#### Functional insights from the GC-poor genomes of two aphid 1

#### parasitoids, Aphidius ervi and Lysiphlebus fabarum 2

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### 23 Abstract

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## 26 Background

Parasitoid wasps have fascinating life cycles and play an important role in trophic
networks, yet little is known about their genome content and function. Parasitoids that
infect aphids are an important group with the potential for biocontrol, and infecting
aphids requires overcoming both aphid defenses and their defensive endosymbionts.

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## 32 Results

33 We present the *de novo* genome assemblies, detailed annotation, and comparative 34 analysis of two closely related parasitoid wasps that target pest aphids: Aphidius ervi 35 and Lysiphlebus fabarum (Hymenoptera: Braconidae: Aphidiinae). The genomes are 36 small (139 and 141 Mbp), highly syntenic, and the most AT-rich reported thus far for 37 any arthropod (GC content: 25.8% and 23.8%). This nucleotide bias is accompanied by 38 skewed codon usage, and is stronger in genes with adult-biased expression. AT-richness 39 may be the consequence of reduced genome size, a near absence of DNA methylation, 40 and age-specific energy demands. We identify expansions of F-box/Leucine-rich-repeat 41 proteins, suggesting that diversification in this gene family may be associated with their 42 broad host range or with countering defenses from aphids' endosymbionts. The 43 absence of some immune genes (Toll and Imd pathways) resembles similar losses in 44 their aphid hosts, highlighting the potential impact of symbiosis on both aphids and 45 their parasitoids.

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## 47 Conclusions

48	These findings are of fundamental interest for insect evolution and beyond. This will
49	provide a strong foundation for further functional studies including coevolution with
50	respect to their hosts, the basis of successful infection, and biocontrol. Both genomes
51	are available at https://bipaa.genouest.org.
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54	Keywords: Parasitoid wasp, aphid host, Aphidius ervi, Lysiphlebus fabarum, GC content,
55	de novo genome assembly, DNA methylation loss, chemosensory genes, venom
56	proteins, Toll and Imd pathways
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## 71 Background

72 Parasites are ubiquitously present across all of life (Poulin 2007; Windsor 1998). Their 73 negative impact on host fitness can impose strong selection on hosts to resist, tolerate, 74 or escape potential parasites. Parasitoids are a special group of parasites whose 75 successful reproduction is fatal to the host (Godfray 1994; Quicke 2014). The 76 overwhelming majority of parasitoid insects are hymenopterans that parasitize other 77 terrestrial arthropods, and they are estimated to comprise up to 75% of the species-78 rich insect order Hymenoptera (Forbes et al. 2018; Godfray 1994; Heraty 2009; 79 Pennacchio & Strand 2006). Parasitoid wasps target virtually all insects and 80 developmental stages (eggs, larvae, pupae, and adults), including other parasitoids 81 (Chen & van Achterberg 2018; Godfray 1994; Müller et al. 2004; Poelman et al. 2012). 82 Parasitoid radiations appear to have coincided with those of their hosts (Peters et al. 83 2017), and there is ample evidence that host-parasitoid relationships impose strong 84 reciprocal selection, promoting a dynamic process of antagonistic coevolution (Dupas 85 et al. 2003; Kraaijeveld et al. 1998; Vorburger & Perlman 2018).

86 Parasitoids of aphids play an economically important role in biological pest 87 control (Boivin et al. 2012; Heimpel & Mills 2017), and aphid-parasitoid interactions are 88 an excellent model to study antagonistic coevolution, specialization, and speciation 89 (Henter & Via 1995; Herzog et al. 2007). While parasitoids that target aphids have 90 evolved convergently several times, their largest radiation is found in the braconid 91 subfamily Aphidiinae, which contains at least 400 described species across 50 genera 92 (Chen & van Achterberg 2018; Shi & Chen 2005). As koinobiont parasitoids, their 93 development progresses initially in still living, feeding, and developing hosts, and ends 94 with the aphids' death and the emergence of adult parasitoids. Parasitoids increase

95 their success with a variety of strategies, including host choice (Chau & Mackauer 2000; 96 Łukasik et al. 2013), altering larval development timing (Martinez et al. 2016), injecting 97 venom during stinging and oviposition, and developing special cells called teratocytes 98 (Burke & Strand 2014; Colinet et al. 2014; Falabella et al. 2003; Poirié et al. 2014; Strand 99 2014). In response to strong selection imposed by parasitoids, aphids have evolved 100 numerous defenses, including behavioral strategies (Gross 1993), immune defenses 101 (Schmitz et al. 2012), and symbioses with heritable endosymbiotic bacteria whose 102 integrated phages can produce toxins to hinder parasitoid success (Oliver et al. 2010; 103 Oliver & Higashi 2018; Vorburger & Perlman 2018).

104 The parasitoid wasps Lysiphlebus fabarum and Aphidius ervi (Braconidae: 105 Aphidiinae) are closely related endoparasitoids (Figure 1). In the wild both species are 106 found infecting a wide range of aphid species although their host ranges differ, with A. 107 ervi more specialized on aphids in the Macrosiphini tribe and L. fabarum on the Aphidini 108 tribe (Kavallieratos et al. 2004; Monticelli et al. 2019). In both taxa, there is evidence 109 that parasitoid success is hindered by the presence of defensive symbionts in the aphid 110 haemocoel, including the bacteria Hamiltonella, Regiella, and Serratia (Oliver et al. 111 2003; Vorburger et al. 2010). Studies employing experimental evolution in both species 112 have shown that wild-caught populations can counter-adapt to cope with aphids and 113 the defenses of their endosymbionts, and that the coevolutionary relationships 114 between parasitoids and the aphids' symbionts likely fuel diversification of both 115 parasitoids and their hosts (Dennis et al. 2017; Dion et al. 2011; Rouchet & Vorburger 116 2014). While a number of parasitoid taxa are known to inject viruses and virus-like 117 particles into their hosts, there is thus far no evidence that this occurs in parasitoids 118 that target aphids; emerging studies have identified abundant RNA viruses in L.

*fabarum* (Lüthi *et al.* submitted; Obbard *et al.* in revision), but whether this impacts
their ability to parasitize is not yet fully understood.

121 These two closely related parasitoids differ in several important life history 122 traits, and are expected to have experienced different selective regimes as a result. 123 Aphidius ervi is has successfully been introduced widely (Nearctic, Neotropics) as a 124 biological control agent (far more than L. fabarum). Studies on both native and 125 introduced populations of A. ervi have shown ongoing evolutionary processes with 126 regard to host preferences, gene flow, and other life history components (Henry et al. 127 2008; Hufbauer et al. 2004; Zepeda-Paulo et al. 2015; Zepeda-Paulo et al. 2013). A. ervi 128 is known to reproduce only sexually, whereas *L. fabarum* is capable of both sexual and 129 asexual reproduction. In fact, wild *L. fabarum* populations are more commonly 130 composed of asexually reproducing (thelytokous) individuals (Sandrock et al. 2011). In 131 asexual populations, diploid *L. fabarum* females produce diploid female offspring via central fusion automixis (Belshaw & Quicke 2003). While they are genetically 132 133 differentiated, sexual and asexual populations appear to maintain gene flow and thus 134 both reproductive modes and genome-wide heterozygosity are maintained in the species as a whole (Mateo Leach et al. 2009; Sandrock et al. 2011; Sandrock & 135 136 Vorburger 2011). Aphidius. ervi and L. fabarum are also expected to have experienced 137 different selective regimes with regard to their cuticular hydrocarbon profiles and 138 chemosensory perception. Lysiphlebus target aphid species that are ant-tended, and 139 ants are known to prevent parasitoid attacks on "their" aphids (Rasekh et al. 2010). To 140 counter ant defenses, L. fabarum has evolved the ability to mimic the cuticular 141 hydrocarbon profile of the aphid hosts (Liepert & Dettner 1993, 1996). With this, they 142 are able to circumvent ant defenses and access this challenging ecological niche, from which they also benefit nutritionally; they are the only parasitoid species thus far
documented to behaviorally encourage aphid honeydew production and consume this
high-sugar reward (Rasekh *et al.* 2010; Völkl 1992; Völkl 1997).

146 We present here the genomes of A. ervi and L. fabarum, assembled de novo 147 using a hybrid sequencing approach. The two genomes are highly syntenic and strongly 148 biased towards AT nucleotides. We have examined GC content in the context of host 149 environment, nutrient limitation, and gene expression. By comparing these two 150 genomes we identify key functional specificities in genes underlying venom 151 composition, oxidative phosphorylation, cuticular hydrocarbon composition, and 152 chemosensory perception. In both species, we identify losses in key immune genes and 153 an apparent lack of key DNA methylation machinery. These are functionally important 154 traits associated with success infecting aphids and the evolution of related traits across 155 all of Hymenoptera.

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## 157 Results and Discussion

## 158 *Two* de novo *genome* assemblies

The genome assemblies for A. ervi and L. fabarum were constructed using hybrid 159 approaches that incorporated high-coverage short read (Illumina) and long-read (Pac 160 161 Bio) sequencing, but were assembled with different parameters (Supplementary Tables 162 1, 2). This produced two high quality genome assemblies (A. ervi N50 = 581kb, L. 163 fabarum N50 = 216kb) with similar total lengths (A. ervi: 139MB, L. fabarum: 141MB) but different ranges of scaffold-sizes (Table 1, Supplementary Table 3). These assembly 164 lengths are within previous estimates of 110-180Mbp for braconids, including A. ervi 165 166 (Ardila-Garcia et al. 2010; Hanrahan & Johnston 2011). Both assemblies are available in NCBI (SAMN13190903-4) and can be accessed via the BioInformatics Platform for
Agroecosystem Arthropods (BIPAA, https://bipaa.genouest.org), which contains the
full annotation reports, predicted genes, and can be searched via both keywords and
blast.

171 We constructed linkage groups for the *L. fabarum* scaffolds using phased SNPs 172 from the haploid (male) sons of a single female wasp from a sexually reproducing 173 population. This placed the 297 largest scaffolds (>50% of the nucleotides, 174 Supplementary Table 5, Supplementary Figure 1, Additional File 1) into the expected 175 six chromosomes (Belshaw & Quicke 2003). With this largely contiguous assembly, we 176 show that the two genomes are highly syntenic, with >60k links in alignments made by 177 NUCmer (Kurtz et al. 2004) and >350 large syntenic blocks that match the six L. fabarum 178 chromosomes to 28 A. ervi scaffolds (Supplementary Figures 2 and 3).

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	A. ervi	L. fabarum
Assembly statistics		
Total length (bp)	138,951,524	140,705,580
Longest scaffold (bp)	3,671,467	2,183,677
scaffolds	5,778	1,698
scaffolds $\geq$ 3,000 bp	1,503	1,698
N50 (bp)	581,355	216,143
GC %	25.8%	23.8%
Annotation statistics		
Exons	95,322	74,701
Introns	74,978	59,498
CDS	20,344	15,203
% genome covered by CDS	17.8%	14.9%
GC % in CDS	31.9%	29.8%
GC % of 3 <sup>rd</sup> position in CDS	15.5%	10.7%
CDS with transcriptomic support	77.8%	88.3%

180 Table 1: Assembly and draft annotation statistics

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183 Within the two assemblies, we used the Maker2 annotation pipeline to predict coding genes (CDS) for the two genomes, and these were functionally annotated 184 185 against the NCBI nr database (NCBI), matches to gene ontology (GO) terms, and 186 predictions for known protein motifs, signal peptides, and transmembrane domains 187 (Supplemental Table 6). In A. ervi there were 20,344 predicted genes comprising 188 27.8Mbp, while in *L. fabarum* there were 15,203 genes across 21.9 Mbp (Table 1). 189 These numbers are on par with those predicted in other hymenopteran genomes 190 (Table 2), and comparisons among taxa suggest that the lower number of predicted 191 genes in L. fabarum are more likely due to their loss than to a gene gain in A. ervi. 192 However, it is important to recognize that predictive annotation is imperfect and any 193 missing genes should be specifically screened with more rigorous methods. In both 194 species, there was high transcriptomic support for the predicted genes (77.8% in A. ervi 195 and 88.3% in *L. fabarum*). The two genome annotations appear to be largely complete; 196 at the nucleotide level, we could match 94.8% (A. ervi) and 76.3% (L. fabarum) of the 197 1,658 core orthologous BUSCO genes for Insecta in both species (Supplementary Table 198 4). Within the predicted genes, protein-level matches to the BUSCO genes were 199 improved in *L. fabarum* (95.9%) and slightly lower for *A. ervi* (93.7%). These numbers 200 suggest that low GC content did not negatively impact gene prediction (Supplementary 201 Table 4).

