruskal-Wallis comparisons of Shannon diversity indices
177 also detected no inter-sample variation in diversity of F_1 adult cuticle surface or internal
178 microbiota when every pair of collection site was considered (Table 2b and Suppl. 1).

179 **Laboratory-colonized F_1 *An. albimanus* larvae comprised a rich and diverse microbiota**
180 **that differed by geographic location**

181 Overall, ASVs from larval internal microbiota were assigned to 180 bacterial taxa, and cuticle
182 surface microbiota to 194 bacterial taxa (suppl. 2.). A majority of these taxa across all locations
183 (ranging from 118-139 taxa) were shared between the internal and cuticle surface microbiota
184 (Fig 3a), as well as across collection sites ($n=110$ for cuticle surface and $n=117$ for internal
185 microbiota) (Fig 3b). While a majority of the identified microbial taxa were shared between both
186 microbial niches, their abundance was generally higher in internal (Fig 4a) compared to cuticle
187 surface (Fig 4b) microbiota. Although a majority of identified microbial taxa in both internal and
188 cuticle surface microbiota were shared across all locations, their abundance differed by location
189 (Fig 4a, 4b and 5).

190 In general, larval internal microbiota was dominated by ASVs identified as an uncharacterized
191 *Enterobacteriaceae*, *Leucobacter*, *Thorsellia*, and *Chryseobacterium* (Fig 4a), together making
192 up over 50% of ASVs (Suppl. 2). In contrast, *Acidovorax*, uncharacterized *Comamonadaceae*,
193 and *Paucibacter* (Fig 4b) made up over 50% of ASVs detected on the larval cuticle surface
194 (Suppl. 2).

195 A few predominant bacterial taxa were present in the larval internal microbial niche across all
196 three collection sites: unclassified *Enterobacteriaceae*, *Thorsellia*, *Rhizobium*, *Xantobacter*,
197 *Acidovorax* and *Pirellula* (Fig 4a.), with remaining taxa showing different patterns of abundance
198 between collection sites (Fig 5 and Suppl. 2). For example, ASVs assigned to the genus
199 *Azoracus* were predominant in larvae from El Terrero, while *Singulisphaera*, *Paucibacter*,
200 *Ancylobacter*, *Gemmobacter*, and *Rayranella* were predominant in those from Las Cruces 3.
201 Predominant in larvae from Las Cruces 3 and Las Cruces 4 were *Azospirillum*, *Bosea* and
202 *Microbacterium*; and in those from El Terrero and Las Cruces 3 were *Terrimicrobium* and
203 *Legionella* (Fig 5). Although outside of the cut off limit set for differential abundance, ASVs
204 assigned to the bacterial genera *Leucobacter* were predominant in the internal microbiota of
205 larvae from Las Cruces 4 and El Terrero, but predominant in only three of the 14 sample
206 replicates from Las Cruces 3 (Fig 4a). Similarly, *Chryseobacterium* was predominant in Las
207 Cruces 4 and 3, but only predominant in six of the 16 pools of larvae from El Terrero. Bacterial
208 taxa that were unique to each location comprised <8% of all taxa in larval internal microbiota
209 (Suppl. 3b), and were below the threshold for inclusion in the heatmap and differential
210 abundance testing (Suppl. 2).

211 Unlike the internal microbial niche, no microbial taxa was predominant in larval cuticle surface
212 microbiota across all three collection sites. However, some taxa showed notable patterns of
213 abundance between locations (Fig 5). These included the genus *Azoarcus*, which was detected
214 at low to moderate frequencies in 13 of 15 pools of larvae from El Terrero, at low frequency in a
215 single pool of larvae from Las Cruces 3, and was not detected at all in Las Cruces 4 (Fig 4b and
216 6). Similarly, ASVs assigned to the genus *Spirosoma* were detected at moderate frequencies in
217 all pools of larvae from Las Cruces 4, but only in a few pools from the other two locations. ASVs
218 assigned to the genus *Paucibacter* were present at relatively higher abundance in larvae from

219 both Las Cruces 3 and El Terrero compared to those from Las Cruces 4. Those assigned to the
220 genus *Acidovorax* were predominant in larvae from Las Cruces 3 and Las Cruces 4 in contrast
221 to El Terrero. ASVs assigned to *Microbacterium*, *Bdellovibrio* and *Pelomonas* were present at
222 moderate frequencies in larvae from both Las Cruces 3 and Las Cruces 4 but were not detected
223 in El Terrero (Fig. 5). Bacterial taxa that were unique to each collection site comprised <8% of
224 larval cuticle surface microbiota (Fig 3b), and were below the threshold for inclusion in the
225 heatmap and differential abundance testing (Suppl. 2).

226

227 **Laboratory-colonized adult F₁ *An. albimanus* were comprised of sparse internal and**
228 **cuticle surface microbiota that were dominated by ASVs assigned to the genus *Asaia***
229 ASVs from adult internal microbiota were assigned to 62 microbial taxa and cuticle surface
230 microbiota were assigned to 106 microbial taxa. Two of these ASVs which were only present in
231 the cuticle surface microbiota were classified as archaea, while all other remaining ASVs were
232 classified as bacteria (Suppl. 2). Unlike larval microbiota, less than half of the assigned taxa
233 across all locations (ranging from 19-37 taxa) were shared between internal and cuticle surface
234 microbiota (Fig 3a), and only 18 taxa on the cuticle surface and 19 internal taxa were shared
235 across all maternal collection sites (Fig 3b).

236 Overall, ASVs assigned to the bacterial genus *Asaia* dominated both adult internal and cuticle
237 surface microbiota (Fig 6), constituting at least 70% of taxa in each microbial niche (Suppl. 2). A
238 majority of identified taxa in adult internal, but not cuticle surface, microbiota was detected
239 across all three collection sites, with a few of these taxa present in high abundance across all
240 collection sites (Fig. 6). Across all three collection sites, ASVs assigned to the genera
241 *Acinetobacter*, *Gluconobacter*, *Pantoea* and *Pseudomonas* were present in moderate to high
242 abundance in adult internal microbiota in addition to *Asaia* (Fig 6).

243 No site-specific microbial taxa were identified in adult cuticle surface microbiota, as the ASV
244 distribution did not meet the criteria for this type of analysis. This was compounded by
245 dominance (>70%) of ASVs that were assigned to the bacterial genus *Asaia* (Suppl. 2). In
246 addition, the cuticle surface microbiota of adults originating from Las Cruces 4 comprised 43% of
247 all adult cuticle surface microbial taxa (Fig 3b), although a majority were of low abundance.

248 **Discussion**

249 The scientific community is increasingly investigating the role of mosquito microbiota in fighting
250 mosquito-borne diseases [9]. The successful transition of mosquito microbiome research from
251 laboratory to field requires a comprehensive understanding of the dynamics underlying the
252 composition of the microbiota of field-collected mosquitoes and their progeny. We provide a
253 comprehensive characterization of the internal and cuticle surface microbiota of laboratory-
254 reared F₁ progeny from field-caught adult *An. albimanus* that were collected from different
255 locations. Our results show that while location-driven heterogeneity in the microbial community
256 structure of both internal and cuticle surface microbiota was present in the F₁ larvae, this
257 heterogeneity was only evident in the cuticle surface microbiota of adult progeny from the same
258 generation. This work provides comprehensive fundamental data for studying how parentage
259 and environmental conditions differentially or concomitantly affect mosquito microbiome
260 composition.

