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Genomes of the Hymenoptera

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Although Hymenoptera is the second-most sequenced arthropod order, with 52 publically archived genomes (71 with ants, reviewed elsewhere), these genomes do not capture the breadth of this very diverse order (Figure 1, Table 1). These sequenced genomes represent only 15 of the 97 extant families. Although at least 55 other genomes are in progress in an additional 11 families (see Table 2), stinging wasps represent 35 (67%) of the available and 42 (76%) of the in progress genomes. A more comprehensive catalog of hymenopteran genomes is needed for research into the evolutionary processes underlying the expansive diversity in terms of ecology, behavior, and physiological traits within this group. Additional sequencing is needed to generate an assembly for even 0.05% of the estimated 1 million Hymenopteran species, with premier level assemblies for at least 0.1% of the >150,000 named species dispersed across the order. Given the haplodiploid sex determination in Hymenoptera, haploid male sequencing will help minimize genome assembly issues to enable higher quality genome assemblies.

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Introduction

The order Hymenoptera with fossils dating back to the Triassic [1], is both ancient and hyper-diverse, with over 150,000 described and one million estimated species [2,3]. One of the ‘big four’ insect orders, Hymenoptera comprise diverse species including sawflies and wood wasps (‘Symphyta’), parasitoid wasps (‘Parasitica’), and stinging wasps (Aculeata), which includes the ubiquitous and ecologically dominant ants, bees and social wasps. With an astonishing diversity of biologically interesting traits Hymenoptera have significant economic impact (e.g. biological control and pollination) and thus was one of the first insect orders to benefit from genome sequencing [4]. Despite additional genomes sequenced (Table 1) or in preparation (Figure 1, Table 2), genomic resources remain lacking for most major lineages, which comprise 28 superfamilies, 97 families, and 8422 genera. Herein we discuss insights gained and opportunities for improvements for Hymenopteran genomics (noting that ants are reviewed separately).

State of hymenopteran genomic resources

Features of hymenopteran genomes

Hymenopteran genomes possess some notable and unique features. Hymenoptera are haplodiploid: unfertilized eggs produce haploid males and fertilized eggs produce diploid females [5]. Haplodiploidy engenders interesting biology related to sex determination [6], control of sex ratios, and relatedness [7], but is also useful for whole genome sequencing. Enough DNA can be extracted from a single large haploid male to provide material for whole genome sequencing without genetic variation (e.g. [8]). Another notable feature of some Hymenoptera is extremely high recombination rates, especially in social species such as the honey bee, where recombination rates are among the highest known for any organism [9].

Among sequenced examples, hymenopteran genomes are moderate in size (80% are between 180 and 340 Mb) with a few exceptions [10,11,12**]. Most possess 12,000–20,000 genes (note counts are highly annotation-pipeline and assembly contiguity dependent [13*]), with a relatively low content of repetitive and transposable elements. One unusual feature is low GC content, which ranges from 30 to 45% depending on the species [8]. Although the reason for low GC content is not

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Table 1

Hymenoptera genomes in INSDC. Family Formicidae 19 genomes are omitted (reviewed elsewhere).