A survey of transposable Elements (TEs) identified a similar overall number of putative TE elements in the two assemblies (*A. ervi*: 67,695 and *L. fabarum*: 60,306, Supplementary Table 7). Despite this similarity, the overall genomic coverage by TEs is larger in *L. fabarum* (41%, 58 Mbp) than in A. *ervi* (22%, 31 Mbp) and they differ in the TE classes that they contain (Supplementary Table 7, Supplementary Figures 4, 5). The

207	spread of reported TE coverage in arthropods is quite large, even among Drosophila
208	species (ca. 2.7% - 25%, Drosophila 12 Genomes et al. 2007). Within parasitoids,
209	reported TE content also varies, and relatively low coverage in the parasitoid
210	Macrocentrus cingulum in comparison to Nasonia vitripennis (24.9% vs 40.6% Yin et al.
211	2018) was attributed the smaller genome size of <i>M. cinculum</i> (127.9Mbp and
212	295.7Mbp, respectively, Table 3). However, the variation we observe here suggests
213	that differences in predicted TE content may be evolutionary quite labile, even within
214	closely related species with the same genome size.

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Table 2: Assembly summary statistics compared to other parasitoid genomes. All species are from the family
 Braconidae, except for N. vitripennis (Pteromalidae). Protein counts from the NCBI genome deposition.

Parasitoid species	Assembly	Total Length (Mbp)	Scaffold Count	Scaffold N50 (bp)	Predicted genes (CDS)	GC (%)	NCBI BioProject
Aphidius ervi	A. ervi_v3	139.0	5,778	581,355	20,344	25.8	This paper
Lysiphlebus fabarum	L. fabarum_v1	140.7	1,698	216,143	15,203	23.8	This paper
Fopius arisanus	ASM80636v1	153.6	1,042	51,867	18,906	39.4	PRJNA258104 (Geib <i>et al.</i> 2017)
Diachasma alloeum	Dall1.0	388.8	3,968	44,932	19,692	39.1	PRJNA284396 (Tvedte <i>et al.</i> 2019)
Microplitis demolitor	Mdem 2	241.2	1,794	27,508	18,586	33.1	PRJNA251518 (Burke <i>et al.</i> 2018)
Cotesia vestalis	ASM95615v1	186.1	9,156	46,055	-	30.4	PRJNA271135
Macrocentrus cingulum	MCINOGS1.0	127.9	12,056	65,089	11,993	35.6	PRJNA361069 (Yin <i>et</i> <i>al.</i> 2018)
Nasonia vitripennis	Nvit_2.1	295.7	6,169	18,840	24,891	40.6	PRJNA13660 (Werren <i>et al.</i> 2010)

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## 219 GC content

The *L. fabarum* and *A. ervi* genomes are the most GC-poor of insect genomes sequenced to date (GC content: 25.8% and 23.8% for *A. ervi* and *L. fabarum*, respectively, Table 3, Supplementary Figure 6). This nucleotide bias is accompanied by strong codon bias in the predicted genes, meaning that within the possible codons for

each amino acid, the two genomes are almost universally skewed towards the codon(s)
with the lowest GC content (measured as Relative Synonymous Codon Usage, RSCU,
Figure 2). These patterns are much more extreme than RSCU found in other
hymenopterans, which are known to prefer codons that end in –A or –U (Behura &
Severson 2013). This codon bias has functional consequences; work in other taxa has
shown that codon usage is tied to both expression efficiency and mRNA stability
(Barahimipour *et al.* 2015).

231 Low GC content could be a consequence of the relatively small size of these 232 genomes. Genome size and GC content are positively correlated in a diverse set of taxa 233 including bacteria (Almpanis et al. 2018; McCutcheon et al. 2009), plants (Šmarda et al. 234 2014; Veleba et al. 2016), and vertebrates (Vinogradov 1998). This widespread pattern 235 may be driven by GC-rich repetitive elements that are more abundant in larger 236 genomes, stronger selection on thermal stability in larger genomes, or thermal stability associated with the environment (Šmarda et al. 2014; Vinogradov 1998). The apparent 237 238 lack of DNA methylation in this system may also contribute to low GC content (see 239 below and Bewick et al. 2017). Methylation is a stabilizing factor with regard to GC 240 content (Mugal et al. 2015), so its absence could relax selection on GC content and 241 allow it to decline. However, neither the absence of methylation nor codon bias are 242 unique to these taxa, suggesting that some additional selective factors or genetic drift 243 may have further shaped the composition of these two genomes.

We used two approaches to investigate whether environmental constraints could drive extremely low GC content, but found no evidence for such constraints. There is reason to expect that environment could contribute to the low GC content of these genomes; in taxa including bacteria (Foerstner *et al.* 2005) and plants (Šmarda *et* 

248 al. 2014) the environment has been shown to influence GC content via limitation in 249 elements including nitrogen. These two wasps parasitize aphids exclusively, and aphids 250 themselves have relatively low genome-wide GC content. This includes the pea aphid 251 (Acyrtosiphon pisum), which is a frequent host of A. ervi and also has notably low GC 252 content (29.8%, Li et al. 2019). This is not limited to A. pisum, with other aphid 253 genomes' GC content ranging between 26.8% - 30% (Additional File 2), perhaps related 254 to their high-sugar, low-nitrogen, sap diet. One way to explore the restrictions imposed 255 by nutrient limitation is to look at the expressed genes, since selective pressure should 256 be higher for genes that are more highly expressed (Ran & Higgs 2010; Seward & Kelly 257 2016). For our first test, we explored potential constraints in the most highly expressed 258 genes in both genomes. In both species, the most highly expressed 5% of genes had 259 higher GC content and higher nitrogen content, although the higher number of 260 nitrogen molecules in G's and C's means that these two measures cannot be entirely 261 disentangled (Additional File 3, Supplementary Figure 7). This is in line with 262 observations across many taxa, and with the idea that GC-rich mRNA has increased 263 expression via its stability and secondary structure (Kudla et al. 2009; Plotkin & Kudla 264 2011). For a second approach to examining constraints, we compared codon usage 265 between our genomes and taxa associated with this parasitoid-host-endosymbiont 266 system (Supplementary Table 8). We found no evidence of similarity in codon usage 267 (scaled as RSCU) nor in nitrogen content (scaled per amino acid) between parasitoids 268 and host aphids, the primary endosymbionts *Buchnera* nor, with the secondary 269 endosymbiont Hamiltonella (Supplementary Figures 8-10). Together, these tests do not 270 support environmental constraints as the driver of low GC content in these two 271 genomes.

272 In contrast, we did find evidence for reduced GC content in genes expressed at 273 different parasitoid life-history stages. We found higher GC content in larvae-biased 274 genes in *L. fabarum* (Figure 3). This was true when we compared the 10% most highly 275 expressed genes in adults (32.6% GC) and larvae (33.2%, p=1.2e-116, Figure 3, 276 Additional File 3), and this pattern holds even more strongly for genes that are 277 differentially expressed between adults (upregulated in adults: 28.7% GC) and larvae (upregulated in larvae: 30.7% GC, p=2.2e-80. Note that the most highly expressed 278 279 genes overlap partially with those that are differentially expressed, Additional File 3). 280 At the same time, we found no evidence that nitrogen content differs in either of these 281 comparisons (Figure 3). While the magnitude of these differences is not very large, 282 subtle differences in gene content are hypothesized to be the result of selection in 283 other systems (Acquisti et al. 2009). It seems plausible that GC content differences 284 among genes expressed at different life history stages could be selected in a process 285 analogous to the small changes in gene expression that are linked to large phenotypic 286 differences within and between species (Romero et al. 2012). One explanation for 287 lower GC content in adult-biased genes could be differences in energy demands and availability of resource across life stages. Given the extreme codon bias in these 288 289 genomes (Figure 2), using codons that match this bias is expected to be more efficient 290 and accurate, resulting in lower energy consumption and faster turnover (Chaney & 291 Clark 2015; Galtier et al. 2018; Kudla et al. 2006; Rao et al. 2013). Expressing AT-rich 292 genes is slightly more energy-efficient in itself, and this could favor otherwise neutral 293 mutations from GC to AT (Rocha & Danchin 2002). There is good motivation for adults to have a greater demand for energy efficiency. Adult parasitoids usually feed on 294 295 carbohydrate rich but protein and lipid poor resources like nectar, while performing

costly tasks including flying, mating, and laying eggs. Meanwhile, parasitoid larvae are
feeding on their aphid host's tissue, and likely benefit further from nutrients coming
from the aphids' endosymbionts, while their only task is to grow as fast as possible
(Cheng *et al.* 2011; Miao *et al.* 2004; Pennacchio *et al.* 1999).

300 This supports the idea that selection at the level of gene expression is shaping 301 the GC content of these genomes. Nonetheless, further work should more explicitly 302 test both nutrient limitation and how selective pressures differ across life-history 303 stages. While we do not have the power to test for GC-biased gene conversion with 304 two taxa, the even lower third position GC content (15.5% and 10.7%, Table 1) suggests 305 that this should be tested in relation to other parasitoids (Galtier et al. 2018). Further 306 explanations to be considered include effective population size, translational efficiency, 307 and mutational bias (Behura & Severson 2013; Bentele et al. 2013; Galtier et al. 2018). 308 Altogether, these patterns raise important questions about how codon biases impact 309 genome content, and whether synonymous mutations are always functionally neutral 310 (Plotkin & Kudla 2011; Powell & Moriyama 1997).

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### 312 Orphan genes in the assembly

To examine genes that may underlie novel functional adaptation, we identified sequences that are unique within the predicted genes in the *A. ervi* and *L. fabarum* genomes. We defined orphan genes as predicted genes with transcriptomic support and with no identifiable homology based on searches against the NCBI *nr, nt,* and Swissprot databases. With this, we identified 2,568 (*A. ervi*, Additional File 4) and 968 (*L. fabarum*, Additional File 5) putative orphans (Supplementary Table 9). The evolutionary origin of these orphan genes is not known (Gold *et al.* 2018; Van Oss &

320 Carvunis 2019), but their retention or evolution could be important to understanding
321 specific functions or traits in these taxa. The higher number of orphan genes in *A. ervi*322 partially explains the absolute difference in the number of annotated genes between
323 both taxa.

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### 325 Gene family expansions

326 To examine gene families that may have undergone expansions in association with 327 functional divergence and specialization, we identified groups of orthologous genes 328 that have increased and decreased in size in the two genomes, relative to one another. 329 We identified these species-specific gene-family expansions using the OMA standalone 330 package (Altenhoff et al. 2018). OMA predicted 8,817 OMA groups (strict 1:1 orthologs) 331 and 8,578 HOGs (Hierarchical Ortholog Groups, Additional File 6). Putative gene-family 332 expansions would be found in the predicted HOGs, because they are calculated to allow 333 for >1 member per species. Among these, there were more groups in which A. ervi 334 possessed more genes than L. fabarum (865 groups with more genes in A. ervi, 223 335 with more in *L. fabarum*, Supplementary Figure 11, Additional File 6). To examine only 336 the largest gene-family expansions, we looked further at the HOGs containing >20 337 genes (10 HOG groups, Supplementary Figure 12). Strikingly, the four largest 338 expansions were more abundant in A. ervi and were all identified as F-box proteins/ 339 Leucine-rich-repeat proteins (LRR, total: 232 genes in A. ervi and 68 in L. fabarum, 340 Supplementary Figure 12, Additional File 6). This signature of expansion does not 341 appear to be due to fragmentation in the A. ervi assembly: the size of scaffolds 342 containing LRRs is on average larger in A. ervi than in L. fabarum (Welch two-sampled 343 t-test, p=0.001, Supplementary Figure 13).