261 Previous studies on other mosquito species have also shown a loss of field-acquired internal
262 mosquito microbiota following several generations of laboratory colonization [37, 38]. Until now,
263 however, the laboratory generation at which this loss occurs was largely undescribed. Our study
264 represents an initial step in filling this knowledge gap and provides further details regarding the
265 dynamics of changes in both the internal and cuticle surface microbial niches. The homogeneity
266 in internal microbiota of F₁ adults is suggestive of a loss of field-acquired microbiota, which may
267 have implications for studies that rely on laboratory progeny in lieu of field populations [25, 27,
268 28]. Previous findings showed that rearing mosquitoes in water from the field could preserve the
269 field-derived internal microbiota in laboratory-colonized adult progeny of *An. gambiae s. l.* for
270 several generations [37]. In addition, other studies, have shown that the microbiota in larval
271 habitat water significantly influences the internal microbiota of emerging adult mosquitoes [40,

272 41]. Thus, preserving wild-type microbiota in F_1 progeny could be an avenue for studying
273 mosquito-microbe dynamics in the field.

274 Maternal egg-smearing has been proposed as a mechanism through which adult female
275 mosquitoes and other insects transfer microbes to their progeny [42-45]. This egg-smearing
276 could explain the heterogeneity observed in internal and cuticle surface microbiota between F_1
277 larvae with different maternal origins. On the other hand, a lack of this heterogeneity in the
278 internal microbiota of adult progeny could be attributed to the physiological changes that occur
279 during metamorphosis and adult eclosion, whereby elimination of the larval meconial peritrophic
280 membrane and meconium (midgut and midgut content), along with ingestion of exuvial fluid—
281 which is said to be bactericidal—results in sterile or nearly sterile midguts in newly emerged
282 adults [39, 46]. This could additionally be explained by mechanisms that regulate the
283 composition of adult mosquito internal microbiota [47-49]. Conversely, the heterogeneity in
284 cuticle surface microbiota between adult progeny originating from different maternal sites,
285 suggests that maternally derived microbes in the rearing trays may have colonized adult cuticle
286 surfaces during emergence. The mechanisms underlying the assemblage of the mosquito cuticle
287 surface microbiome are largely undescribed, and thus require further investigation.

288 The low inter-sample variation in microbial diversity observed in this study has largely been
289 described in laboratory mosquito colonies [38, 50]. The microbial composition of laboratory-
290 reared larvae is typically less diverse [50, 51] compared to those of field-derived larvae, but our
291 laboratory-reared larvae exhibited a rich microbial composition that was comparable to those of
292 field populations [39, 40]. In contrast, our adult progeny had a less diverse microbial composition
293 that was reflective of typical laboratory-reared adult mosquitoes [36, 52]. Further suggesting that
294 field-acquired microbiota, although transferred to laboratory progeny, may be lost within one
295 generation of laboratory colonization—particularly at the adult stage.

296 In this study, we detected microbial taxa that have previously been identified in *Anopheles* and
297 other mosquito genera [7, 41, 53, 54]. While a majority of the taxa in F₁ larvae were shared
298 between both the internal and cuticle surface microbial niches, a greater abundance of microbial
299 taxa was detected in the internal microbial niche compared to the cuticle surface. The cuticle
300 surface microbiota of mosquitoes and other hematophagous insects are largely uncharacterized
301 and the mechanisms underlying their assemblage remain unknown. As such, we hypothesize
302 that although both internal and cuticle surface niches are exposed to the same water from which
303 the microbiota is derived, a more conducive and/or selective internal environment could allow for
304 greater proliferation of colonizing bacteria. In F₁ adults however, less than half of the detected
305 microbial taxa were shared between the internal and cuticle surface microbial niches, suggesting
306 differences in physiological conditions that favor microbial colonization, and corroborating
307 findings that point toward microbial regulatory mechanisms within the mosquito midgut [47-49].
308 Although a few microbial taxa overlapped between adult internal and cuticle surface microbial
309 niches and the most abundant taxa were shared, many of the unshared taxa have been
310 previously detected in adult mosquitoes including *Anopheles* [1, 54, 55], indicating that the
311 cuticle surface microbiota characterized in this study are inherently associated with mosquitoes.
312 Like the larval microbiota, there was a higher abundance of microbial taxa in the adult internal
313 microbial niche compared to the cuticle surface, further supporting the hypothesis of a more
314 conducive and/or selective internal environment.

315 With the exception of the adult cuticle surface microbial niche, a majority of all detected microbial
316 taxa overlapped between collection sites in both F₁ larvae and adults, albeit with differing
317 abundances. This reflects restrictions imposed by controlled laboratory environments in the
318 development of mosquito microbiota. In both microbial niches of both larvae and adults,
319 microbial taxa that were specific to collection sites were low in abundance, compared to the

320 moderate to high abundance of those that were shared across all locations. This was particularly
321 true for *Asaia*—notorious for rapidly colonizing laboratory mosquitoes [56]—which constituted at
322 least 70% of both adult internal and cuticle surface microbiota from progeny across all collection
323 sites. These results suggest that field-acquired mosquito microbiota may be lost in as early as
324 the first generation of laboratory colonization.

325 We recognize that not having the microbial community profiles of the mothers from which the F_1
326 progeny were derived is a limitation of this study. However, the findings herein provide empirical
327 data on the composition of laboratory reared F_1 *An albimanus* microbiota from different
328 locations. It provides a foundation for exploring the role of parentage, environmental conditions,
329 and inherent host physiological characteristics on the assemblage of the mosquito microbiome,
330 as well as the fate of field-derived microbes upon laboratory colonization. This is critical for
331 advancing mosquito microbiome studies and their applications beyond laboratory settings.

332 **Methods**

333 The findings presented here extend those of a larger study [8]. Thus, the mosquito collection,
334 processing and sequencing procedures have previously been described in detail [8].

335 **Mosquito collections and laboratory generation of F₁ progeny**

336 Gravid and/or blood-fed adult female *An. albimanus* were sampled across four field sites in the
337 villages of Las Cruces and El Terrero, in La Gomera, department of Escuintla, Guatemala
338 (Figure 1). Field-collected mosquitoes from each location were held in separate paper cups,
339 sustained on 10% sucrose solution, and transported to the insectary at Universidad del Valle de
340 Guatemala in Guatemala City for species identification, oviposition and subsequent rearing of F₁
341 progeny. Mosquitoes that were morphologically identified as *An. albimanus* following
342 identification keys [57] (approximately 300 in total) were subjected to oviposition, and a
343 subsample of the resulting F₁ progeny was used for molecular verification of species identity as
344 described below. A modified oviposition procedure [58] was employed, wherein no more than 70
345 gravid females from the same location were placed in quart size paper ice cream containers with
346 distilled water to a depth of at least 2 cm. The containers were covered with fine mesh fabric that
347 were secured with rubber bands prior to the introduction of gravid field-caught adult *An.*
348 *albimanus*. The mesh was topped with cotton balls soaked in 10% sucrose to sustain the
349 mosquitoes, and subsequently covered with a thick piece of black plastic bag to keep the
350 containers dark and trap in moisture. The oviposition chambers were held under the following
351 insectary conditions; 27±2°C, 80±10% relative humidity, and 12-h light-dark cycle. After at least
352 48 h, the adult females were removed and discarded as they were not needed for the original
353 study. The F₁ eggs were collected, pooled by location and reared separately under identical
354 ambient conditions (as described above) and the following larval feeding regimen. Eggs were

355 washed into 18 x 14 x 3 inch plastic larval trays (approximately 200 eggs per tray) containing
356 distilled water to a depth of at least 2 cm, and 3-4 drops of 10% yeast solution. Hatched larvae
357 were sustained on finely ground Koi fish food (Foster & Smith, Inc. Rhinelander, WI) until
358 pupation. At the third to fourth larval instar stage (L3-L4), half of the larvae were separated and
359 used for bioassays in the original study and subsequently processed for microbiota
360 characterization (n=132). Using a stereo microscope, female pupae from the remaining
361 mosquitoes were separated into 8 oz. paper ice cream cups and placed into cardboard cages for
362 adult eclosion. The resulting F₁ adult virgin female mosquitoes were sustained on 10% sucrose
363 solution until they were 2-5 days old and also used for bioassays in the parent study. These
364 were subsequently processed for microbiota characterization (n=135). Post bioassays, larval
365 and adult samples were preserved in RNALater® solution (Applied Biosystems, Foster City,
366 CA), shipped on dry ice to the US Centers for Disease Control and Prevention (CDC) in Atlanta,
367 USA, and stored at -80 °C until further processing. Samples originating from different geographic
368 locations were handled, stored and processed separately.