Family	Scientific Name	Representative Assembly	Reference
Agaonidae	<i>Ceratosolen solmsi</i>	GCF_000503995.1	[103]
Apidae	<i>Apis cerana</i>	GCF_001442555.1	[104]
Apidae	<i>Apis dorsata</i>	GCF_000469605.1	[105]
Apidae	<i>Apis florea</i>	GCF_000184785.2	[12**]
Apidae	<i>Apis mellifera</i> [†]	GCF_000002195.4	[4,13*]
Apidae	<i>Bombus impatiens</i> [†]	GCF_000188095.1	[106]
Apidae	<i>Bombus terrestris</i> [†]	GCF_000214255.1	[106]
Apidae	<i>Ceratina calcarata</i>	GCF_001652005.1	[107]
Apidae	<i>Eufriesea mexicana</i>	GCF_001483705.1	[12**]
Apidae	<i>Euglossa dilemma</i>	GCA_002201625.1	[10]
Apidae	<i>Habropoda laboriosa</i> [†]	GCF_001263275.1	[12**]
Apidae	<i>Melipona quadrifasciata</i>	GCA_001276565.1	[12**]
Braconidae	<i>Cotesia vestalis</i>	GCA_000956155.1	[108]
Braconidae	<i>Diachasma alloenum</i>	GCF_001412515.1	[109]
Braconidae	<i>Fopius arisanus</i>	GCF_000806365.1	[17]
Braconidae	<i>Macrocentrus cingulum</i>	GCA_002156465.1	[110]
Braconidae	<i>Microplitis demolitor</i>	GCF_000572035.2	[65]
Cephalidae	<i>Cephus cinctus</i>	GCF_000341935.1	[111]
Diprionidae	<i>Neodiprion lecontei</i>	GCF_001263575.1	[112]
Encyrtidae	<i>Copidosoma floridanum</i>	GCF_000648655.1	[113]
Figitidae	<i>Leptopilina clavipes</i>	GCA_001855655.1	[114]
Halictidae	<i>Dufourea novaeangliae</i>	GCF_001272555.1	[12**]
Halictidae	<i>Lasioglossum albipes</i>	GCA_000346575.1	[115]
Megachilidae	<i>Megachile rotundata</i> [†]	GCF_000220905.1	[12**]
Orussidae	<i>Orussus abietinus</i>	GCF_000612105.1	[116]
Pteromalidae	<i>Nasonia giraulti</i>	GCA_000004775.1	[117*]
Pteromalidae	<i>Nasonia longicornis</i>	GCA_000004795.1	[117*]
Pteromalidae	<i>Nasonia vitripennis</i>	GCF_000002325.3	[117*]
Pteromalidae	<i>Trichomalopsis sarcophagae</i>	GCA_002249905.1	[22*]
Tenthredinidae	<i>Athalia rosae</i>	GCF_000344095.1	[116]
Trichogrammatidae	<i>Trichogramma pretiosum</i>	GCF_000599845.1	[113]
Vespidae	<i>Polistes canadensis</i>	GCF_001313835.1	[18]
Vespidae	<i>Polistes dominula</i>	GCF_001465965.1	[8]

[†] Important pollinators.

* Genome unpublished but included in collection.

88 yet understood, it may be related to GC-biased gene
 89 conversion and high recombination rates [14]. Due to
 90 their relatively small size and simple structure, Hyme-
 91 nopteran genomes are readily assembled and highly trac-
 92 table for genome sequencing [15*].

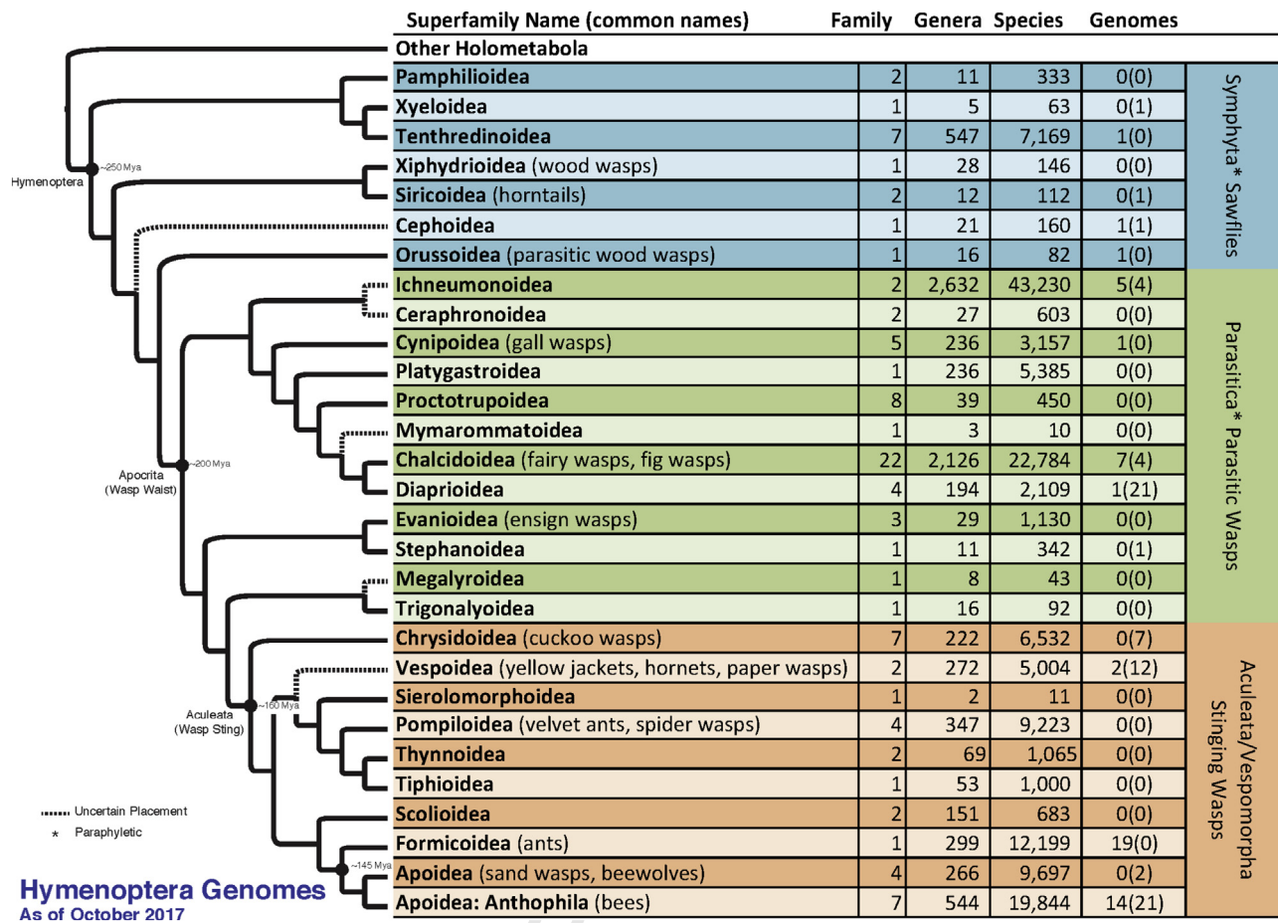
93 Genomes generate biological understanding