344 The LRRs are a broad class of proteins associated with protein-protein 345 interactions, including putative venom components in these parasitoids (Colinet et al. 346 2014). LRRs belong to a larger category of leucine rich repeat pattern recognition 347 receptor proteins, which are an important component of innate immunity and cell-348 surface recognition of bacterial intruders and include toll-like receptors in insects 349 (Soanes & Talbot 2010; Takeda & Akira 2005). While the functions of these proteins are 350 diverse, expansion in F-box/LRR proteins has been shown to have specific function in 351 immunity in parasitic insects. In the Hessian fly (Mayetiola destructor), fly-encoded F-352 box/LRR proteins bind with plant-encoded proteins to form a complex that blocks the 353 plant's immune defenses against the parasitic fly (Zhao et al. 2015). Thus, we 354 hypothesize that this class of proteins has expanded in these parasitoids in relation to 355 recognizing the diverse bacterial defenses of their aphid hosts. Under this hypothesis, 356 we argue that expansion of F-box/LRR proteins contributes to the broad host 357 recognition in both species, and that their greater abundance in A. ervi may be 358 associated with a recent arms race with respect to the immune defenses and protective 359 endosymbionts of their host aphids.

360 The six largest gene families that were expanded in *L. fabarum*, relative to *A*. 361 ervi, were less consistently annotated. Interestingly, they contained two different 362 histone proteins: Histone H2B and H2A (Supplementary Figure 12). All eukaryotic 363 genomes examined to date contain multiple histone genes for the same histone 364 variants found in humans (e.g. 22 genes for H2B or 16 genes for H2A in humans, Singh 365 et al. 2018), and it has recently been suggested that these histone variants are not 366 functionally equivalent but rather play a role in chromatin regulation (Singh et al. 2018). 367 Hence, these variants could also play a role in several L. fabarum specific traits,

including the switch from sexual to asexual reproduction (thelytoky); in mammals, sex
determination has been linked to regulation via histone modification (Kuroki *et al.*2013).

371

### 372 Venom proteins

373 Venom injected at oviposition is crucial for successful reproduction in most parasitoid wasp species (Moreau & Asgari 2015; Poirié et al. 2014). The venom of A. ervi was 374 375 previously analyzed using a combined transcriptomic and proteomic approach (Colinet 376 et al. 2014), and we applied similar methods here to compare the venom composition 377 in *L. fabarum*. The venom gland in *L. fabarum* is morphologically different from *A. ervi* (Supplementary Figure 14). A total of 35 L. fabarum proteins were identified as putative 378 379 venom proteins using 1D gel electrophoresis and mass spectrometry, combined with 380 transcriptomic and the genome data (Supplementary Figure 15, Additional File 7, 381 Dennis et al. 2017). These putative venom proteins were identified based on predicted 382 secretion (for complete sequences) and the absence of a match to typical cellular 383 proteins (e.g. actin, myosin). To match the analysis between the two taxa, the previous 384 A. ervi venom data (Colinet et al. 2014) was analyzed using the same criteria as L. 385 fabarum. This identified 32 putative venom proteins in A. ervi (Additional File 7).

Although these two species differ in their host range (Kavallieratos *et al.* 2004), comparison of venom proteins between species revealed that more than 50% of the proteins are shared between species (Figure 4A and Additional File 7), corresponding to more than 70% of the putative function categories that were predicted (Figure 4B and Additional File 7). Among venom proteins shared between both parasitoids, a gamma glutamyl transpeptidase (GGT1) is the most abundant protein in the venom of

392 both A. ervi (Colinet et al. 2014) and L. fabarum (Additional File 7). This protein has 393 been suggested to be involved in the castration of the aphid host after parasitism 394 (Falabella et al. 2007). As previously reported for A. ervi (Colinet et al. 2014), a second 395 GGT venom protein (GGT2) containing mutations in the active site was also found in 396 the venom of *L. fabarum* (Supplementary Figure 16, 17). Phylogenetic analysis (Figure 397 5) revealed that the A. ervi and L. fabarum GGT venom proteins occur in a single clade 398 in which GGT1 venom proteins group separately from GGT2 venom proteins, thus 399 suggesting that they originated from a duplication that occurred prior to the split from 400 their most recent common ancestor. As previously shown for A. ervi, the GGT venom 401 proteins of A. ervi and L. fabarum are found in one of the three clades described for the non-venomous hymenopteran GGT proteins (clade "A", Figure 5 and Colinet et al. 402 403 2014). Within this clade, venomous and non-venomous GGT proteins had a similar exon 404 structure, except for exon 1 that corresponds to the signal peptide only present in 405 venomous GGT proteins (Supplementary Figure 17). Aphidius ervi and L. fabarum 406 venomous GGT proteins thus probably result from a single imperfect duplication of the 407 non-venomous GGT gene belonging to clade A in their common ancestor, followed by recruitment of the signal peptide coding sequence. This first imperfect duplication 408 409 event would then have been followed by a second duplication of the newly recruited 410 venomous GGT gene before the separation of both species.

The presence of truncated *LRR* proteins was previously reported in venom of *A*. *ervi* (Colinet et al. 2014) and other Braconidae (Mathé-Hubert et al. 2016) that likely interfere with the host immune response. Several *LRR* proteins were found in the venom of *L. fabarum* as well, however these results should be interpreted with caution since the sequences were incomplete and the presence of a signal peptide could not be confirmed (Additional File 7). Moreover, these putative venom proteins were only
identified from transcriptomic data of the venom apparatus and we could not find any
corresponding annotated gene in the genome. This supports the idea that gene-family
expansions in putative F-box/*LRR* proteins (discussed above) are not related to venom
production.

421 Approximately 50% of the identified venom proteins were unique to either A. 422 ervi or L. fabarum, and these could be related to their differing host ranges (Additional 423 File 7). However, most of these proteins had no predicted function, making it difficult 424 to hypothesize their possible role in parasitism success. Among the venom proteins 425 with a predicted function, an apolipophorin was found in the venom of *L. fabarum* but 426 not in A. ervi. Apolipohorin is an insect-specific apolipoprotein involved in lipid 427 transport and innate immunity that is not commonly found in venoms. Among 428 parasitoid wasps, apolipophorin has been described in the venom of the ichneumonid 429 Hyposoter didymator (Dorémus et al. 2013) and the encyrtid Diversinervus elegans (Liu 430 et al. 2017), but its function is yet to be deciphered. Apolipophorin is also present in 431 low abundance in honeybee venom where it could have antibacterial activity (Kim & Jin 432 2015; Van Vaerenbergh et al. 2014). Lastly, we could not find L. fabarum homologs for 433 any of the three secreted cystein-rich toxin-like peptides that are highly expressed in 434 the A. ervi venom apparatus (Additional File 7). However, this may not be definitive 435 since the search for similarities in the genome is complicated by the small size of these 436 toxin-like sequences.

437

438

439

Category	A. ervi	L. fabarum
Venom proteins	32	35
Desaturases*	16	15
Immune genes <sup>+</sup>	216	216
Osiris genes	21	25
Mitochondrial Oxidative Phosphorylation System (OXPHOS)**	75	74
Chemosensory group		
Chemosensory: Odorant receptors (ORs)	228	156
Chemosensory: Ionotropic chemosensory receptors (IRs)	42	40
Chemosensory: Odorant-binding proteins (OBPs)	14	14
Chemosensory: Chemosensory proteins (CSPs)	11	13
Sex determination group		
Sex determination: Core (transformer, doublesex)	4	3
Sex determination: Related genes	6	5
DNA methylation genes	2	2
TOTALS	667	598

#### 440 Table 3: Summary of manual curations of select gene families in the two parasitoid genomes

441 442

- \*Note 1: Includes genes that are partial, ambiguous, or potential pseudogenes †Note2: although the same number, the set of immune genes is not identical in the two genomes.
- 443
- 444

#### 445 Key gene families

446 We manually annotated more than 1,000 genes (667 for A. ervi and 598 for L. fabarum;

447 Table 3) using Apollo, hosted on the BIPAA website (Dunn *et al.* 2019;

448 https://bipaa.genouest.org ; Lee et al. 2013) to confirm and improve the results of the

449 machine annotation. This is especially important for large gene families, which are

450 usually poorly annotated by automatic prediction (Robertson et al. 2018); since such

451 gene families potentially underlie key adaptive differences between the two

452 parasitoids, accurate annotation is needed.

453

#### 454 Desaturases

455 Desaturases are an important gene family that introduce carbon-carbon double bonds 456 in fatty acyl chains in insects (Los & Murata 1998; Sperling et al. 2003). While these

function broadly across taxa, a subset of these genes (specifically acyl-CoA desaturases) 457 458 have been implicated in insect chemical recognition for roles including alkene 459 production and modification of fatty acids (Helmkampf et al. 2015). This gene family is 460 particularly interesting because it has been shown that Lysiphlebus cardui, a close 461 relative of *L. fabarum*, have no unsaturated cuticular hydrocarbons, just as is seen in its 462 aphid host. This allows the parasitoid to go undetected in aphid colonies that are anttended and therefore better parasitize them (Liepert & Dettner 1996). We confirmed 463 464 that the same is true for *L. fabarum*; its CHC profile is dominated by saturated 465 hydrocarbons (alkanes), contains only trace alkenes, and is completely lacking dienes (Supplementary Figure 18, 20). In contrast, A. ervi females produce a large amount of 466 unsaturated hydrocarbons, with a significant amount of alkenes and alkadiens in their 467 468 CHC profiles (app. 70% of the CHC profile are alkenes/alkadienes, Supplementary 469 Figure 19, 20).

470 The loss of one annotated desaturase gene in L. fabarum compared to A. ervi 471 (Table 3) might explain these differences in the composition of their CHC profiles, 472 especially their apparent inability to synthesize dienes. We also note there is little 473 evidence that members of this gene family are clustered in the genome (just three and 474 two desaturase genes in the same scaffolds of *A. ervi* and *L. fabarum*, respectively). 475 Further investigations should verify this loss in *L. fabarum*, identify the ortholog of the 476 missing copy in A. ervi, and test if this potential lost desaturase gene in L. fabarum is 477 involved in the generation of unsaturated CHCs in A. ervi. This would determine if this 478 loss is a key adaptation for mimicry of their aphid hosts' cuticular hydrocarbon profiles 479 in *L. fabarum*.

480

## 481 Immune genes

482 We searched for immune genes in the two genomes based on a list of 367 immunity 483 related genes, collected primarily from the Drosophila literature (Additional File 8). 484 Using blast-based searches, 204 of these genes (59%) were found and annotated in 485 both species. Six were present in only the A. ervi genome and six in only the L. fabarum 486 genome. We compared these with the immune genes used to define the main Drosophila immune pathways (Toll, Imd, and JAK-STAT, Supplementary Table 10) and 487 488 conserved in a large number of insect species (Buchon et al. 2014; Charroux & Royet 2010; Lemaitre & Hoffman 2007). Among these genes there are several well 489 490 characterized pathways. The D. melanogaster Toll pathway is essential for the response 491 to fungi and Gram-positive bacteria (Valanne et al. 2011). It was initially identified as a 492 developmental pathway acting via the nuclear factor kappa B (NF- $\kappa$ B). The Imd/NF-493 kappa-B pathway is pivotal in the humoral and epithelial immune response to Gram-494 negative bacteria. Signaling through *imd* (a death domain protein) ultimately activates 495 the transcription of specific antimicrobial peptides (AMPs, Myllymäki et al. 2014). The 496 JAK-STAT pathway is involved in the humoral and cellular immune response (Morin-Poulard *et al.* 2013). It is activated after a cytokine-like protein called unpaired (*upd*) 497 498 binds to its receptor Domeless (Dome). Activated JAK phosphorylates STAT molecules 499 that translocate into the nucleus, where they bind the promoters of target genes.

In the genome of both wasps, many genes encoding proteins of the Imd and Toll pathways were absent, such as upstream GNBPs (Gram Negative Binding Proteins) and PGRPs (Peptidoglycan Recognition Proteins) and downstream AMPs (Supplementary Table 10, Supplementary Figure 21, Additional File 8). While none of these genes were found in *L. fabarum*, one PGRP related to PGRP-SD, involved in the

505 response to Gram-positive bacteria (Bischoff et al. 2004), and one defensin-related 506 gene were found in A. ervi. The imd gene was also absent in in both wasps; this is 507 noteworthy because *imd* has been present in other hymenopteran genomes analyzed 508 to date. Strikingly, all of the Imd pathway genes, including GNBP- and PGRP-encoding 509 genes, imd, FADD, Dredd and Relish are lacking in aphid genomes (A. pisum, A. gossypii 510 and *D. noxia*, via AphidBase (Legeai et al. 2010) and Gerardo et al (2010)), and *imd* is 511 absent in A. glycines, M. persicae, M. cerisae, R. padi genomes, some of which are hosts 512 for A. ervi and L. fabarum (Kavallieratos et al. 2004). The lack of an Imd pathway in 513 aphids is suggested to be an adaptation to tolerate the obligate bacterial symbiont, 514 Buchnera aphidicola, as well as their facultative endosymbionts that are gram-negative 515 gamma-proteobacteria (e.g. Hamiltonella defensa). These facultative symbionts exhibit 516 defensive activities against microbial pathogens and insect parasitoids (Guo et al. 2017; 517 Leclair et al. 2016; Oliver et al. 2010; Scarborough et al. 2005) and may at least partially 518 compensate for the host aphids innate immune functions. Recent data also suggest 519 that cross-talk occurs between the Imd and Toll pathways to target wider and 520 overlapping arrays of microbes (Nishide et al. 2019). Whether a similar cross-talk occurs 521 in these two Aphidiidae (A. ervi and L. fabarum) needs further study.