369 **Genomic DNA extraction from mosquito legs, whole mosquito samples, and mosquito**
370 **cuticle surfaces**

371 Stored mosquito samples were thawed overnight at 4°C, and the thawed RNALater® solution
372 was discarded. This was followed by a rinse with nuclease-free water to remove any residual
373 RNALater®. Legs from a subsample of individual F₁ adults were removed using sterile forceps
374 and placed in individual sterile 1.5 mL Eppendorf tubes containing 75 µL Extracta DNA Prep
375 solution (Quantabio, Beverly, MA) for DNA extraction following manufacturer's instructions. Both
376 larvae and adult (with and without legs removed) F₁ samples were pooled for whole body and
377 cuticle surface DNA extractions. Each pool comprised 3 individuals, resulting in 44 pools of
378 larvae and 45 pools of adults. Table 1 shows the number of pools (replicates) processed per life

379 stage and collection site. To obtain cuticle surface samples, each pool was submerged in 500 μ L
380 of nuclease free water and washed vigorously by agitating with a vortex mixer for at least 15
381 seconds. The resulting wash water was transferred to new sterile tubes for DNA extraction. We
382 acknowledge that some microbes may have been lost via rinsing off the preservative
383 (RNALater® solution) prior to this step. A comprehensive (comparable or more than the internal
384 community) microbial community was nonetheless recovered from the cuticle surface samples.
385 The washed sample pools were further surface sterilized using two vigorous washes; first in 70%
386 ethanol, then nuclease free water, each with at least 15 seconds agitation on a vortex mixer.
387 This was followed by one gentle rinse with nuclease free water. Genomic DNA from the surface-
388 sterilized samples and wash water (subsequently referred to as internal and cuticle surface,
389 respectively) was isolated using the TissueLyser II and DNeasy Blood and Tissue Kit (QIAGEN,
390 Hilden, Germany) as follows: in individual 2 mL sterile tubes, 180 μ L of buffer ATL (QIAGEN)
391 and 5 mm diameter stainless steel beads (QIAGEN) were added to each sample pool. With the
392 following settings: 30 hz/s for 8 and 15 minutes for internal and cuticle surface samples
393 respectively, samples were homogenized using the TissueLyser (QIAGEN) fitted with two sets of
394 96-well adapter plates that held the tubes. The plates were rotated every minute during
395 homogenization, and afterwards, samples were transferred into new sterile 1.5 mL Eppendorf
396 tubes for the remaining part of the DNA extraction process. 20 μ L of Proteinase K (QIAGEN)
397 was added to each homogenized sample and incubated overnight at 56 °C, after which 200 μ L
398 of buffer AL (QIAGEN) was added and incubated for a further 2 h at the same temperature. The
399 remaining steps were performed according to QIAGEN's spin-column protocol for purification of
400 DNA from animal tissues, and the purified DNA was eluted in 70 μ L of buffer AE (QIAGEN). Two
401 sets of two blank controls (without samples) were processed alongside the internal and cuticle
402 surface samples, and all steps, along with those described below, were performed under sterile

403 conditions. The purified genomic DNA from samples and blank controls were stored at -80 °C
404 until further processing.

405 **Molecular species confirmation of *An. albimanus* and 16S rRNA amplicon sequencing**

406 DNA samples from mosquito legs were used as templates to amplify the second internal
407 transcribed spacer region (ITS2) of the mosquito ribosomal DNA in order to verify morphological
408 identification of *An. albimanus*. This was achieved by conventional PCR using the universal ITS2
409 primers (ITS2 A: TGTGAACTGCAGGACACAT and ITS2 B: TATGCTTAAATTCAGGGGGT) for
410 distinguishing members of *Anopheles* complexes [59]. With a final reaction volume of 25 µL,
411 each PCR comprised ≥100 ng/µL DNA template, 15 µM of each primer, 12.5 µL of 2X AccuStart
412 II PCR SuperMix (Quantabio, Beverly, MA), and PCR grade water to final volume. With the
413 following conditions, initial denaturation at 94 °C for 4 min, then 35 cycles of 94 °C for 30 s,
414 53 °C for 40 s, and 72 °C for 30 s, followed by a final extension for 10 min at 72 °C, the
415 reactions were performed using a T100™ Thermal Cycler (Bio-Rad, USA). Using EtBr-stained
416 agarose gel electrophoresis, the amplified products sized ~500 bp, confirmed all samples as *An.*
417 *albimanus* [60].

418 DNA from internal and cuticle surface samples, along with those from blank controls, were used
419 as templates to amplify the universal bacterial and archaeal 16S rRNA gene. This was also
420 achieved by conventional PCR using universal primers targeting the v3-v4 region of the 16S
421 rRNA gene (341F:
422 **TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG**CCTACGGGNGGCWGCAG, and 805R:
423 **GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAG**ACTACHVGGGTATCTAATCC), each
424 with Illumina® (San Diego, CA USA) adapters (bold typeface). With a final reaction volume of 25
425 µL, each PCR comprised; ≥20 ng/µL DNA template, 5 µM of each primer, 10 µL of 2x KAPA

426 HiFi HotStart PCR mix (Roche, Switzerland), and PCR grade water to final volume. Three
427 negative controls with PCR grade water substituted for DNA template, were processed along
428 with the samples. The reactions were performed using the T100™ Thermal Cycler (Bio-Rad,
429 USA) under the following conditions; initial denaturation at 95°C for 3 min, then 25 cycles of
430 95°C, 55°C, and 72°C for 30s each, followed by a final extension for 5 min at 72°C.
431 The amplification products, sized ~460 bp, were verified by electrophoreses as described above
432 and quantified using a NanoDrop™ spectrophotometer (Thermo Fisher Scientific, Waltham,
433 MA). Amplicons, including PCR products from blanks and negative controls, which yielded no
434 bands following electrophoresis, were purified using Agencourt AMPure XP beads (Beckman
435 Coulter Inc., Indianapolis, IN, USA) at 0.7X (internal) or 0.875X (cuticle surface) sample volume,
436 and eluted in 40 µL of 10 mM Tris buffer (pH 8.5).