94 The Hymenoptera include everything from herbivores to
 95 pollinators to predators to parasites, and the species filling
 96 each of these ecological roles are endowed with a complex
 97 set of physiological and behavioral adaptations. Genomic
 98 studies have informed applied questions related to polli-
 99 nator health [16] and biological control using parasitoid
 100 wasps [17]. Comparative genomics approaches have
 101 unlocked some understanding of the molecular evolu-
 102 tionary basis for these adaptations, including the evolu-
 103 tion of eusociality [8,12**,18], social parasitism [19,20*],
 104 venom function [19,20*,21,22*], and behavioral host spec-
 105 ificity [23,24*].

Potential genomic insights into Hymenoptera biology

106 Pollinator health and management research has greatly
 107 benefited from the availability of reference genomes for
 108 many important pollinators (Table 1). Genomic
 109 approaches to pollinator health allow us to screen for
 110 disease, elucidate the effects of parasites, and investigate
 111 the immune response to environmental stressors and
 112 pathogens. Examples include developing new biomark-
 113 ers of honey bee colony health [25–29], characterizing
 114 environmental stress responses and optimal developmen-
 115 tal temperature regimes for the alfalfa leafcutting bee (an
 116 intensively managed solitary pollinator) [30,31] and
 117 developing other genomic and transcriptomic indicators
 118 of health in wild and managed pollinators [32–39]. Honey
 119 bee genomic nutrition research has been fruitful in gener-
 120 ating insights into mechanisms of bee health [40–43].
 121 Expanding this emerging field to native, unmanaged bees
 122 can provide a comparative perspective on how wild bee
 123 health is influenced by environmental diet restriction due
 124
 125

Figure 1



Hymenoptera phylogeny and genome assemblies. Phylogeny based upon [83**,88**], with dotted lines marking lineages of uncertain placement, branches not to scale (=cladogram). Major groups shown on right, individual superfamilies listed with columns indicating the numbers of Families, Genera, Species, Genomes in NCBI (in Progress).

126 to habitat loss [44]. This approach can also improve our
127 understanding of innate and acquired immunity function
128 among bees, which is critical for management and con-
129 servation efforts [45,46].