522 Overall, our results suggest convergent evolution of loss in immunity genes, and 523 possibly function, between these parasitoids and their aphid hosts. One reason might 524 be that during the early stages of development, parasitoids need host symbionts to 525 supply their basic nutrients, and thus an immune response from the parasitoid larvae 526 might impair this function. Alternatively, but not exclusively, mounting an immune 527 response against bacteria by the parasitoid larvae may be energetically costly and 528 divert resources from its development. This idea of energy conservation would be

529 especially relevant if the GC-loss discussed above is a mechanism to conserve 530 resources. In both cases, the immune response will be costly for the parasitoid. Further 531 work is needed to address whether other unrelated aphid parasitoids are lacking *imd*, 532 upstream activators, and downstream effectors of the immune pathways (a 533 preliminary blast search suggests that *imd* is present in the Aphelinidae Aphelinus 534 abdominalis). This impaired immunity might lead to a decrease in both wasps' 535 responses to pathogenic bacteria, or they may use other defensive components to fight 536 bacterial infections (perhaps some in common with aphids) that await to be discovered. 537 For example, in *L. fabarum*, recent transcriptomic work has shown that detoxifying 538 genes may be a key component of parasitoid success (Dennis et al. in revision), and 539 these could play a role in immunity.

540

## 541 Osiris genes

The Osiris genes are an insect-specific gene family that underwent multiple tandem duplications early in insect evolution. These genes are essential for proper embryogenesis (Smoyer *et al.* 2003) and pupation (Andrade López *et al.* 2017; Schmitt-Engel *et al.* 2015), and are also tied to immune and toxin-related responses (e.g. Andrade López *et al.* 2017; Greenwood *et al.* 2017) and developmental polyphenism (Smith *et al.* 2018; Vilcinskas & Vogel 2016).

We found 21 and 25 putative Osiris genes in the *A. ervi* and *L. fabarum* genomes, respectively (Supplementary Tables 11, 12). In insects with well assembled genomes, there is a consistent synteny of approximately 20 Osiris genes; this cluster usually occurs in a ~150kbp stretch and gene synteny is conserved in all known Hymenoptera genomes (Supplementary Figure 22). The Osiris cluster is largely devoid

of non-Osiris genes in most of the Hymenoptera, but the assemblies of *A. ervi* and *L. fabarum* suggest that if the cluster is actually syntenic in these species, there are interspersed non-Osiris genes (those are black boxes in Supplementary Figures 23 and 24).

557 In support of their role in defense (especially metabolism of xenobiotics and immunity), these genes were much more highly expressed in larvae than in adults 558 (Supplementary Table 12). We hypothesize that their upregulation in larvae is an 559 560 adaptive response to living within a host. Because of the available transcriptomic data, 561 we could only make this comparison in *L. fabarum*. Here, 19 of the 26 annotated Osiris 562 genes were significantly differentially expressed in larvae over adults (Supplementary 563 Table 12, Additional File 9). In both species, transcription in adults was very low, with 564 fewer than 10 raw reads per cDNA library sequenced, and most often less than one 565 read per library.

566

567 OXPHOS

568 In most eukaryotes, mitochondria provide the majority of cellular energy (in the form of adenosine triphosphate, ATP) through the oxidative phosphorylation (OXPHOS) 569 570 pathway. OXPHOS genes are an essential component of energy production, and have 571 increased in Hymenoptera relative to other insect orders (Li et al. 2017). We identified 572 69 out of 71 core OXPHOS genes in both genomes, and identified five putative 573 duplication events that are apparently not assembly errors (Supplementary Table 13, 574 Additional File 10). The gene sets of *A. ervi* and *L. fabarum* contained the same genes 575 and the same genes were duplicated in each, implying duplication events occurred 576 prior to the split from their most recent common ancestor. One of these duplicated 577 genes appears to be duplicated again in *A. ervi*, or the other copy has been lost in *L*.

578 fabarum.

579

### 580 Chemosensory genes

581 Genes underlying chemosensory reception play important roles in parasitoid mate and 582 host localization (Comeault et al. 2017; Nouhaud et al. 2018). Several classes of 583 chemosensory genes were annotated separately (Table 4): odorant receptors (ORs) are 584 known to detect volatile molecules, odorant-binding proteins (OBPs) and 585 chemosensory proteins (CSPs) are possible carriers of chemical molecules to sensory 586 neurons, and ionotropic receptors (IRs) are involved in both odorant and gustatory 587 molecule reception. With these manual annotations, further studies can now be made 588 with respect to life history characters including reproductive mode, specialization on 589 aphid hosts, and mimicry.

590

## 591 Chemosensory: Soluble proteins (OBPs and CSPs)

592 Hymenoptera have a wide range of known OBP genes, with up to 90 in *N. vitripenis* (Vieira et al. 2012). However, the numbers of these genes appear to be similar across 593 594 parasitic wasps, with 14 in both species studied here and 15 recently described in D. 595 alloeum (Tvedte et al. 2019). Similarly, CSP numbers are in the same range within 596 parasitic wasps (11 and 13 copies here, Table 4). Interestingly, two CSP sequences (one 597 in A. ervi and one in L. fabarum) did not have the conserved cysteine motif, 598 characteristic of this gene family. So although they were annotated here, further work 599 should investigate if and how these genes function.

600

## 601 <u>Chemosensory: Odorant receptors (ORs)</u>

602	In total, we annotated 228 putative ORs in <i>A. ervi</i> and 156 in <i>L. fabarum</i> (Table 4). This
603	is within the range of OR numbers annotated in other hymenopteran parasitoids,
604	including: 79 in <i>M. cingulum</i> (Ahmed <i>et al.</i> 2016), 225 in <i>N. vitripennis</i> (Robertson <i>et</i>
605	al. 2010), and 187 in D. alloeum (Tvedte et al. 2019). Interestingly, we annotated a
606	larger set of ORs in A. ervi than in L. fabarum. One explanation is that A. ervi generally
607	has more annotated genes than <i>L. fabarum,</i> and whatever broad pattern underlies
608	the reduction in the gene repertoire of <i>L. fabarum</i> also affected OR genes. One
609	functional explanations for a lower number of OR genes in <i>L. fabarum</i> is that the <i>A.</i>
610	ervi strain sequenced of was derived from several field strains that parasitized
611	different hosts on different host plants, and the ability to parasitize a broader host
612	range could select for more OR genes (Monticelli et al. 2019).
613	
614	Chemosensory: Ionotropic chemosensory receptors (IRs)
615	In total, we annotated 38 putative IRs in <i>A. ervi</i> and 37 in <i>L. fabarum</i> (Table 4). Three
616	putative co-receptors (IR 8a, IR 25a and IR 76b) were annotated both species, one of
617	which (IR 76b) was duplicated in <i>A. ervi</i> . This bring the total for the IR functional group
619	to 12 and 10 genes for A any and L fabarum respectively. This is within the range of

618 to 42 and 40 genes for *A. ervi* and *L. fabarum*, respectively. This is within the range of

619 IRs known from other parasitoid wasps such as *Aphidius gifuensis* (23 IRs identified in

620 antennal transcriptome, Braconidae, Kang et al. 2017), D. alloeum (51 IRs, Braconidae,

- 621 Tvedte *et al.* 2019) and *N. vitripennis* (47 IRs, Pteromalidae, Robertson *et al.* 2010). A
- 622 phylogenetic analysis of these genes showed a deeply rooted expansion in the IR genes
- 623 (Supplementary Figure 25). Thus, in contrast to the expansion usually observed in
- 624 hymenopteran ORs compared to other insect orders, IRs have not undergone major

625 expansions in parasitic wasps, which is generally the case for a majority of insects with

626 the exception of Blattodea (Harrison *et al.* 2018)

627

### 628 Sex determination

The core sex determination genes (transformer, doublesex) are conserved in both 629 species (Supplementary Table 14, Additional File 11). Notably, A. ervi possesses a 630 631 putative *transformer* duplication. This scaffold carrying the duplication (scaffold2824) 632 is only fragmentary, but a transformer duplicate has also been detected in the 633 transcriptome of a member of the A. colemani species complex, suggesting a conserved 634 presence within the genus (Peters et al. 2017). In A. ervi, transformer appears to have 635 an internal repeat of the CAM-domain, as is seen in the genus Asobara (Geuverink et 636 al. 2018). In contrast, there is no evidence of duplication in sex determination genes in 637 L. fabarum. This supports the idea that complementary sex determination (CSD) in 638 sexually reproducing *L. fabarum* populations is based on up-stream cues that differ 639 from those known in other CSD species (Matthey-Doret et al. 2019), whereas the CSD locus known from other hymenopterans locus is a paralog of transformer (Heimpel & 640 641 de Boer 2007).

In addition to the core sex determination genes, we identified homologs of several genes related to sex determination (Supplementary Table 15). We identified *fruitless* in both genomes, which is associated with sex-specific behavior in taxa including *Drosophila* (Yamamoto 2008). Both genomes also have homologs of *sex-lethal* which is the main determinant of sex in *Drosophila* (Bell *et al.* 1988). *Drosophila* has two homologs of this gene, and the single version in Hymenoptera may have more in common with the non-sex-lethal copy, called *sister-of-sex-lethal*. We identified 649 homologs of the gene CWC22, including a duplication in A. ervi; this duplication is 650 interesting because a duplicated copy of CWC22 is the primary signal of sex 651 determination in the house fly Musca domestica (Sharma et al. 2017). Lastly, there was 652 a duplication of *RBP1* in both genomes. The duplication of *RBP1* is not restricted to 653 these species, nor is the duplications of CWC22, which appears sporadically in 654 Braconidae. Together, these annotations add to our growing knowledge of duplications 655 of these genes, and provide possibilities for further examinations of the role of 656 duplications and specialization in association with sex determination.

657

## 658 DNA Methylation genes

659 DNA methyltransferase genes are thought to be responsible for the generation and 660 maintenance of DNA methylation. In general, DNA methyltransferase 3 (DNMT3) 661 introduces *de novo* DNA methylation sites and DNA methyltransferase 1 (*DNMT1*) 662 maintains and is essential for DNA methylation (Jeltsch & Jurkowska 2014; Provataris et al. 2018). A third gene, EEF1AKMT1 (formerly known as DNMT2), was once thought 663 to act to methylate DNA but is now understood to methylate tRNA (Provataris et al. 664 665 2018). In both A. ervi and L. fabarum, we successfully identified homologs DNMT3 and 666 EEF1AKMT1. In contrast, DNMT1 was not detected in either species (Table 4, Supplementary Table 16). This adds to growing evidence that these genes are not 667 conserved across family Braconidae, as DNMT1 appears to be absent in several other 668 669 braconid genera, including Asobara tabida, A. japonica, Cotesia sp., and F. arisanus 670 (Bewick et al. 2017; Geuverink 2017). However, DNMT1 is present in some braconids, 671 including *M. demolitor*, and outside of Braconidae these genes are otherwise strongly

672 conserved across insects. In contrast, DNMT3, present here, is more often lost in 673 insects (Provataris *et al.* 2018).