437 All purified products, including those of blanks and negative controls, were used as templates for
438 sequencing library preparation. This was accomplished via index PCR using the Nextera XT
439 Index kit v2 sets, A, B and D (Illumina, San Diego, CA). With a final reaction volume of 50 µL,
440 each index PCR comprised; 25 µL NEBNext High-Fidelity 2X PCR master mix (New England
441 Biolabs Inc., Ipswich, MA), 5 µL of each index primer, 10 µL of purified PCR products (0–
442 20 ng/µL) as template, and PCR grade water to final volume. The PCRs were performed under
443 the following reaction conditions; 98°C for 30s, then 8 cycles of 98°C for 10s, 55°C and
444 65°C for 30s each, and a final extension at 65°C for 5 min, and resulting libraries were
445 also cleaned using Agencourt AMPure XP beads (Beckman Coulter Inc., Indianapolis, IN, USA)
446 at 1.2X sample volume and eluted in 25 µL of 10 mM Tris buffer (pH 8.5). The libraries were
447 subsequently analyzed for size and concentration, normalized to 2 nM, pooled and denatured
448 following Illumina guidelines for loading onto flowcells. Sequencing was performed on an
449 Illumina HiSeq 2500 machine, using 2x250 cycle paired-end sequencing kits.

450 **Preprocessing of sequencing reads**

451 Sequencing outputs were demultiplexed and converted to the fastq format for downstream
452 analysis using the bcl2fastq (v2.19) conversion software (Illumina®). A total of 115,250,077
453 demultiplexed paired-end sequencing reads, with a maximum length of 250 bp were initially
454 imported into the 'quantitative insights into microbial ecology' pipeline, QIIME2 v.2017.7.0 [61],
455 and further sequencing read processing and analysis were performed in v.2018.2.0 of the
456 pipeline. Using the DADA2 plugin in QIIME 2 [62], the `denoise-paired` command with the
457 following options; `trunc_len_f: 244, trunc_len_r: 244, and n_reads_learn:`
458 `500000`, was used to correct errors, remove chimeras and merge paired-end reads. The
459 resulting amplicon sequence variants (ASVs; n=30,956,883) were further filtered to remove
460 potentially extraneous ASVs (those with <100 counts) and ASVs that were associated with
461 blanks and negative controls. This last filtering step resulted in 17,225,776 ASVs, ranging from
462 3,277 to 223,222 per sample (mean 96,774), that were used for downstream comparison of
463 bacterial composition and taxonomic analysis. Suppl. 3. shows sequencing reads and ASV
464 summary statistics.

465 **Diversity indices**

466 Analysis of microbial diversity within (alpha diversity) and between (beta diversity) samples were
467 performed in QIIME2 using the Shannon diversity index and Bray-Curtis dissimilarity index,
468 respectively. The Shannon diversity indices were calculated using rarefied ASVs counts per
469 sample, in which ASVs per sample were selected randomly without replacement at an even
470 depth (Suppl. 4) for ten iterations. The resulting average Shannon indices are presented and
471 were compared between samples using pairwise Kruskal-Wallis tests with Benjamini-Hochberg
472 false discovery rate (FDR) corrections for multiple comparisons.

473 The Bray-Curtis dissimilarity indices were computed with or without rarefaction, and resulting
474 indices were compared between samples using pairwise PERMANOVA tests (999 permutations)
475 with FDR corrections. There were no discernable differences between results of rarefied and
476 non-rarefied data. Thus, results of Bray-Curtis dissimilarity indices using non-rarefied data were
477 visualized by Principal Co-ordinates Analysis (PCoA) plots in R [63] using the phyloseq R
478 package [64].

479 Significance for both pair-wise analyses was set to $q < 0.05$ (i.e. post FDR p -value corrections).

480 **Taxonomic analysis and differentially abundant microbial taxa**

481 Taxonomic analysis of ASVs was performed using QIIME2's pre-trained Naïve Bayes classifier
482 [65] and `q2-feature-classifier` plugin [66]. Prior to analysis, the classifier was trained on the
483 QIIME-compatible 16S SILVA reference (99% identity) database v.128 [67], and using the
484 `extract-reads` command of the `q2-feature-classifier` plugin, the reference sequences
485 were trimmed to the v3-v4 region (425 bp) of the 16S rRNA gene. The relative abundance of
486 annotated ASVs across samples were subsequently visualized using the `qiime feature-`
487 `table heatmap` plugin based on Bray-Curtis distances with the plugin's default clustering
488 method. Only annotated ASVs with counts ≥ 2000 (larvae) or ≥ 1000 (adults) were included in
489 the heatmaps.

490 Differentially abundant microbial taxa across locations were identified using QIIME2's analysis of
491 composition of microbiomes (ANCOM) [68] plugin. The cut-off for differential abundance was set
492 to an effect size of $\log F \geq 20$ and $W \geq 20$, i.e. a taxon was differentially abundant across collection
493 sites if the ratio of its abundance to those of at least 20 other taxa (25% of all included taxa)
494 differed significantly across sites.

495 Prior to each analysis, ASV frequency data was normalized by \log_{10} transformation following the
496 addition of pseudocounts of 1. To ensure that filtering of low frequency reads did not
497 compromise our findings, all downstream analyses were also performed with these reads
498 included, and results (Suppl 5) were consistent with those of the clean/filtered dataset. Results
499 of the clean dataset are subsequently discussed.

500 The outputs of data analyses were aesthetically formatted using Inkscape [69].

501 **Declarations**

502 **Ethics approval and consent to participate:** Not applicable

503 **Consent for publication:** Not applicable

504 **Availability of data and material:** The raw sequencing reads generated from this project,
505 including those from negative controls (blanks), have been deposited in the National Center for
506 Biotechnology Information (NCBI), Sequence Read Archive under the BioProject
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515 rearing; ND[†] & MS performed molecular analysis and sequencing; ND[†] analyzed the data; ND[†]
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760 **Supporting Information**

761 **Suppl. 1.** Shannon alpha diversity plots

762 **Suppl. 2.** Frequency and relative abundance of microbial taxa detected in internal and cuticle

763 surface microbiota of F₁ laboratory colonized *An. albimanus* larvae and adults

764 **Suppl. 3.** Summary statistics of sequencing reads and amplicon sequence variants (ASVs) used

765 for downstream analysis

766 **Suppl. 4.** Rarefaction depth and plots of sequencing reads

767 **Suppl 5.** Results of downstream analysis including low frequency (potentially extraneous) reads