130 The vast majority of Hymenoptera species are parasitoids
131 (green in Figure 1 and Table 2). The majority of insect
132 species are attacked by at least one species of parasitic
133 Hymenoptera [47–51], with complex and intimate host-
134 parasitoid interactions [52] and narrow host ranges [53]
135 which may drive ecological speciation [54,55]. Parasitoids
136 are economically important as biological control agents of
137 invasive pests, reducing pest abundance and impact,
138 providing a safe, cost-effective alternative to insecticides
139 [56], so understanding the genetic architecture and evo-
140 lution of traits like climatic adaptation and host specificity
141 is critical. Such research is revealing diverse mechanisms
142 by which parasitoid wasps overcome the host immune
143 system, including venoms, immunosuppressive factors
144 such as polydnviruses and virus-like particles,

145 specialized embryonic cells (teratocytes), and larval secre-
146 tions [57–59]. Parasitoids can also co-opt host endocrine
147 systems, disrupting host development [60] or manipulat-
148 ing host behavior [61,62]. Recent findings have revealed
149 evolutionary changes in some wasp lineages in association
150 with microbes that affect the interactions with their host,
151 suggesting much remains to be discovered. Turnover of
152 integrated viral genomes within Braconidae and Ichnue-
153 monidae wasp genomes are more complex than once
154 anticipated [63,64*,65]. Genomes in progress for three
155 parasitoids with a common host have revealed that each
156 has separate mechanisms to overcome host immunity and
157 other defenses (M. Strand, personal communication)
158 demonstrating that comparative genomics can reveal
159 more information than phylogeny alone as illustrated
160 by the rich collection of research stemming from genomes
161 in the genus *Nasonia* (Pteromalidae; [66] introduces the
162 collection), though annotation of these genomes lever-
163 aged the extensive history of research on *Nasonia* genetics
164 [67]. Comparable research on species from other families

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Table 2

Q8 Hymenoptera Families with Genera and Species counts and Genomes in NCBI, otherwise published (Pub), in progress (Prog), and proposed here (Prop). Classification citations (Class) follow [2,3] with updates from [47,83**,118]. Counts of genera and species are modified from [2] with richness sources cited (Rich). *20 are low coverage genomes.