674 This absence of DNMT1 helps explains previous estimates of very low DNA methylation in A. ervi (0.5%, Bewick et al. 2017). We confirmed these low levels of 675 676 methylation in A. ervi by mapping this previously generated bisulfite sequencing data (Bewick et al. 2017) to our genome assembly. We aligned >80% of their data (total 677 94.5Mbp, 625,765 reads). The sequence coverage of this mapped data was low: only 678 679 63,554 methylation-available cytosines were covered and only 1,216 were represented 680 by two or more mapped reads. Nonetheless, of these mapped cytosines, the vast 681 majority (63,409) were never methylated, just 143 sites were always methylated, and 682 two were variably methylated. Methylation-available cytosine classes were roughly 683 equally distributed among three cytosine classes (CG: 0.154%, CHG: 0,179%, and CHH: 684 0.201%). This methylation rate is less than the 0.5% estimated by Bewick (2017) and 685 confirms a near absence of DNA methylation in A. ervi. Given the parallel absence of 686 DNMT1 in *L. fabarum*, it seems likely that both species sequenced here may have very low levels of DNA methylation, and that this is not a significant mechanism in these 687 688 species.

This stark reduction in DNA methylation is interesting, given that epigenetic mechanisms are likely important to insect defenses, including possible responses to host endosymbionts (Huang *et al.* 2019; Vilcinskas 2016, 2017). As with the immune pathways discussed above, this could reflect a loss that is adaptive to developing within endosymbiont-protected hosts. It is also interesting that while one epigenetic mechanism seems to be absent in both *A. ervi* and *L. fabarum*, we see an increase in histone variants in *L. fabarum* (based on the OMA analysis of gene family expansion),

- and these histones could function in gene regulation. However, whether there is a
- 697 functional or causal link between these two observations is yet to be tested.

## 698699 Table 4: Summary of annotation of putative DNA methylation genes

Species	Gene	Scaffold	e-value (Nasonia)	
A. ervi		scaffold94	1.00E-66	
<i>L. fabarum</i> EEF1AKMT1 homolog		tig00000449	5.00E-63	
A. ervi	DNA	scaffold45	5.00E-138	
<i>L. fabarum</i> methyltransferase 3		tig00002022	9.00E-117	
A. ervi	DNA	no homolog detected		
L. fabarum methyltransferase 1		no homolog detected		

700

701

## 702 Conclusions

703 These two genomes have provided insight into adaptive evolution in parasitoids that 704 infect aphids. Both genomes are extremely GC-poor, and their extreme codon bias 705 provides an excellent system for examining the chemical biases and selective forces 706 that may overshadow molecular evolution in eukaryotes. We have also highlighted 707 several groups of genes that are key to functional evolution across insects, including 708 venom, sex determination, response to bacterial infection (F-box/LRR proteins), and 709 near absence of DNA methylation. Moreover, the absence of certain immune genes 710 (e.g. from the Imd and Toll pathways) in these two species is similar to losses in host 711 aphids, and raises intriguing questions related to the effects of aphids' symbiosis on 712 both aphid and parasitoid genomics.

Parasitoid wasps provide an excellent model for studying applied and basic biological questions, including host range (specialist vs generalist), reproductive mode (sexual vs asexual), antagonistic coevolution, genome evolution, and epigenetic regulation, to mention just a few. Our new genomic resources will open the way for a broad set of future research, including work to understand host specialization, adaptive changes associated with climate, and the potential loss of diapause in *A. ervi* (Tougeron *et al.* 2019; Tougeron *et al.* 2017). Lastly, the genomes of these two non-social Hymenoptera provide a valuable comparison for understanding processes specific to social insects with complex caste structure, and are a first but essential step to better understand the genetic architecture and evolution of traits that are important for a parasitic life style and their use in biological control.

- 724
- 725
- 726 Methods
- 727 \*More complete methods are available in the Supplementary Material
- 728 Insect collection and origin

729 Aphidius ervi

730 Aphidius ervi samples used for whole-genome sequencing came from two different, 731 sexually reproducing, isofemale lines established from parasitized aphids (recognizable 732 as mummies) from fields of cereals and legumes in two different geographic zones in Chile: Region de Los Rios (S 39° 51′, W 73° 7′) and Region del Maule (S 35° 24′, W 71° 733 734 40<sup>'</sup>). Mummies (parasitized aphids) of *Sitobion avenae* aphids were sampled on wheat 735 (Triticum aestivum L.) while mummies of Acyrtosiphon pisum aphids were sampled on 736 Pisum sativum L. (pea aphid race). Aphid mummies were isolated in petri dishes until 737 adult parasitoids emerged. These two parasitoid lineages were separated in two cages 738 with hosts ad libitum and were propagated for approximately 75 generations under 739 controlled conditions as described elsewhere (Ballesteros et al. 2017; Sepúlveda et al. 740 2016). A further reduction of genetic variation was accomplished by establishing two

741 isofemale A. ervi lines, which were maintained as described previously and propagated 742 for approximately 10 generations before adult parasitoids (male and female) were 743 collected live and stored in 1.5 ml centrifuge tubes containing ethanol (95%) at -20°C. 744 Aphidius ervi samples used for CHC analysis (below) were purchased from Katz 745 Biotech AG (Baruth, Germany). Species identification was confirmed with COI 746 barcoding following Hebert et al. (2003). Wasps sacrificed for CHC analysis were 747 sampled from the first generation reared in the lab on Acyrtosiphon pisum strain LL01 748 (Peccoud et al. 2009), which were mass-reared on Vicia faba cv. Dreifach Weisse.

749

750 Lysiphlebus fabarum

751 *Lysiphlebus fabarum* samples used for whole-genome sequencing came from a single, 752 asexually reproducing, isofemale line (IL-07-64). This lineage was first collected in 753 September 2007 from Wildberg, Zürich, Switzerland as mummies of the aphid Aphis 754 fabae fabae, collected from the host plant *Chenopodium album*. In the lab, parasitoids 755 were reared on A. f. fabae raised on broad bean plants (Vicia faba) under controlled 756 conditions [16 h light: 8 h dark, 20°C] until sampling in September 2013, or 757 approximately 150 generations. Every lab generation was founded by ca. 10 individuals 758 that were transferred to fresh host plants containing wasp-naïve aphids. Approximately 759 700 individuals were collected for whole-genome sequencing from a single generation 760 in December 2013 and flash frozen at -80°C. To avoid sequencing non-wasp DNA, 761 samples were sorted over dry ice to remove any contaminating host aphid or plant 762 material.

For linkage group construction, separate *L. fabarum* collections were made
from a sexually reproducing lineage. Here, we collected all sons produced by a single

virgin female, sampled from the control lineage in a recently employed evolution experiment (H-lineage; Dennis *et al.* 2017). Wasps were stored on ethanol until RADseq library construction. Lastly, a third population was sampled for the proteomic analysis of the venom-apparatus (below); these females came from the geneticallydiverse starting population used to found the evolution experiment of Dennis *et al.* (2017), and were sampled in December 2014.

771

## 772 DNA extraction and library preparation

773 Aphidius ervi

774 DNA was extracted from adult haploid males of A. ervi in seven sub-samples (ca. 120 775 males each), reared in *S. avenae*. Total DNA was extracted using the DNEasy Plant Mini 776 Kit (QIAGEN) following the manufacturer's instructions. DNA was quantified by 777 spectrophotometry (Epoch Microplate Spectrophotometer, Biotek) and fluorometry 778 (Qubit 3.0; Qubit DNA High sensitivity Assay Kit, Invitrogen), and quality was assessed 779 using 1% agarose gel electrophoresis. DNA samples were sent on dry ice to MACROGEN 780 (Seoul, South Korea) and were used to produce Illumina paired-end (PE) and mate-pair 781 (MP) libraries for sequencing. A PE library was constructed from one of the seven sub-782 samples (120 individuals, 1µg DNA) sheared by ultrasonication (Covaris) company, 783 average sheared insert size: 350bp). The remaining DNA samples were pooled (6 784 samples, 720 individuals) and used for MP sequencing (3kb, 5kb and 8kb insert sizes), 785 which were prepared with the Nextera mate-pair protocol (Illumina). All libraries were 786 sequenced using an Illumina HiSeq 2000 sequencer (MACROGEN).

787 Long read PacBio (Pacific Biosciences) RS II sequencing was performed from a
788 single DNA extraction of 270 *A. ervi* females, reared on *A. pisum*. Genomic DNA was

789 extracted using the Wizard genomic DNA purification kit (Promega) according to 790 manufacturer instructions and quantified spectrophotometrically using a NanoDrop 791 2000 (Thermo Scientific). Input DNA was mechanically sheared to an average size 792 distribution of 10Kb (Covaris gTube, Kbiosciences) and the resulting library was size 793 selected on a Blue Pippin Size Selection System (Cat #BLU0001, Sage Science) to enrich 794 fragments > 8Kb. Quality and quantity were checked on Bioanalyzer (Agilent 795 Technologies) and Qubit, respectively. Four SMRT RSII cells with P6 chemistry were 796 sequenced at GenoScreen, France.

797

798 Lysiphlebus fabarum

799 DNA was extracted from adult female *L. fabarum* in 10 sub-samples (50-100 wasps 800 each) using the QIAmp DNA mini Kit (Qiagen) according to the manufacturer's 801 instructions, with the inclusion of an overnight tissue digestion at 56 °C. Extracted DNA 802 was then pooled and used to produce Illumina PE and MP, and PacBio libraries. The PE 803 library was prepared using the Illumina Paired-End DNA protocol; the average fragment 804 size was 180 base pair (bp). The MP library (5kb insert) was generated with the Nextera 805 mate-pair protocol (Illumina). Both libraries were sequenced on the Illumina MiSeq in 806 Paired-End mode at the University of Zürich.

Long-read libraries for PacBio RS II sequencing were produced using the DNA Template Prep Kit 2.0 (Pacific Biosciences). Input DNA was mechanically sheared to an average size distribution of 10Kb (Covaris gTube, Kbiosciences) and the resulting library was size selected on a Blue Pippin Size Selection System (Sage Science) machine to enrich fragments > 8Kb; quality and quantity were checked on the Bioanalyzer and Qubit, respectively. Ten SMRT Cells were sequenced at the University of Zürich.

813

### 814 Genome assembly

815 Aphidius ervi

Library quality was checked with FastQC ver. 0.11.3 (Andrews et al. 2010). Paired-end 816 817 libraries were processed with Trimmomatic ver. 0.35 (Bolger et al., 2014) for trimming 818 Illumina adapters/primers, low quality bases (Q <25, 4bp window) and discarding 819 sequences shorter than 50bp or without its mate-pair. In the case of Mate-Pair libraries, 820 removal of improperly oriented read-pairs and removal of Nextera adapters was 821 performed using NextClip (Leggett *et al.* 2014). Filtered PE and MP libraries were used 822 for genome assembly with Platanus ver. 1.2.1 with default parameters (Kajitani et al. 823 2014), gap closing was performed with GapCloser (Luo et al. 2012). Scaffolding with 824 PacBio reads was performed using a modified version of SSPACE-LR v1.1 (Boetzer & 825 Pirovano 2014), with the maximum link option set by –a 250. Finally, the gaps of this 826 last version were filled with the Illumina reads using GapCloser.

827

## 828 Lysiphlebus fabarum

Library quality was also checked with FastQC (Andrews *et al.* 2010). Illumina reads were filtered using Trimmomatic to remove low quality sequences (Q<25, 4bp window), to trim all Illumina primers, and to discard any sequence shorter than 50bp or without its mate-pair. NextClip was used to remove all improperly oriented read pairs.

Raw PacBio reads were error-corrected using the quality filtered Illumina data with the program Proovread (Hackl *et al.* 2014). These error-corrected reads were then used for *de novo* assembly in the program *canu* v1.0 (Koren et al. 2017). Since our PacBio reads were expected to have approximately 30X coverage (based on the presumed size of 128MB), *Canu* was run with the recommended settings for low coverage data (corMhapSensitivity=high corMinCoverage=2 errorRate=0.035), and with the specification that the genome is approximately 128Mbp. The resulting assembly was polished using Pilon (Walker *et al.* 2014) to correct for both single nucleotide and small indel errors, using mapping of both the MP and PE data, generated with bwa-mem (Li & Durbin 2009).

843

#### 844 Linkage map construction: L. fabarum

For linkage map construction, we followed the methodology described in Wang *et al.* (2013) and Purcell *et al.* (2014). In brief, we genotyped 124 haploid male offspring from one sexual female using ddRADseq. Whole-body DNA was high-salt extracted (Aljanabi & Martinez 1997), digested with the *EcoRI* and *Msel* restriction enzymes, and ligated with individual barcodes (Parchman *et al.* 2012; Peterson *et al.* 2012). Barcoded samples were purified and amplified with Illumina indexed primers by PCR (Peterson *et al.* 2012) and quality-checked on an agarose gel.