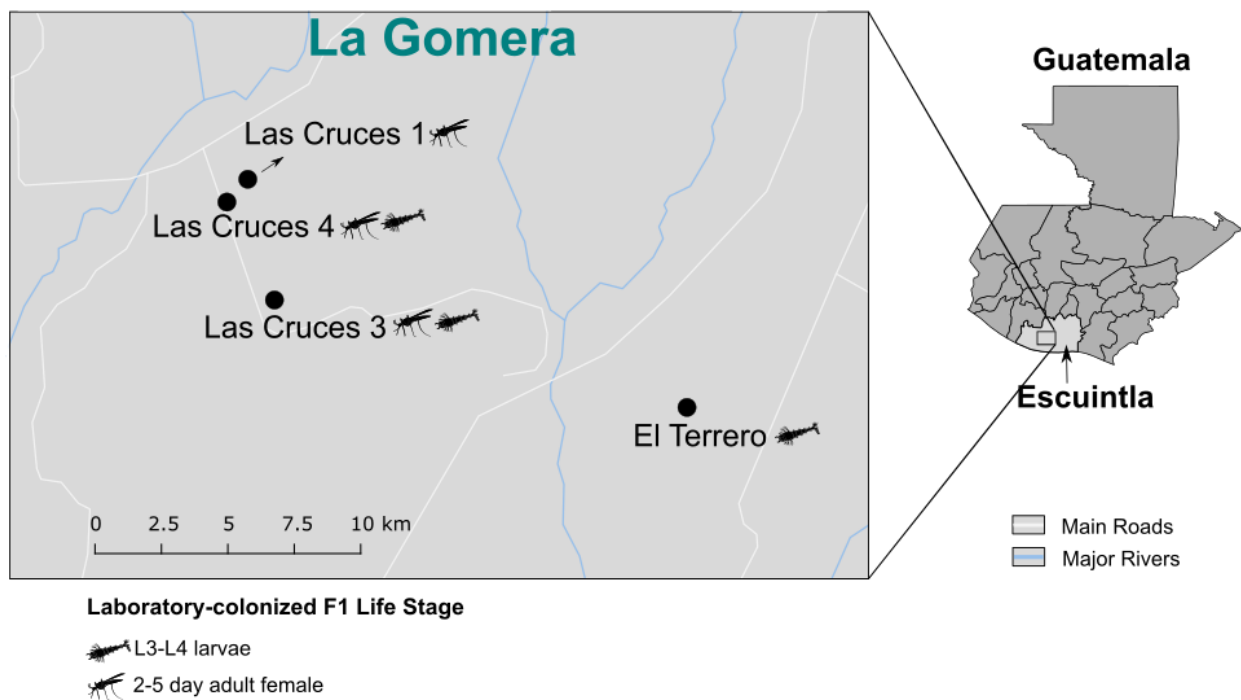
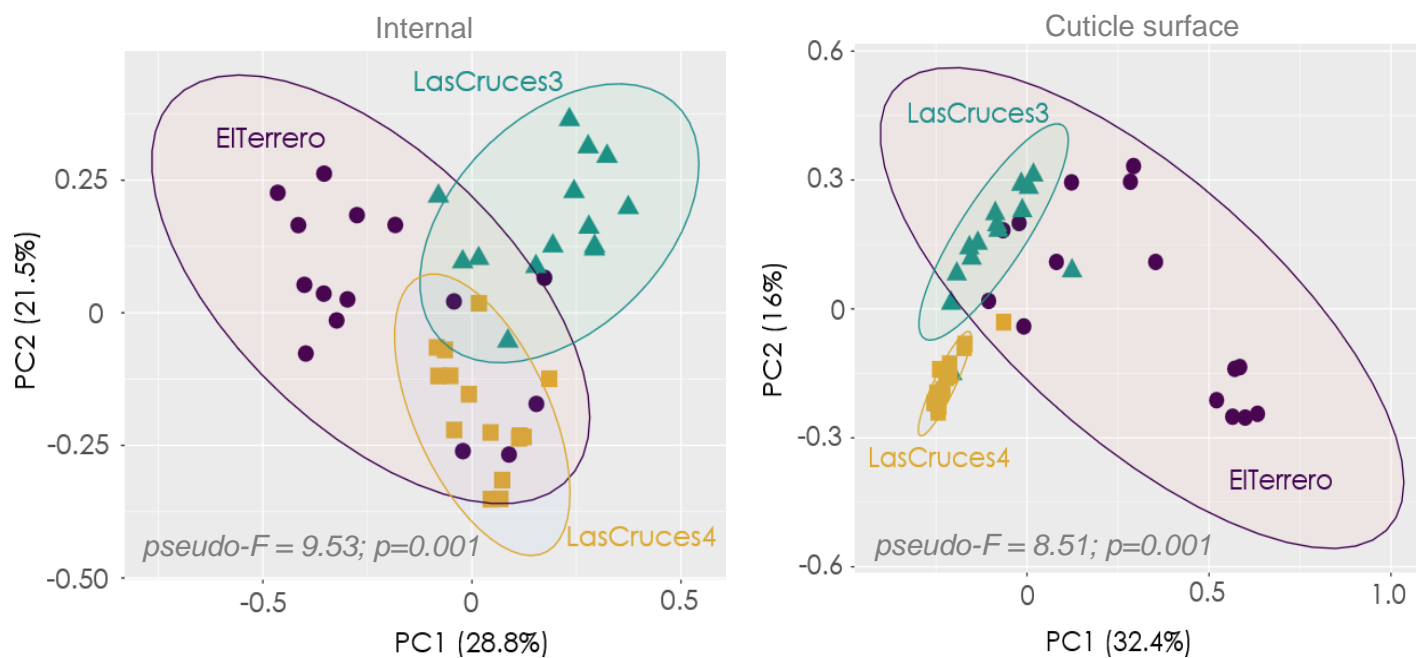


Figure 1: Map showing the geographical origins of F₁ *Anopheles albimanus*. F₁ laboratory-colonized mosquitoes were derived from gravid and/or blood fed females collected from each location. Mosquito icons show the geographical origin of the F₁ life stages studied.

LARVAE



ADULTS

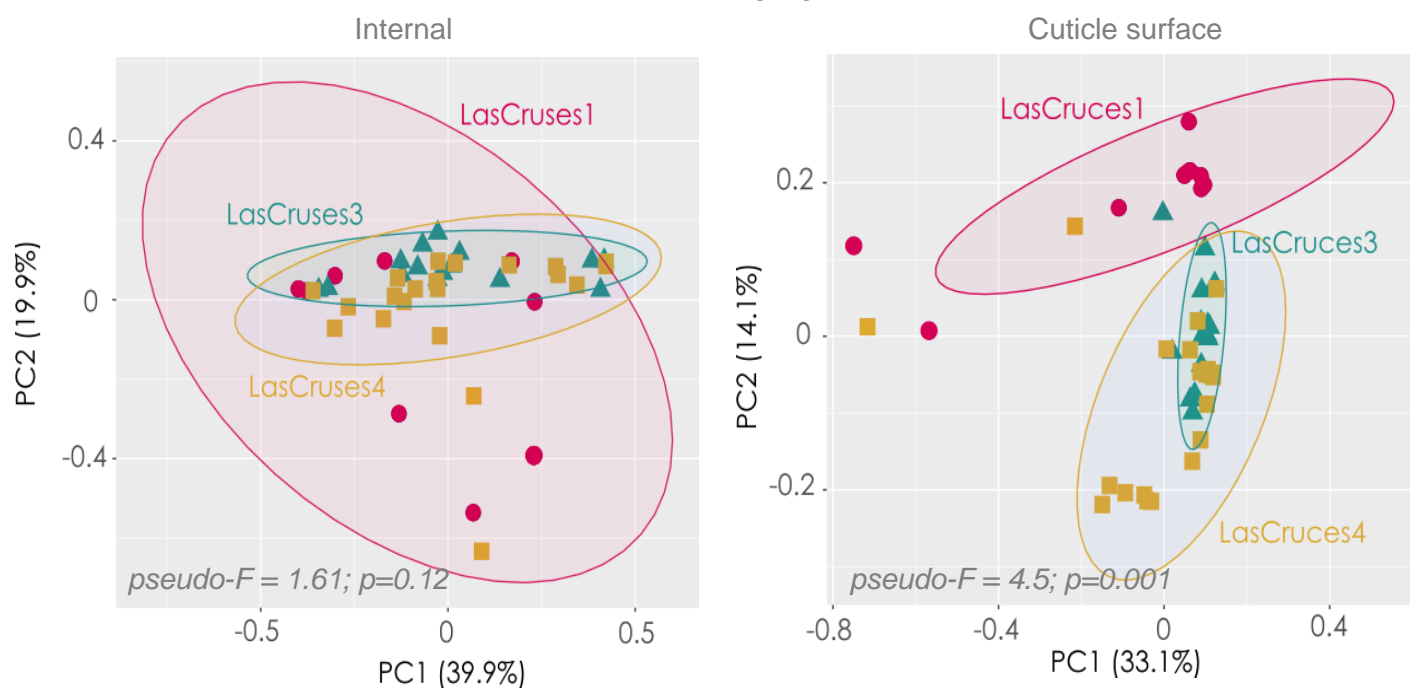


Figure 2. Principal coordinate analysis (PCoA) ordinations of internal and cuticle surface microbiota from F₁ laboratory-colonized *An. albimanus*. The PCoA plots are based on Bray-Curtis distances between the microbiota of mosquitoes with differing collection sites. Each point on the plot represents the microbial composition of a pool of three individuals, and mosquito pools are color-coded by their origin. For larvae, the first two principal component (PC) axes captured 50% (internal) and 48% (cuticle surface) of the variance in the data, with both internal

and cuticle surface microbiota clustering distinctly by collection site. For adults, the first two PC axes captured 59% (internal) and 47% (cuticle surface) of the variance in the data, with cuticle surface but not internal microbiota clustering distinctly by collection site. PERMANOVA statistics are presented at the bottom of each plot.