	Superfamily	Family	Genomes					Species	Rich	Class		
			Total	NCBI	Pub	Prog	Prop				Genera	
Aculeata/Vespomorpha	Thynnoidea	Chyphotidae	–	–	–	–	1	2	65	[119]	[118]	
		Thynnidae	–	–	–	–	2	67	1000	[120]	[118]	
	Tiphioidea	Tiphidae	–	–	–	–	2	53	1000	[1]	[118]	
	Pompiloidea	Pompilidae	–	–	–	–	3	125	4855	[1]	[118]	
		Mutillidae	–	–	–	–	2	200	4252	[1]	[118]	
		Sapygidae	–	–	–	–	1	12	66	[121]	[118]	
	Sierolomorpoidea	Myrmosidae	–	–	–	–	1	10	50	[120]	[118]	
		Sierolomorphidae	–	–	–	–	1	2	11	[122]	[83**]	
	Vespoidea	Rhopalosomatidae	–	–	–	–	1	4	72	[1]	[118]	
		Vespidae	14	2	–	12	–	268	4932	[1]	[118]	
	Formicoidea	Formicidae	20	19	1	n/a	n/a	299	12,199	[123]	[118]	
	Apoidea	Ampulicidae	1	–	–	1	–	6	200	[124]	[118]	
		Heterogynaidae	–	–	–	–	1	1	8	[124]	[118]	
	Apoidea: Anthophila	Sphecidae	–	–	–	–	1	17	716	[124]	[118]	
		Crabronidae*	1	–	–	1	3	242	8773	[124]	[118]	
		Apidae	12	11	–	1	–	209	5749	[125]	[118]	
		Andrenidae	1	–	–	1	1	77	2917	[125]	[118]	
		Megachilidae	3	1	–	2	–	76	4096	[125]	[118]	
		Melittidae	–	–	–	–	1	15	187	[125]	[118]	
		Halictidae	19	2	–	17	–	79	4327	[125]	[118]	
		Stenotritidae	–	–	–	–	1	2	21	[125]	[118]	
		Colletidae	–	–	–	–	2	86	2547	[125]	[118]	
		Scolioidea	Scoliidae	–	–	–	–	1	143	560	[1]	[118]
	Chrysoidea	Bradynobaenidae s.s.	–	–	–	–	1	8	123	[119]	[118]	
		Bethylidae	1	–	–	1	1	84	2340	[1]	[118]	
		Chrysididae	6	–	–	6	–	81	2500	[1]	[118]	
		Dryinidae	–	–	–	–	2	41	1605	[1]	[118]	
		Embolemidae	–	–	–	–	1	2	39	[1]	[118]	
		Plumariidae	–	–	–	–	1	7	22	[1]	[118]	
		Sclerogibbidae	–	–	–	–	1	3	20	[1]	[118]	
		Scolecbythidae	–	–	–	–	1	4	6	[1]	[118]	
		Megalyroidea	Megalyridae	–	–	–	–	1	8	43	[2]	[3]
			Trigonalidae	–	–	–	–	1	16	92	[2]	[3]
	Evanoidea	Aulacidae	–	–	–	–	1	2	185	[2]	[3]	
		Evaniidae	–	–	–	–	1	21	449	[2]	[3]	
	Stephanoidea	Gasteruptiidae	–	–	–	–	1	6	496	[2]	[3]	
		Stephanidae	1	–	–	1	–	11	342	[2]	[3]	
	Chalcidoidea	Eulophidae	1	–	–	1	2	334	4969	[2]	[3]	
		Pteromalidae	5	4	–	1	–	619	3544	[2]	[3]	
		Torymidae	1	–	–	1	–	82	900	[2]	[3]	
Agaonidae		1	1	–	–	–	20	762	[2]	[3]		
Leucospidae		–	–	–	–	1	4	134	[2]	[3]		
Eurytomidae		–	–	–	–	2	97	1453	[2]	[3]		
Azotidae		–	–	–	–	1	1	92	[2]	[2]		
Chalcididae		–	–	–	–	2	90	1469	[2]	[3]		
Cynipencyrtidae		–	–	–	–	1	1	1	[2]	[2]		
Encyrtidae		1	1	–	–	1	493	4058	[2]	[3]		
Eriaporidae		–	–	–	–	1	5	22	[2]	[2]		
Trichogrammatidae		1	1	–	–	–	97	881	[2]	[3]		
Aphelinidae		–	–	–	–	1	29	1078	[2]	[3]		
Eucharitidae		–	–	–	–	1	57	427	[2]	[3]		
Eupelmidae		–	–	–	–	1	51	931	[2]	[3]		
Mymaridae		1	–	–	1	1	96	1437	[2]	[3]		
Ormyridae		–	–	–	–	1	3	125	[2]	[3]		
Perilampidae		–	–	–	–	1	17	284	[2]	[3]		
Rotoitidae		–	–	–	–	1	2	2	[2]	[3]		
Signiphoridae		–	–	–	–	1	4	78	[2]	[3]		
Tanaostigmatidae	–	–	–	–	1	9	93	[2]	[3]			
Tetracampidae	–	–	–	–	1	15	44	[2]	[3]			
Diapriidae	Diapriidae	23	1	–	21*	–	190	2048	[2]	[3]		
	Ismaridae	–	–	–	–	1	1	29	[2]	[2]		

Table 2 (Continued)