852 Pooled samples were sequenced on the Illumina HiSeq2500. Raw single-end libraries were quality filtered and de-multiplexed using the process radtags routine 853 854 within Stacks v1.28 with default parameters (Catchen et al. 2011), and further filtered 855 for possible adapter contamination using custom scripts. Genotyping was performed by 856 mapping all samples against the *L. fabarum* draft genome assembly using bowtie2 857 (Langmead & Salzberg 2012) with rg-id, sensitive and end-to-end options. Genotypes 858 were extracted using samtools mpileup (Li et al. 2009) and bcftools (haploid option, Li 859 2011). We filtered the resulting genotypes for a quality score >20 and removed loci 860 with >20% missing data and/or a minor allele frequency <15% using VCFtools v0.1.12b 861 (Danecek *et al.* 2011). After filtering, 1,319 biallelic SNPs in 90 offspring remained.

For constructing linkage groups, we followed Gadau (2009) to account for the 862 unknown phase of the maternal genotype. In short, we duplicated the haploid male 863 864 genotypes and reversed the phase for one duplicated set and removed one of the 865 mirror linkage group sets after mapping. We generated the map using MSTmap (Wu et al. 2008) on the data with following parameters: population type DH, 866 distance function kosambi, no map dist 15.0, no map size 2, missing threshold 1.00, 867 868 and the cut off p value 1e-6. The cut-off p-value was adjusted to create a linkage map 869 of five linkage groups, however the biggest group had a gap of >70 cM, indicating a false 870 fusion of two groups, which we split in two groups. This result corresponded to the six 871 chromosomes previously described for L. fabarum (Belshaw & Quicke 2003), these 872 were visualized with AllMaps (Tang *et al.* 2015). Initial mapping showed that 14 SNPs at 873 one end of tig0000000 mapped to Chromosome1, while the majority of the contig 874 (>150,000 bp) mapped to Chromosome 2. Thus, these SNPs were removed from the 875 linkage maps, and it is advised that subsequent drafts of the *L. fabarum* genome should 876 split this contig around position 153,900.

877

### 878 Genome completeness and synteny

879 Completeness of the two assemblies was assessed by identifying Benchmarking
880 Universal Single-Copy Orthologs (BUSCOs) using the BUSCO v3.0.2 pipeline in genome
881 mode (Simão *et al.* 2015). We identified single copy orthologs based on the
882 Arthropoda\_db9 (1,066 genes, training species: *Nasonia vitripennis*).

883 Synteny between the two genomes was assessed using the NUCmer aligner, 884 which is part of the MUMmer v3.23 package (Kurtz *et al.* 2004). For this, we used the

*L. fabarum* chromosomes as the reference, and included the scaffolds not incorporated
into chromosomes (total 1,407 pieces). The *A. ervi* assembly was mapped to this using
the default settings of NUCmer.

888

# 889 Predictive gene annotation

890 For both assembled genomes, gene predictions were generated using MAKER2 (Holt & 891 Yandell 2011). Within MAKER2, predictive training was performed in a three step 892 process. A first set of genes was predicted by similarity to known proteins or contigs 893 from RNAseq in the same species (described below). This gene set was used thereafter 894 for training both Augustus (Keller et al. 2011) and SNAP (Korf 2004), in two steps, with 895 the results of the first training re-used to train the software in the second round. 896 Transcriptomic evidence was provided separately for each species. For A. ervi, six 897 separate de novo transcriptome assemblies from Trinity (Grabherr et al. 2011) were constructed, one each for the adults reared on different hosts (NCBI PRJNA377544, 898 899 Ballesteros et al. 2017). For each transcript, we only included variants based on filtering 900 with RSEM v 1.2.21 using the option -fpkm cutoff 1.0, --isopct cutoff=15.00. This 901 resulted in 452,783 transcripts. For L. fabarum, we utilized a joint transcriptome, built 902 using RNAseq data (NCBI PRJNA290156) collected from adults (Dennis et al. 2017) and 903 4-5 day old larvae (Dennis et al. in review). Peptide evidence came from the 904 Hymenoptera genomes database (http://hymenopteragenome.org, Acromyrmex 905 echiniator v3.8, Apis mellifera v3.2, Nasonia vitripennis v1.2), from the BioInformatics Platform of Agroecosystems Arthropod database (https://bipaa.genouest.org, 906 *Hyposoter didymator* v1.0), and *Drosphila melanogaster* (http://flybase.org, v6.13), and 907 908 SwissProt (October 2016) databases. Summary statistics were generated with GAG

909 (Hall et al. 2014). Transcriptomic support for the predicted genes was estimated by

- 910 mapping available transcriptomic data (same as above) to the respective genomes
- 911 using STAR (Dobin *et al.* 2013) in the "quantMode".
- 912

# 913 Functional annotation

The putative functions of the proteins predicted by the above pipeline were identified based on blastp (v2.5.0) matches against Genbank *nr* (non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF) release 12/2016 and interproscan v5 against Interpro (1.21.2017). GO terms associations were collected from blast *nr* and interproscan results with blast2GO (v2.2). Finally, transmembrane domains were identified with Hidden Markov Models (HMM) in tmhmm v2.0c, and peptide signals with signalP (euk v4.1, Emanuelsson *et al.* 2007; Nielsen 2017).

921

#### 922 Transposable elements

923 Transposable elements (TE) were predicted using the REPET pipeline (Flutre et al. 924 2011), combining *de novo* and homology-based annotations. *De novo* prediction of TEs was restricted to scaffolds larger than the scaffold N50 for each species. Within these, 925 926 repetitive elements were identified using a blast-based alignment of each genome to 927 itself followed by clustering with Recon (Bao & Eddy 2002), Grouper (Quesneville et al. 928 2005) and Piler (Edgar & Myers 2005). For each cluster, a consensus sequence was 929 generated by multiple alignment of all clustered elements with MAP (Huang 1994). The 930 resulting consensus was then scanned for conserved structural features or homology 931 to nucleotide and amino acid sequences from known TEs (RepBase 20.05, Bao et al. 932 2015; Jurka 1998) using BLASTER (tblastx, blastx, Flutre et al. 2011) or HMM profiles of

933 repetitive elements (Pfam database 27.0) using hmmer3 (Mistry et al. 2013). Based on 934 identified features, repeats were classified using Wicker's TE classification as 935 implemented in the PASTEclassifier (Hoede et al. 2014). The resulting de novo TE library 936 for the genome was then filtered to retain only the elements with at least one perfect 937 match in the genome. Subsequently, all TEs in the genomes were annotated with 938 REPET's TE annotation pipeline. Reference TE sequences were aligned to the genome 939 using BLASTER, Repeat Masker (Smit et al. 2013-2015) and CENSOR (Kohany et al. 940 2006). The resulting HSPs were filtered using an empirical statistical filter implemented 941 in REPET (Flutre et al. 2011) and combined using MATCHER (Quesneville et al. 2005). 942 Short repeats were identified using TRF (Benson 1999) and Mreps (Kolpakov et al. 943 2003). Elements in genomic sequences with homology with known repbase elements 944 (RepBase 20.05) were identified with BLASTER (blastx, tblastx) and curated by 945 MATCHER. Finally, redundant TEs and spurious SSR annotations were filtered and 946 separate annotations for the same TE locus were combined using REPET's "long join 947 procedure".

948

# 949 GC content and codon usage

We examined several measures of nucleotide composition, at both the nucleotide and protein level. Whole genome GC content was calculated by totaling the numbers of A, C, T, and G in the entire assembly. In the predicted coding sequences, this was also calculated separately for each predicted gene and third position GC composition was calculated separately in the predicted coding sequences. In all cases, this was done with the sscu package in R (Sun 2016). Relative Synonymous Codon Usage (RSCU) was extracted from the entire CDS using the seqinR package in R (Charif & Lobry 2007), and 957 visualized with a PCA (R packages factoextra, reshape, and ggplot2, Kassambara & 958 Mundt 2016; Wickham 2007, 2009). To examine GC content in coding genes of other 959 insects, we downloaded the 118 available CDS in the RefSeg database of NCBI (date: 960 October 2018) and again calculated per-gene GC content. 961 To examine the GC content of life-stage biased transcripts, we compared GC 962 content in the genes that are significantly (FDR < 0.05) differentially expressed between previously generated transcriptomes from adult (Dennis et al. 2017) and larval (Dennis 963 964 et al. in revision) L. fabarum, as well in the 10% most highly expressed genes in adults

965 and larvae.

966

### 967 Orphan genes

968 We identified orphan genes as those for which we could not find orthologs in any other 969 sequenced genomes. To do this, we first used OrthoFinder (Emms & Kelly 2015) to 970 generate clusters of orthologous and paralogous genes among the predicted genes 971 (CDS) from the genomes of A. ervi and L. fabarum, as well as five other sequenced 972 parasitoids (Diachasma alloeum, Fopius arisanus, Macrocentrus cingulum, Microplitis demolitor and Nasonia vitripennis). OrthoFinder produces a set of genes that were not 973 974 assigned to any orthogroup. We identified species specific genes, which we are calling 975 orphan genes, by removing all genes that had hits to any other genes in the *nt*, *nr*, and 976 swissprot NCBI database (June 2019). Within these putative orphans, we only retained 977 those with transcriptomic support.

978

979

980

## 981 Gene family expansions

982 We examined gene families that have expanded and contracted in A. ervi and L. 983 fabarum relative to one another using the OMA standalone package (v2.2.0, default 984 values, Altenhoff et al. 2018). OMA was used to compute orthologs (OMA groups) and 985 Hierarchical Orthologous Groups (HOGs) for the predicted proteins of *L. fabarum* and 986 A. ervi: 15,203 and 20,344, respectively. While OMA groups consist of strict 1:1 987 orthologs between OGS1 and OGS3, HOGs contain all orthologs and paralogs of a given 988 predicted gene family. HOGs were parsed with a custom Perl script to identify all gene 989 families in which one of the wasp species contained more members than the other. We 990 focused on only the groups that contained more than 20 genes (ten groups, 991 Supplementary Figure 12). These were identified by blastx against the *nr* database in 992 NCBI.

993

#### 994 Venom proteins

995 The *L. fabarum* venom proteomic analysis was performed from 10 extracted venom 996 glands (Supplementary Figure 14). The 16 most visible bands in 1D gel electrophoresis were cut, digested with trypsin and analyzed by mass spectrometry. All raw data files 997 998 generated by mass spectrometry were processed to generate mgf files and searched 999 against: (i) the L. fabarum proteome predicted from the genome (L. fabarum 1000 annotation v1.0 proteins) and (ii) the L. fabarum de novo transcriptome (Dennis et al. 1001 2017) using the MASCOT software v2.3 (Perkins et al. 1999). The mass spectrometry 1002 proteomics data have been deposited to the ProteomeXchange Consortium 1003 (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository 1004 (Hanrahan & Johnston 2011), with the ID PXD015758.

1005 Sequence annotation was performed based on blast similarity searches. Signal 1006 peptide prediction was performed with SignalP (Emanuelsson et al. 2007; Nielsen 1007 2017). Searches for protein domains was performed with PfamScan (Finn *et al.* 2013) 1008 and venom protein genes were identified using the blast tools in Apollo (Dunn et al. 1009 2019; Lee et al. 2013). Multiple amino acid sequence alignments were made with 1010 MUSCLE (Edgar 2004a, b). Phylogenetic analysis was performed using maximum 1011 likelihood (ML) with PhyML 3.0 (Guindon et al. 2010). SMS was used to select the best-1012 fit model of amino acid substitution for ML phylogeny (Lefort et al. 2017).

1013

# 1014 Manual gene curation

1015 The two genome assemblies were manually curated for a number of gene families of 1016 interest. This improved their structural and functional annotation for more in-depth 1017 analysis. Manual curation, performed in Apollo included the inspection of stop/start 1018 codons, duplications (both true and erroneous), transcriptomic support, and 1019 concordance with the predicted gene models.

1020

# 1021 Desaturases

1022 Desaturase genes in both genomes were automatically identified and annotated with 1023 GeMoMa (Keilwagen *et al.* 2016) using desaturase gene annotations from *Diachasma* 1024 *alloeum, Fopius arisanus,* and *Microplitis demolitor,* retrieved from NCBI's protein 1025 database as queries (retrieved May 2017). Additionally, all desaturase genes were 1026 manually inspected.