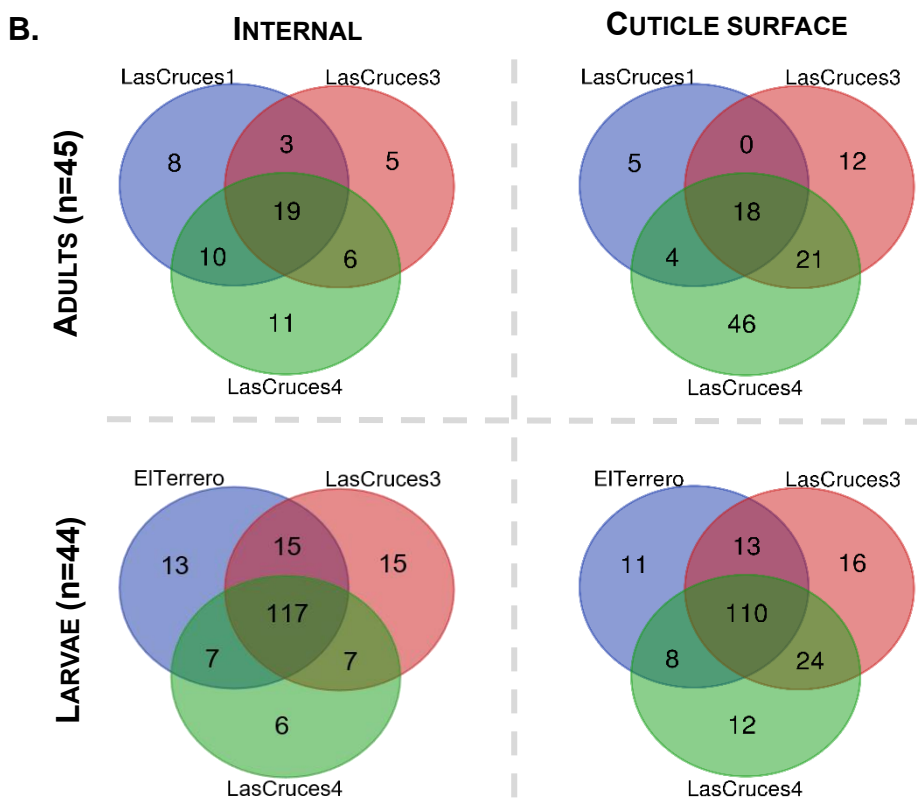
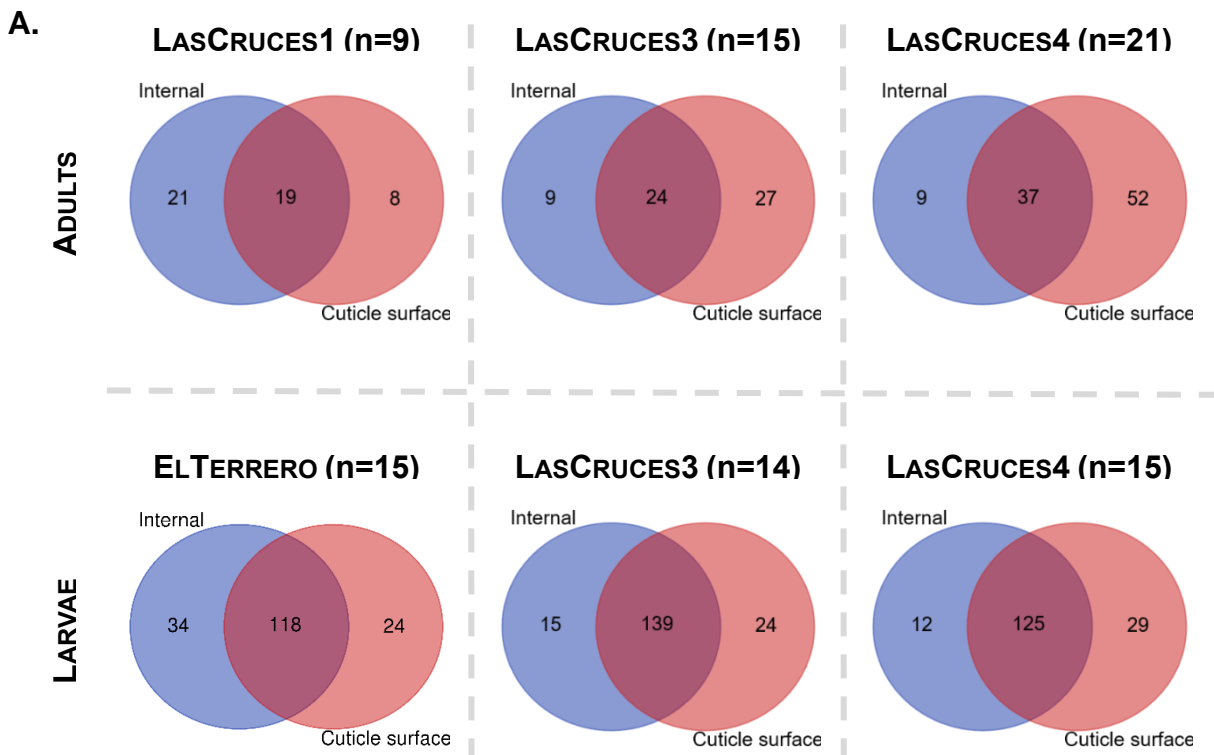
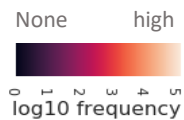


Figure 3. Number of unique and shared microbial taxa between microbial niches (A); number of unique and shared microbial taxa between collection sites (B). The number of taxa shown in the Venn diagram represent bacterial taxa except in the cuticle surface microbiota of adults from Las Cruces 4, where two archaeal taxa were present. n = pools of mosquito samples analyzed per location or microbial.