	Superfamily	Family	Genomes					Species	Rich	Class		
			Total	NCBI	Pub	Prog	Prop				Genera	
	Monomachidae	–	–	–	–	1	2	30	[2]	[3]		
	Maamingidae	–	–	–	–	1	1	2	[2]	[3]		
Mymarommatoidea	Mymaromatidae	–	–	–	–	1	3	10	[2]	[3]		
Proctotrupoidea	Pelecinidae	–	–	–	–	1	1	3	[2]	[3]		
	Vanhorniidae	–	–	–	–	1	3	5	[2]	[3]		
	Roproniidae	–	–	–	–	1	2	20	[2]	[3]		
	Austroniidae	–	–	–	–	1	1	3	[2]	[2]		
	Heloridae	–	–	–	–	1	1	12	[2]	[3]		
	Peradeniidae	–	–	–	–	1	1	2	[2]	[3]		
	Proctotrupidae	–	–	–	–	1	28	403	[2]	[3]		
	Proctorenyxidae	–	–	–	–	1	2	2	[2]	[3]		
Platygastroidea	Platygastridae	–	–	–	–	3	236	5385	[2]	[3]		
Cynipoidea	Cynipidae	–	–	–	–	2	74	1412	[2]	[3]		
	Figitidae	1	1	–	–	1	148	1571	[2]	[3]		
	Liopteridae	–	–	–	–	1	10	153	[2]	[3]		
	Ibaliidae	–	–	–	–	1	3	20	[2]	[3]		
	Austrocynipidae	–	–	–	–	1	1	1	[2]	[3]		
Ichneumonoidea	Braconidae	5	4	–	–	1	4	1057	19,205	[2]	[3]	
	Ichneumonidae	3	–	–	–	3	9	1575	24,025	[2]	[3]	
Ceraphronoidea	Megaspilidae	–	–	–	–	1	15	304	[2]	[3]		
	Ceraphronidae	–	–	–	–	1	12	299	[2]	[3]		
Symphyta*	Orussoidea	Orussidae	1	1	–	–	–	16	82	[2]	[3]	
	Cephoidea	Cephoidea	2	1	–	–	–	21	160	[2]	[3]	
	Xiphydrioidae	Xiphydriidae	–	–	–	–	–	1	28	146	[2]	[3]
	Siricoidea	Siricidae	–	–	–	–	–	1	11	111	[2]	[3]
		Anaxyelidae	–	–	–	–	–	1	1	1	[2]	[3]
	Xyeloidea	Xyelidae	1	–	–	–	–	5	63	[2]	[3]	
	Tenthredinoidea	Argidae	–	–	–	–	–	1	58	897	[2]	[3]
		Blasticotomidae	–	–	–	–	–	1	2	12	[2]	[3]
		Cimbicidae	–	–	–	–	–	1	16	182	[2]	[3]
		Diprionidae	–	–	–	–	–	1	11	136	[2]	[3]
		Heptamelidae	–	–	–	–	–	1	2	38	[126,127]	[3]
		Pergidae	–	–	–	–	–	1	60	442	[2]	[3]
		Tenthredinidae	1	1	–	–	–	2	398	5462	[2]	[3]
	Pamphilioidea	Pamphiliidae	–	–	–	–	–	1	1	42	[2]	[3]
		Megalodontesidae	–	–	–	–	–	1	10	291	[2]	[3]
		Family counts	26	15	1	20	81					
		Sum totals	128	52	1	75*	108	8422	152,658			

will be needed to fully realize the potential of genomics for parasitic Hymenoptera.

Mechanisms and evolution of social behavior are an emphasis of Hymenoptera genomic research. Aculeata are known for sophisticated social behavior within colonies of some ants, bees, and wasps, with eusociality arising multiple times within this clade [68,69]. Glimpses into the molecular basis for this cooperative lifestyle were among the most anticipated findings from the initial sequencing of the honey bee genome [4] while more recent comparative genomic approaches have placed these initial findings within a broader phylogenetic context [8,12^{**},18]. Comparative methods are particularly useful for understanding the molecular basis for hymenopteran behavior given the impressive diversity of social niches [70^{*},71]. Although social wasps have been a model lineage for understanding the evolution of sociality [72], there are currently only two published genomes for this group, both within the paper wasp genus *Polistes* [8,18].

While most of the focus of hymenopteran genomics has been on social evolution, the Hymenoptera offer many opportunities to further investigate the molecular basis for other aspects of development and animal behavior including maternal care, social parasitism, and foraging/hunting behavior. In some cases, the conceptual framework and predictions are already in place (e.g., social parasitism [73], development [74–79]), others can be enhanced by those generated for other taxonomic groups (e.g., venom evolution in snakes [80], genomic basis of diet shifts [81,82]).

Taxa that are underrepresented in genomics

Given the deep evolutionary distances between major hymenopteran families [83^{**}], selecting appropriate taxa for comparison presents a substantial challenge. Crown-group Hymenoptera originated 250–300 mya and spans evolutionary distances more than 3-fold those of modern birds [1,83^{**},84]. Bees shared a common ancestor ~100 mya, and bees + apoid wasps diverged from ants

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203 ~145 mya, similar to the split between marsupial and
204 placental mammals (>160 mya) [85]. Increasing the num-
205 ber of taxa sampled within each lineage, as well as the
206 total number of lineages sampled will provide higher
207 resolution to interpret genomic signatures of key pheno-
208 types. In addition, we suggest specific groups that may be
209 particularly appropriate for additional comparisons.