1027To measure the production of desaturases in *A. ervi*, wasps were freeze-killed1028and stored separately by sex at - 20 °C. For CHC extraction, single individuals were

1029 covered with 50 µl of MS pure hexane (UniSolv) in 2 ml GC vials (Agilent Technologies,) 1030 and swirled for 10 minutes on a Thermo-shaker (IKA KS 130 Basic, Staufen). The hexane 1031 extracts where then transferred to a fresh conical 250 µl GC insert (Agilent 1032 Technologies), where the hexane was completely evaporated under a constant flow of 1033  $CO_2$ . The dried extract was then resuspended in 5  $\mu$ l of a hexane solution containing 1034 7.5 ng/µl of n-dodecane (EMD Millipore Corp.) as an internal standard. 3 µl of the 1035 extract were then injected into a GC-QQQ Triple Quad (GC: 7890B, Triple Quad: 7010B, 1036 Agilent) with a PAL Autosampler system operating in electron impact ionization mode. 1037 The split/splitless injector was operated at 300 °C in Pulsed splitless mode at 20 psi until 1038 0.75 min with the Purge Flow to Split Vent set at 50 mL/min at 0.9 min. Separation of 1039 compounds was performed on a 30 m x 0.25 mm ID x 0.25  $\mu$ m HP-1 1040 Dimethylpolysiloxane column (Agilent) with a temperature program starting from 60 1041 °C, held for 2 min, and increasing by 50 °C per min to 200 °C, held for 1 min, followed 1042 by an increase of 8 °C per min to 250 °C, held again for 1 min, and finally 4 °C per min 1043 to 320 °C, held for 10 min. Post Run was set to 325 °C for 5 min. Helium served as carrier 1044 gas with a constant flow of 1.2 ml per min and a pressure of 10.42 psi. Initially CHC 1045 peaks were identified and the chromatogram was generated using the Qualitative 1046 Analysis Navigator of the MassHunter Workstation Software (vB.08.00 / Build 1047 8.0.8208.0, Agilent). CHC quantification was performed using the Quantitative Analysis 1048 MassHunter Workstation Software (vB.09.00 / Build 9.0.647.0, Agilent). Peaks were 1049 quantified using their diagnostic (or the neighboring most abundant) ion as quantifier 1050 and several characteristic ions in their mass spectra as qualifiers to allow for 1051 unambiguous detection by the quantification software. The pre-defined integrator 1052 Agile 2 was used for the peak integration algorithm to allow for maximum flexibility. All

1053 peaks were then additionally checked for correct integration and quantification, and,

1054 where necessary, re-integrated manually. Percentages were based on the respective

- 1055 averages of four individual female CHC extracts.
- 1056

#### 1057 Immune genes

1058 The list of immune genes to be searched against the A. ervi and L. fabarum genomes 1059 was established based on *Drosophila melanogaster* lists from the Lemaitre laboratory (lemaitrelab.epfl.ch/fr/ressources, adapted from De Gregorio et al. 2001; De Gregorio 1060 1061 al. 2002) from the interactive et and flv web site 1062 (www.sdbonline.org/sites/fly/aignfam/immune.htm and Buchon et al. 2014). Each D. 1063 melanogaster protein sequence was used in blast similarity searches against the two 1064 predicted wasp proteomes. The best match was retained, and its protein sequence was 1065 used to perform a new blast search using the NCBI non-redundant protein sequence database to confirm the similarity with the *D. melanogaster* sequence. When both 1066 1067 results were concordant, the retained sequence was then searched for in Nasonia 1068 vitripennis and Apis mellifera proteomes to identify homologous genes in these species.

1069

# 1070 Osiris genes

1071 Osiris gene orthologs were determined with a two-part approach: candidate gene 1072 categorization followed by phylogenetic clustering. Candidate Osiris genes were 1073 generated using HMM (with hmmer v3.1b2, Wheeler & Eddy 2013) and local alignment 1074 searching (blast, Altschul *et al.* 1990). A custom HMM was derived using all 24 well 1075 annotated and curated Osiris genes of *Drosophila melanogaster*. Next, an HMM search 1076 was performed on the *A. ervi* and *L. fabarum* proteomes, extracting all protein models

1077 with P < 0.05. Similarly, all D. melanogaster Osiris orthologs were searched in the 1078 annotated proteomes of *A. ervi* and *L. fabarum* using protein BLAST (e < 0.05). The top 1079 BLAST hit for each ortholog was then searched within each parasitoid genome for 1080 additional paralogs (e < 0.001). All unique candidates from the above approaches were 1081 then aligned using MAFFT (Katoh & Standley 2013), and an approximate maximum-1082 likelihood phylogeny was constructed using FastTree (Price et al. 2009) via the CIPRES 1083 science gateway of Xsede (Miller et al. 2015). The species used were: the fruit fly (D. 1084 *melanogaster*), the tobacco hornworm moth (*Manduca sexta*), the silkworm moth 1085 (Bombyx mori), the flour beetle (Tribolium castaneum), the jewel wasp (Nasonia 1086 vitripennis), the honeybee (Apis mellifera), the buff tail bumble bee (Bombus terrestris), 1087 the red harvester ant (Pogonomyrmex barbatus), the Florida carpenter ant 1088 (*Camponotus floridanus*), and Jerdon's jumping ant (*Harpeqnathos saltator*).

1089

1090 **OXPHOS** 

1091 Genes involved in the oxidative phosphorylation pathway (OXPHOS) were identified in 1092 several steps. Initial matches were obtained using the nuclear-encoded OXPHOS 1093 proteins from Nasonia vitripennis (Gibson et al. 2010; J. D. Gibson unpublished) and 1094 Drosophila melanogaster (downloaded from www.mitocomp.uniba.it: Porcelli et al. 1095 2007). These two protein sets were used as queries to search the protein models 1096 predicted for A. ervi and L. fabarum (blastp, Altschul et al. 1997). Here, preference was 1097 given to matches to N. vitripennis. Next, genes from the N. vitripennis and D. 1098 *melanogaster* reference set that did not have a match in the predicted proteins were 1099 used as queries to search the genome-assembly (blastn), in case they were not in the 1100 predicted gene models. Gene models for all matches were then built up manually,

1101 based on concurrent evidence from the matches in both *A. ervi* and *L. fabarum* and 1102 their available expression evidence. The resulting protein models were aligned to one 1103 another and to *N. vitripennis* using MAFFT (Katoh & Standley 2013) to identify missing 1104 or extraneous sections. These results were used as queries to search the *N. vitripennis* 1105 proteins to ensure that all matches are reciprocal-best-blast-hits. Gene naming was 1106 assigned based on the existing *N. vitripennis* nomenclature. Potential duplicates were 1107 flagged based on blast-matches back to *N. vitripennis* (Additional Data 10).

1108

# 1109 Olfactory genes

# 1110 Odorant-binding proteins (OBPs) and chemosensory Proteins (CSPs)

1111 To identify OBPs based on homology to known sequences, we retrieved 60 OBP amino 1112 acid sequences from other Braconidae (namely Fopius arisanus and Microplitis 1113 demolitor) from GenBank. To this, we added seven OBPs found in a previous 1114 transcriptome of A. ervi (Patrizia Falabella, unpublished, EBI SRI Accessions: 1115 ERS3933807- ERS3933809). To identify CSPs, we used CSP amino acid sequences from 1116 more Hymenoptera species (Apis mellifera, Nasonia vitripennis, Fopius arisanus and 1117 Microplitis demolitor). These sets were used as query against A. ervi and L. fabarum 1118 genomes using tblastn (e-value cutoff 10e-3 for OBPs and 10e-2 for CSPs). Genomic 1119 scaffolds that presented a hit with at least one of the query sequences were selected. 1120 To identify precise intron/exon boundaries, the Braconidae OBP and CSP amino acid 1121 sequences were then aligned on these scaffolds with Scipio (Keller et al. 2008) and 1122 Exonerate (Slater & Birney 2005). These alignments were used to generate gene 1123 models in Apollo. Gene models were manually curated based on homology with other 1124 Hymenoptera OBP and CSP genes and on RNAseq data, when available. Lastly, the deduced amino acid sequences of *A. ervi* and *L. fabarum* OBP and CSP candidates were then used as query for another tblastn search against the genomes in an iterative process to identify any additional OBPs. Since both OBPs and CSPs are secreted proteins, the occurrence of a signal peptide was verified using SignalP (Emanuelsson *et al.* 2007; Nielsen 2017).

1130

## 1131 Odorant receptors (ORs)

ORs were annotated using available OR gene models from *Diachasma alloeum*, *Fopius arisanus*, and *Microplitis demolitor* retrieved from NCBIs protein database (retrieved May 2017). Preliminary OR genes models for *A.ervi* and *L. fabarum* were predicted with exonerate (v2.4.0), GeMoMa (v1.4, Keilwagen 2016), and combined with EVidence Modeler (v.1.1.1, Haas *et al.* 2008). These preliminary models were subsequently screened for the 7tm\_6 protein domain (with PfamScan v1.5) and manually curated in WebApollo2.

1139 In an iterative approach, we annotated the IRs using known IR sequences from 1140 Apis melifera, Drosophila melanogaster, Microplitis demolitor and Nasonia vitripennis 1141 as queries to identify IRs in the genomes of *A. ervi* and *L. fabarum*. The hymenopteran 1142 IR sequences served as input for the prediction of initial gene model with Exonerate 1143 (Slater & Birney 2005) and GeMoMa (Keilwagen et al. 2016). Then, we inspected and 1144 edited homologous gene models from each tool in the Apollo genome browser to 1145 adjust for proper splice sites, start and stop codons in agreement with spliced RNA-Seq 1146 reads. After a first round of prediction, we repeated the whole process and provided 1147 the amino acid sequences of curated IR genes as queries for another round of 1148 predictions to identify any remaining paralogous IRs.

Multiple sequence alignments of the IRs were computed with hmmalign (Eddy 1998) using a custom IR HMM to guide the alignments (Harrison *et al.* 2018). Gene trees were generated with FastTree v2 (Price *et al.* 2010) using the pseudocount option and further parameters for the reconstruction of an exhaustive, accurate tree (options: -pseudo -spr 4 -mlacc 2 -slownni). Resulting trees were visualized with iTOL v4 (Letunic & Bork 2019), well supported IR clusters and expansions were highlighted by color (branch support > 0.9).

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# 1157 Sex Determination

1158 Ortholog searches were performed with tblastn (Altschul et al. 1997) against the 1159 genomic scaffolds. Hits with an e-value smaller than 1e-20 were assessed, apart from 1160 transformer and doublesex where any hit was surveyed. Doublesex, Transformer-2 and 1161 Transformer peptide sequences of Asobara tabida (NCBI accessions MF074326-1162 MF074334) were used as queries for the core sex determination genes. This braconid 1163 species is the closest relative whose sex determination mechanism has been examined 1164 (Geuverink et al., 2018). The putative transformerB sequence of A. ervi was blasted for 1165 verification against the transcriptome of Aphidius colemani (Peters et al. 2017) and a 1166 highly conserved fragment was detected (GBVE01021531). Peptide sequences of sex 1167 determination related genes to use as queries were taken from *Nasonia vitripennis*: 1168 Fruitless (NP 001157594), Sex-Lethal homolog (XP 016836645), pre-mRNA-splicing 1169 factor CWC22 homolog (XP 001601117) and RNA-binding protein 1-like 1170 (XP 008202465). Hidden Markov models were not used as gene models because the 1171 ensuing peptide predictions did not contain all putative homologs (e.g. transformerB in 1172 A. ervi) due to fragmentation of the scaffolds containing the candidate genes.

# 1174 DNA methylation genes

1175	The genomes were searched with tblastn (Altschul et al. 1997) for the presence of
1176	potential DNA methyltransferase genes using peptide sequences from Apis mellifera
1177	and N. vitripennis as queries. These species differ in their copy number of DNMT1, with
1178	two copies (NP_001164522, XP_006562865) in the honeybee A. mellifera (Wang et al.
1179	2006) and three copies (NP_001164521 ,XP_008217946, XP_001607336) in the wasp
1180	N. vitripennis (Werren et al. 2010). DNMT2, currently characterized as EEF1AKMT1
1181	(EEF1A Lysine Methyltransferase 1), has become redundant in the list of DNA
1182	methyltransferase genes as it methylates tRNA instead, but was surveyed here as a
1183	positive control ( <i>N. vitripennis</i> NP_001123319, <i>A. mellifera</i> XP_003251471). DNMT3
1184	peptide sequences from N. vitripennis (XP_001599223) and from A. mellifera
1185	(NP_001177350) were used as queries for this gene. Low levels of methylation were
1186	confirmed by mapping the whole genome bisulfite sequencing data generated by
1187	Bewick <i>et al.</i> (2017) back to the <i>A. ervi</i> genome assembly.