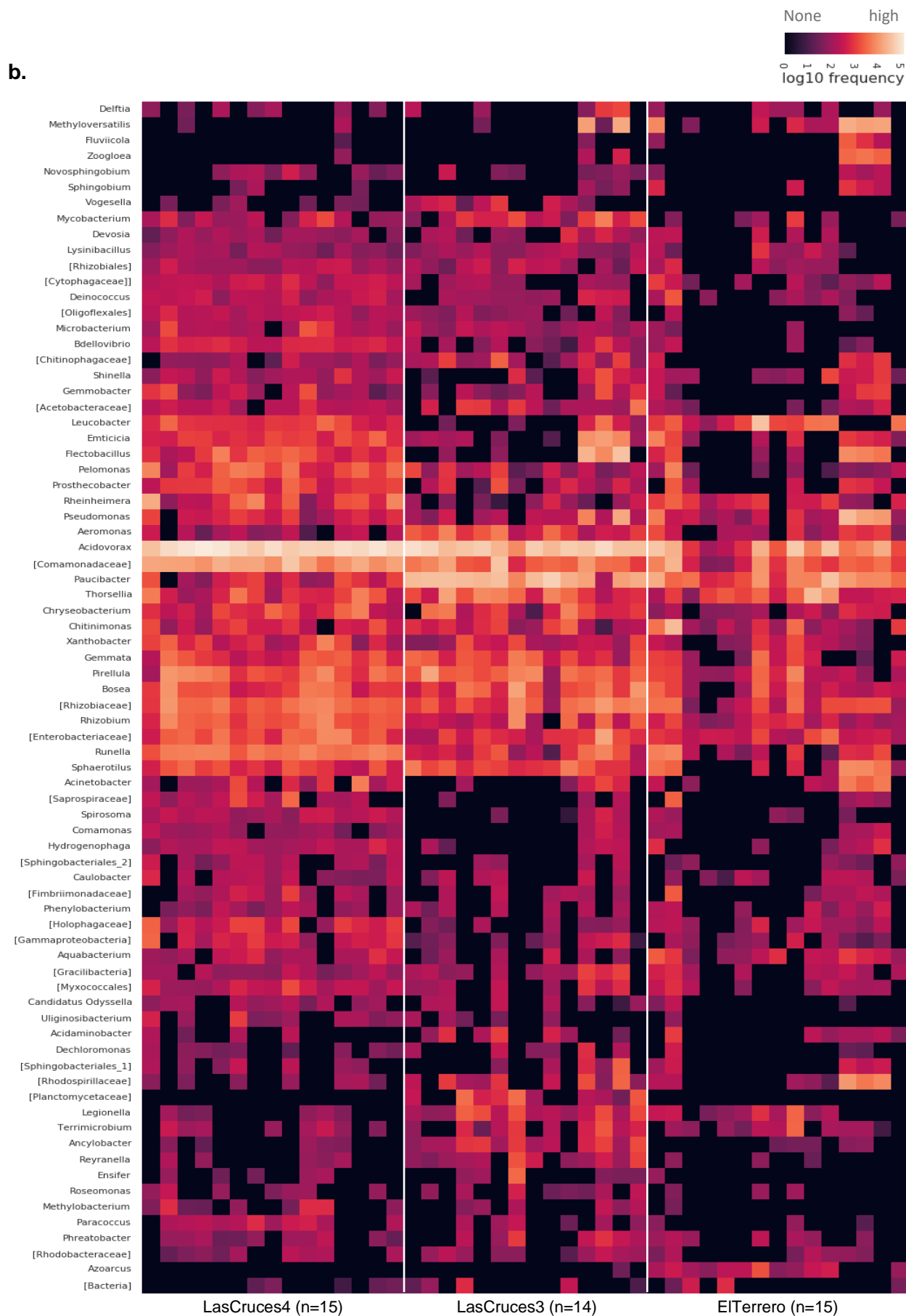


Figure 4. Frequency of ASVs from the internal (a) and cuticle surface (b) microbiota of laboratory-colonized *An. albimanus* F₁ larvae originating from different collection sites.

ASVs were annotated to the genus level or the lowest possible taxonomic level (in square brackets) and are clustered by the average nearest-neighbors chain algorithm. Only taxonomically annotated ASVs with frequencies ≥ 2000 are presented.

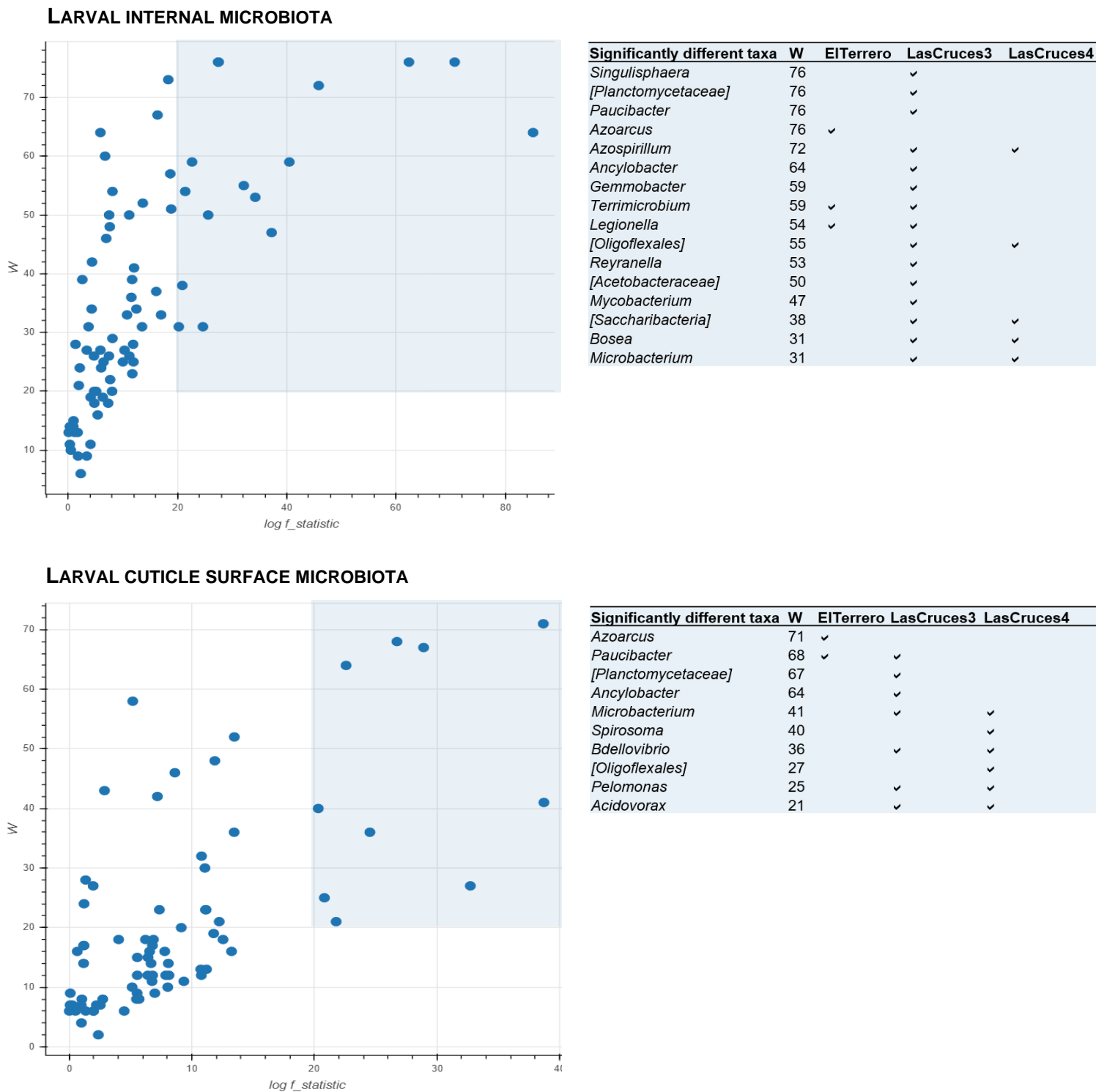


Figure 5. Volcano plots of differentially abundant bacterial taxa in F₁ laboratory colonized *An. albimanus* larvae originating from three different collection sites. The plots show results of analysis of composition of microbiomes (ANCOM) tests for differentially abundant microbial taxa between collection site, with an effect size set to log F₂₀, and a cut-off of

differential abundance set to $W \geq 20$ (i.e., a taxon was differentially abundant across collection sites if the ratio of its abundance to those of at least 20 other taxa (25% of all included taxa) differed significantly across sites). Differentially abundant taxa are highlighted (blue shaded area) and the taxa names and locations in which they were most abundant are presented in the adjoining tables.