210 Expanding sampling of Hymenoptera genomes to
211 include one or several genomes per family would provide
212 a useful framework for future evolutionary studies within
213 the order. Although significant recent progress has been
214 made resolving higher-level phylogenetic relationships
215 [83^{**},86,87,88^{**}], important uncertainties remain at the
216 superfamily (Figure 1) and family levels, especially out-
217 side of the Aculeata. Having genomic data for all 97 fami-
218 lies would help overcome common phylogenetic prob-
219 lems, such as insufficient data, incomplete lineages
220 sorting, base composition bias, and long-branch attraction.
221 Multiple carefully chosen representatives of each family
222 would even further reduce issues related to poor taxon
223 sampling [83^{**}]. While producing phylogenies with entire
224 genomes is still too expensive for most multi-taxon stud-
225 ies, having a high quality genome for each family would
226 enable more economical reduced-representation phylo-
227 genomics across more taxa [89,90]. Having a complete
228 sampling of Hymenoptera genomes at the family level,
229 would also provide a better evaluation of genome size
230 variation and gene family evolution over the entire order.
231 For developmental biology examinations, species should
232 include samples that are readily available for experiments
233 such as embryological and larval time series sampling for
234 gene expression and localization.

235 Parasitoids are an immensely diverse group for those
236 interested in biological control applications and evolu-
237 tionary biology [91,92], however the genomes of only
238 14 species in five families have been published, with
239 draft genomes of ~30 more species (one additional fami-
240 ly) in progress (Figure 1, Table 1). To study polyembry-
241 onic development [78], genomes from the four families
242 have this trait and comparative genomics across these
243 families would be informative. Two have been
244 sequenced (Table 1, *M. cingulum* (Braconidae) and *C.*
245 *floridanum* (Encyrtidae)); thus including at least one
246 polyembryonic species from the Platygasteridae and Dry-
247 nidae families is a priority that would fit in the 0.05%
248 species sample. Because different families of parasitoids
249 use different taxa of hosts with very different biologies,
250 understanding mechanisms parasitoids use in overcoming
251 host immunity and host choice will benefit by having
252 genomes of at least one parasitoid species per family.
253 Including additional samples to allow comparisons of
254 parasitoids sharing the same host, such as *Drosophila*,
255 and pairing with genomes of the hosts will greatly enable
256 comparative studies across lineages and host/parasite
interactions [93].

257 Comparative genomic approaches to understanding the
258 evolution of eusociality will benefit from increasing the
259 depth of coverage within key wasp and bee families,
260 especially the Vespidae, Apidae and Halictidae. Each
261 of these families includes a diversity of social lifestyles,
262 along with closely related solitary individuals. In a few
263 key lineages such as vespid wasps and carpenter bees
264 (Apidae: Xylocopinae), there are also opportunities to
265 compare genomes within lineages in which the entire
266 spectrum of sociality occurs, from solitary to weakly social
267 to highly eusocial [71]. These highly informative species,
268 though underrepresented in previous genome sequenc-
269 ing efforts, have currently projects underway to fill these
270 gaps, with eventual insight into the full spectrum of
271 sociality evolution [94^{**}]. The sweat bees (Halictidae)
272 are also particularly useful for social evolution compara-
273 tive studies, representing at least two independent origins
274 of eusociality, a dozen lineages that have reverted back to
275 a solitary lifestyle [95–97], and with several species in the
276 genera *Halictus*, *Lasioglossum*, and *Megalopta* with faculta-
277 tive expression of eusociality (where individuals or popu-
278 lations vary in solitarily or socially nesting). Understand-
279 ing the genomic basis for *intra-specific* variation in social
280 behavior can illuminate the evolutionary processes that
281 may have been particularly important at the origins of
282 eusociality. We propose including ten additional samples
283 to enhance comparative genomics of social behavior
284 corresponding to environmental factors, such as latitude
285 or elevation [98–100], genetic influences [70^{*}] and *within*
286 population variation due to foundress reproductive strat-
287 egy [101] with dense sampling at fine phylogenetic scales.