# 1197 List of abbreviations

- 1198 A, T, C, G, and U: Adenine, Thymine , Cytosine, Guanine, and Uracile, nucleotides
- **bp**: Base Pair
- 1200 BIPAA: BioInformatics Platform for Agroecosystem Arthropods (bipaa.genouest.org)
- **BUSCO**: Benchmarking Universal Single-Copy Orthologs
- 1202 CDS: Predicted Coding Sequence
- **CSD**: Complementary Sex Determination
- 1204 CHC: Cuticular Hydrocarbons
- **DNMT**: DNA Methyltransferase genes
- **CSP**: Chemosensory Protein
- 1207 GO: Gene Ontology
- 1208 HMM: Hidden Markov Model
- 1209 HOG: Hierarchical Ortholog Group
- 1210 IR: Ionotropic Receptor
- *LRR*: Leucine Rich Repeat Proteins
- 1212 Mbp: Mega Base Pairs, or 1,000,000bp
- 1213 MP: Mate-pair sequence data
- 1214 NCBI: National Center for Biotechnology Information
- **N50**: A measure of genome completeness. The length of the scaffold containing the
- 1216 middle nucleotide
- **OXPHOS**: Oxidative Phosphorylation
- **OBP**: Odorant-binding Protein
- **OR**: Odorant Receptor
- **PE**: Paired-end sequence data
- 1221 RSCU: Relative Synonymous Codon Usage
- **TE**: Transposable Element

# 1243 Availability of data and materials

1244	Both genomes are available from the NCBI Genome database (PRJNA587428, A. ervi:
1245	SAMN13190903, L. fabarum: SAMN13190904). The assemblies, predicted genes, and
1246	annotations are also available at https://bipaa.genouest.org. Raw Illumina and PacBio
1247	sequence data used to construct genomes is available in NCBI SRA for both A. ervi
1248	(SAMN12878248) and L. fabarum (accessions SAMN10617865, SAMN10617866,
1249	SAMN10617867), and is further detailed in Supplementary Tables 1 and 2. Venom
1250	protein data are available via ProteomeXchange with identifier PXD015758.
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# 1918 FIGURES IN MAIN TEXT (title)

1919

1920 Figure 1. Aphid parasitoid life cycle: Generalized life cycle of *Aphidius ervi* and 1921 *Lysiphlebus fabarum*, two different parasitoid wasps that target aphid hosts.

1922

Figure 2. Codon usage in predicted genes: Proportions of all possible codons, as used in the predicted genes in *A. ervi* (top) and *L. fabarum* (bottom). Codon usage was measured as relative synonymous codon usage (RSCU), which scales usage to the number of possible codons for each amino acid (RSCU). Codons are listed at the bottom and are grouped by the amino acid that they encode. The green line depicts GC contend (%) of the codon.

1929

1930Figure 3. GC and nitrogen content of expressed genes: We observe significant1931differences (p-values from two-sided t-test) in the GC content between adult and larval1932L. fabarum in: (A) the most highly expressed 10% of the genes and (B) genes that are1933differentially expressed between adults and larvae. In contrast, there is no difference1934in the nitrogen content of the same set of genes (C, D).

1935

Figure 4. Overlap in Venom proteins between *A. ervi* and *L. fabarum*: Overlap in venom
proteins (A) and venom protein putative function (B) between *A. ervi* and *L. fabarum*

1938

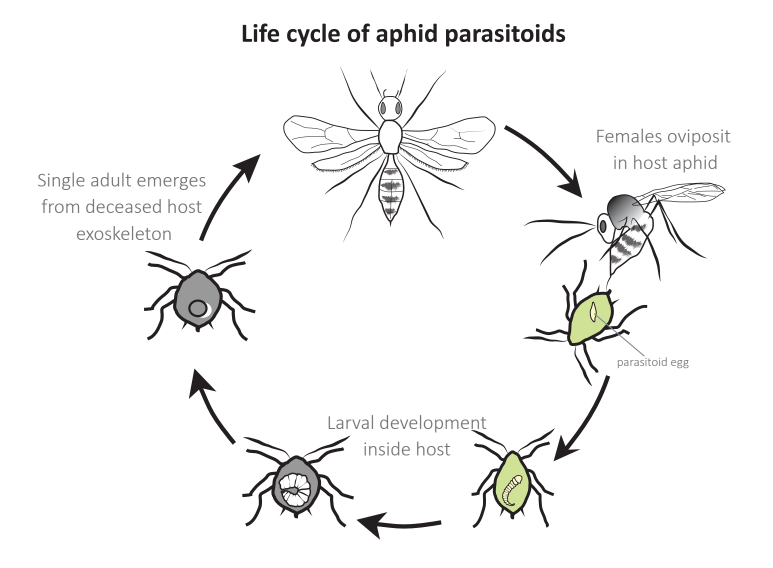
1939 Figure 5: Phylogeny of hymenopteran GGT sequences. A. ervi/L. fabarum and N. 1940 vitripennis/P. puparum venom GGT sequences are marked with blue and orange rectangles respectively. Letters A, B and C indicate the major clades observed for 1941 1942 hymenopteran GGT sequences. Numbers at corresponding nodes are aLRT values. 1943 Only aLRT support values greater than 0.8 are shown. The outgroup is human GGT6 sequence. 1944 1945 1946 1947 1948 1949 1950 1951 1952 1953 1954 1955

# 1956 List of additional data files

# 1957

- 1958 Additional Data 1: details of genetic positions used to construct linkage groups for *L*.
- 1959 fabarum.
- 1960 Additional Data 2: Genbank numbers and taxa information for genome (CDS) graphed
- 1961 in Supplemental Figure 6.
- 1962 Additional Data 3: file detailing (a) the most highly expressed genes in both taxa and
- 1963 (b) differential expression between adult and larval *L. fabarum*.
- 1964 Additional Data 4: fasta file of orphan genes for A. ervi
- 1965 Additional Data 5: fasta file of orphan genes for *L. fabarum*
- 1966 Additional Data 6: Summary of OMA output, including details of *LRR* genes
- 1967 Additional Data 7: Annotation of venom genes in *L. fabarum* and *A. ervi*
- 1968 Additional Data 8: Details of immune gene annotation
- 1969 Additional Data 9: Expression details of Osiris genes in *L. fabarum* and *A. ervi*
- 1970 Additional Data 10: Details of annotated OXPHOS genes, including duplications in the
- 1971 assembly
- 1972 Additional Data 11: Details of sex determination gene annotations
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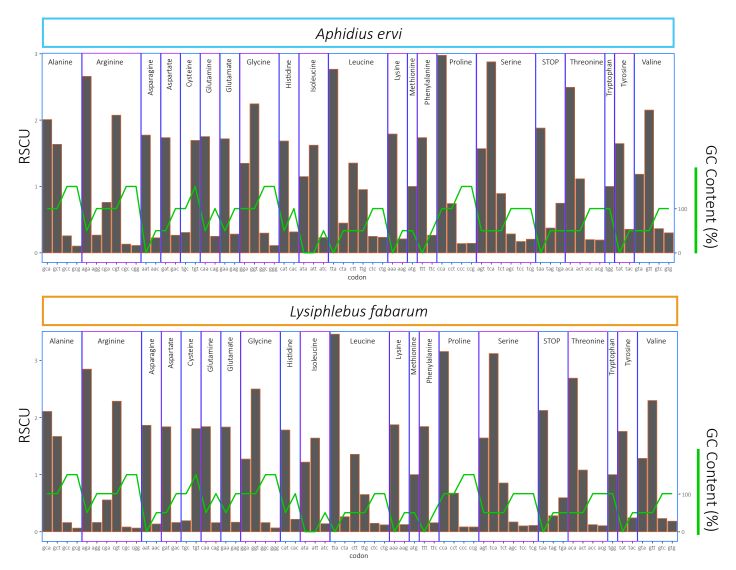
# Figure 1



# Life history characteristics

	Aphidius ervi	Lysiphlebus fabarum
Host insects	Aphididae	Aphididae
Reproductive mode	Sexual	Asexual or sexual
Host is ant tended	No	Yes, usually
Native range	Europe	Europe
Primary host aphid tribe	Macrosiphini	Aphidini

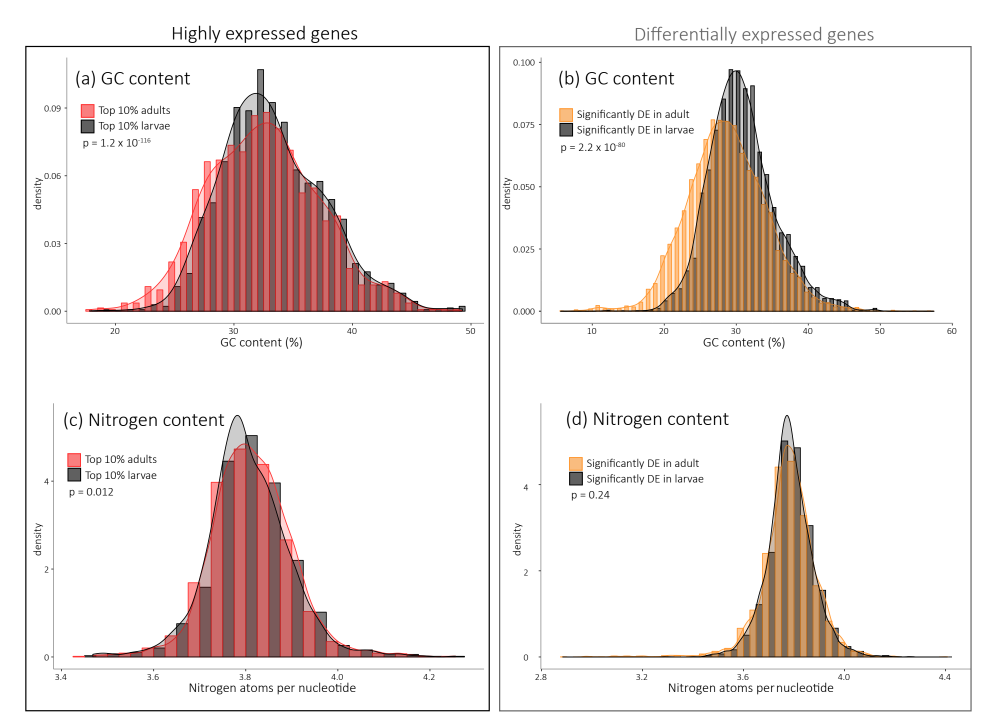
# Figure 2

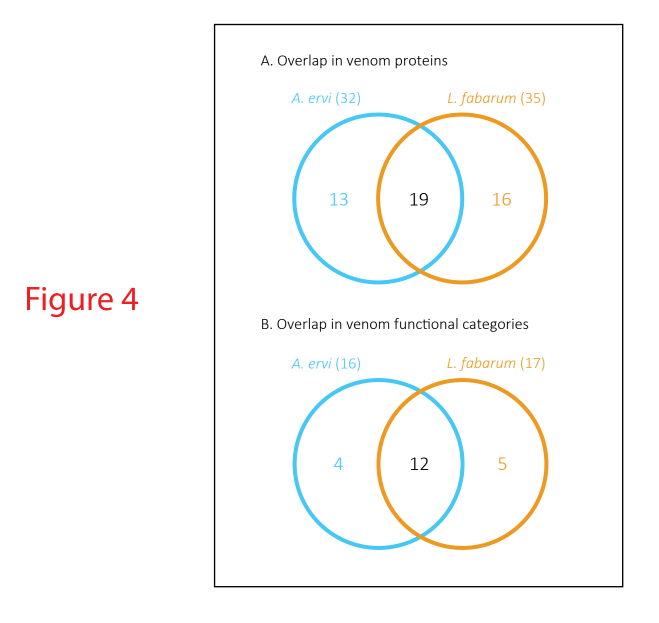


# Codon usage and GC content

# Figure 3

# Genes expressed in larval and adult L. fabarum





# Figure 5