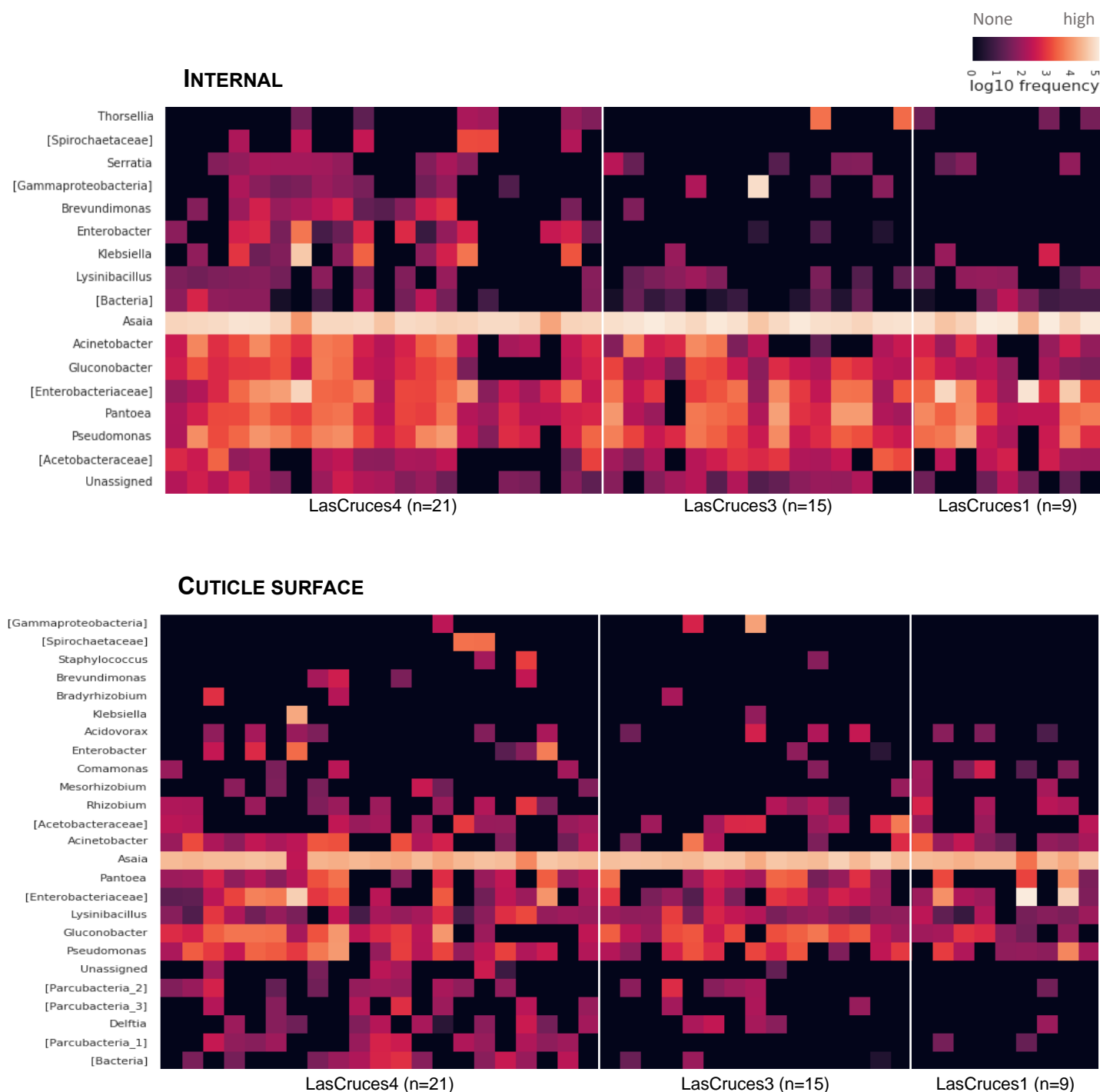


Figure 6. Frequency of ASVs from the internal and cuticle surface microbiota of laboratory-colonized F₁ adult *An. albimanus* originating from different locations. ASVs were annotated to the genus level or the lowest possible taxonomic level (in square brackets) and are clustered by the average nearest-neighbors chain algorithm. Only taxonomically annotated ASVs with frequencies ≥ 1000 are presented.

Table 1. Descriptive statistics of laboratory colonized F₁ *An. albimanus* processed, per life stage and collection site.

Collection site	Number of mosquitoes (number of pools)	
	L3-L4 larvae	Adult ♀s
El Terrero	45 (15)	-
Las Cruces 1	-	27 (9)
Las Cruces 3	42 (14)	45 (15)
Las Cruces 4	45 (15)	63 (21)

Table 2. Pairwise alpha and beta diversity comparisons of laboratory colonized F₁ *An. albimanus* microbiota from different collection sites. a. Pairwise beta (Bray-Curtis) diversity comparison showed significant differences in larval internal and cuticle surface microbiota between collection sites. In contrast, only adult cuticle surface but not internal microbiota were significantly different across collection sites. **b.** Pairwise alpha (Shannon) diversity comparison showed significant differences in larval internal but not cuticle surface microbiota between collection sites (two of the three pairs). In contrast, there was no significant difference in adult internal or cuticle surface microbiota between collection sites. Pairwise alpha and beta diversity comparisons were conducted using Kruskal-Wallis and PERMANOVA (999 permutations) tests respectively, with Benjamini-Hochberg FDR correction (q-value). Significance was determined at $q < 0.05$. n = No. of pools processed, and each pool comprised three individual mosquitoes.

a.

	Group 1	Group 2	Internal			Cuticle surface		
			pseudo-F	p-value	q-value	pseudo-F	p-value	q-value
Adults	LasCruces1 (n=9)	LasCruces3 (n=15)	2.21	0.088	0.161	5.80	0.001	0.003
	LasCruces1 (n=9)	LasCruces4 (n=21)	1.92	0.107	0.161	4.97	0.004	0.005
	LasCruces3 (n=15)	LasCruces4 (n=21)	0.85	0.443	0.443	3.07	0.005	0.005
Larvae	EITerrero ^a (n=15)	LasCruces3 (n=14)	9.81	0.001	0.001	4.48	0.001	0.001
	EITerrero ^a (n=15)	LasCruces4 (n=15)	7.98	0.001	0.001	11.43	0.001	0.001
	LasCruces3 (n=14)	LasCruces4 (n=15)	10.88	0.001	0.001	11.37	0.001	0.001

^aTwo pools were excluded from the analysis of cuticle surface microbiota following rarefaction

b.

	Group 1	Group 2	Internal			Cuticle surface		
			H	p-value	q-value	H	p-value	q-value
Adults	LasCruces1 (n=9)	LasCruces3 (n=15)	0.00	0.98	0.98	1.09	0.30	0.41
	LasCruces1 (n=9)	LasCruces4 (n=21)	0.49	0.48	0.72	1.23	0.27	0.41
	LasCruces3 (n=15)	LasCruces4 (n=21)	1.77	0.18	0.55	0.67	0.41	0.41
Larvae	EITerrero (n=15)	LasCruces3 (n=14)	6.63	0.01	0.02	4.45	0.03	0.10
	EITerrero (n=15)	LasCruces4 (n=15)	6.94	0.01	0.02	1.87	0.17	0.26
	LasCruces3 (n=14)	LasCruces4 (n=15)	0.49	0.48	0.48	1.10	0.29	0.29