288 Development of genomic resources for taxa from the non-
289 social aculeate wasp families will provide a critical picture
290 of the genomic substrate from which eusociality evolved,
291 and additionally provide insight into the genomic signa-
292 tures of other behavioral innovations, such as central place
293 foraging (i.e., nesting), maternal care, and social parasit-
294 ism. Particularly useful families include the spider wasps
295 (Pompilidae), velvet ants (aka ‘cow killers’) (Mutillidae),
296 thread-waisted wasps such as mud daubers (Sphecidae),
297 and cicada killers and bee wolves (Crabronidae), including
298 an independent origin of wasp eusociality). These groups
299 are closely related to vespids and bees, and exhibit
300 remarkable diversity in parental care, dietary breadth,
301 parasitism, and venom function. Additionally, because
302 wasp families are cosmopolitan and conspicuous, their
303 genomes can be studied in the context of well-described
304 behavior and natural history information.

Challenges

305 Several challenges may impede sequencing and assem-
306 bly, as well as annotation, of the genomes of Hymenop-
307 tera including sample quantity, heterozygosity, and avail-
308 ability; functional interpretation, and unknowns. Though
309 obtaining high quality DNA may be difficult because
310 samples are lacking (especially for parasitoids), for
311

Chalcidoidea, the US National Science Foundation has funded a phylogenetics project that could make available high molecular weight DNA from 388 species in 294 genera in 24 families (John M. Heraty, personal communication). For bees, pinned specimens for many groups may be available from the U.S. National Pollinating Insect Collection at the USDA-ARS-PWA Pollinating Insect Research Unit. In addition, ARS scientists are routinely collecting new material in North America and could provide genome quality specimens (recently collected and identified) to interested collaborators for some families found in North America. Additionally, some available samples (also many parasitoids) are very small and yield ≤ 100 ng of DNA per wasp. Although enough DNA for short-read sequencing library preparation, extractions from many individuals are needed to provide sufficient DNA for long-read technologies and heterozygosity in these pooled samples contributes to assembly challenges. Theoretically this can be solved by inbreeding for multiple generations, however some species cannot be inbred (e.g. those with complementary sex determination, in which homozygosity causes mortality or sterile diploid males). Concerning annotation, genes that underlie interesting differences in biology often evolve rapidly, making it difficult to find homologs in species where gene functions are well known. Therefore there is a need for experimentally determined functions (e.g. tissue specific expression and gene knockout/knockin). This is particularly true where differences in expression underlie differences in biology and distant actuators/enhancers and epigenetic modifications, may underlie differences in some key traits (e.g. diapause differences among *Nasonia vitripennis* populations [102*]). Finally, newly sequenced hymenopteran species may have genomic features that reduce genome assembly quality including large size, complex structure and high GC content.

A complete catalog is possible

Although many strategies can be envisioned for future hymenopteran genomics, we suggest a three-pronged strategy focused on breadth, biology and diversity. Increasing the breadth of genome coverage across taxa of this hyper-diverse order will provide long-term scientific benefits, we propose obtaining high quality genomes from at least one representative of each family (81 samples). For the 15 families with more than 50 genera or more than 100 species we propose sampling an additional 0.05% of the species (27 samples, Table 2). For families of particular interest such as pollinators and parasitic wasps, we propose adding 42 species, bringing the total to 150 or 0.1% of the species. Premier genomes are less expensive than in the past, but still expensive so this may not be possible for all of these samples. Short read assemblies or 30x comparative mapping studies can be used to fill in the phylogenetic sampling with additional closely related species. Including transcript sequencing (with high quality long read data) is a cost effective adjunct to enhance

the value of lower quality assemblies. With available samples, robust technologies and more reliable methods, this is a feasible task with the potential to impact studies for years into the future.

Uncited reference

[128*].

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.cois.2017.11.008](https://doi.org/10.1016/j.cois.2017.11.008).

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421 broad range of group sizes, degree of cooperation, and adaptations to
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667 likely to be successful for only the most conserved genomic elements like
668 ultraconserved elements (UCE). This excludes the most rapidly evolving
669 genomic features from analysis, and biases results toward conserved
670 genes. The authors of this study used UCE phylogenomics to generate a
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696 placed at ~281 mya. Sawflies and parasitoid wasps were each found to
697 be monophyletic, Trigonaloidea were identified as the sister group to
698 Aculeata and the evolutionary root of bees was located within the
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social trait.