

Draft genome of the globally widespread and invasive Argentine ant (*Linepithema humile*)

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Ants are some of the most abundant and familiar animals on Earth, and they play vital roles in most terrestrial ecosystems. Although all ants are eusocial, and display a variety of complex and fascinating behaviors, few genomic resources exist for them. Here, we report the draft genome sequence of a particularly widespread and well-studied species, the invasive Argentine ant (*Linepithema humile*), which was accomplished using a combination of 454 (Roche) and Illumina sequencing and community-based funding rather than federal grant support. Manual annotation of >1,000 genes from a variety of different gene families and functional classes reveals unique features of the Argentine ant's biology, as well as similarities to *Apis mellifera* and *Nasonia vitripennis*. Distinctive features of the Argentine ant genome include remarkable expansions of gustatory (116 genes) and odorant receptors (367 genes), an abundance of cytochrome P450 genes (>110), lineage-specific expansions of yellow/major royal jelly proteins and desaturases, and complete CpG DNA methylation and RNAi toolkits. The Argentine ant genome contains fewer immune genes than *Drosophila* and *Tribolium*, which may reflect the prominent role played by behavioral and chemical suppression of pathogens. Analysis of the ratio of observed to expected CpG nucleotides for genes in the reproductive development and apoptosis pathways suggests higher levels of methylation than in the genome overall. The resources provided by this genome sequence will offer an abundance of tools for researchers seeking to illuminate the fascinating biology of this emerging model organism.

Hymenoptera | invasive species | transcriptome | chemoreception | sociality

Ants are pivotal players in the Earth's terrestrial ecosystems. They include >14,000 described species, comprise about half of all insect biomass in the tropics, are the most important predators of insects and other arthropods, and turn and aerate more soil than earthworms (1). The diversity of lifestyles displayed by ants is equally impressive, including minute, acorn-dwelling species (and the ants that parasitize them), fungus-growing leafcutters with complex division of labor, trap-jaw ants that exhibit the fastest animal movements (2), slave-making ants that kidnap and enslave the young of other ants, and the countless teeming multitudes of army ants that prowl the leaf litter and subterranean habitats (1).

Some ant species have been introduced to new geographic ranges by human activities, and a few have emerged as damaging

and destructive invasive species (3). The Argentine ant (*Linepithema humile*) is one of the most widely distributed of these invaders and is established in nearly every Mediterranean-type climate in the world (4). Introduced Argentine ants form enormous "supercolonies," often across hundreds or thousand of kilometers (5), and workers from the largest supercolonies on different continents even accept each other as cononymates (6). At a local scale, the absence of aggression among workers within supercolonies allows them to direct resources toward colony growth (7) and attain high population densities. These introduced populations are then able to outcompete and eliminate native ants, which imperils plants and animals that normally interact with the native ants (5). In contrast, Argentine ants in their native South American range display high levels of aggression and genetic structure among colonies across substantially smaller spatial scales (8, 9). Argentine ants are also significant pests and thus are targets of heavy insecticidal control, leading to environmental contamination and harm to nontarget organisms (10).

Despite their importance and prominence, there are essentially no genomic resources for Argentine ants. Here we report the transcriptome (GenBank accession no. 46575) and de novo ge-

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Data deposition: The sequences reported in this paper have been deposited in the GenBank database [accession nos. 46575 (transcriptome), 45799 (genome), and 45805 (genome project)].

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nome sequence of the Argentine ant (GenBank accession no. 45799), generated using a combination of Roche 454 and Illumina sequencing. Project data are archived at the Hymenoptera Genome Database (http://hymenopteragenome.org/linepithema/genome_consortium). This project (GenBank accession no. 45805) represents an organizational advance in emerging model organism genomics (11), as it was not supported by a federal grant or genome sequencing center but, rather, was accomplished using small sums of discretionary funds provided by members of the insect genomics and Argentine ant research communities. This genome sequence provides a powerful resource for future analysis of gene families and phenotypes and candidate SNP markers for future population genetic and association mapping studies.

Results and Discussion

Genomic Features. We sequenced the Argentine ant genome ($2n = 16$) to $\sim 23\times$ coverage using source material from a single nest of the large California supercolony (Table 1 and *SI Appendix S1, Table S1*). The combined assembly of Roche 454 ($\sim 9\times$ coverage) and Illumina ($\sim 14\times$) sequencing resulted in 215.6 megabases (Mb) of scaffolded sequence (86% of the 250.8-Mb genome; ref. 12). We also recovered 12,516 bp of the mitochondrial genome in three scaffolds (*SI Appendix S1, Fig. S1*). We assembled the combined 454 and Illumina data using the Roche gsAssembler (454 Life Sciences) and Celera CABOG assemblers (13) (Table 1, *SI Appendix S1, Table S1*). Early 454-only assemblies contained homopolymer errors, but addition of Illumina sequence data produced marked improvements (Table 1) with an overall error rate estimated at 0.012% (*SI Appendix*). Overall, the Celera assembler yielded the best assembly, and addition of the Illumina data dramatically increased contig and scaffold length (Table 1). We also generated a 454 transcriptome sequence from ants of mixed age, caste, and geographic location and used it to train hidden Markov models for a custom MAKER (14) annotation pipeline (*SI Appendix S1, Table S2*).

The assembled genome appears to be relatively complete. First, our genome assembly captured 99% (246 of 248) of the core CEGMA genes (15), and 96% of them (239 of 248) were complete. Second, annotation of the cytoplasmic ribosomal protein genes revealed 83 genes, including the full set of 79 cytoplasmic ribosomal proteins (16, 17), and 4 duplicated genes (*RpS16*, *RpS23*, *RpS28*, and *RpS30*) (*SI Appendix S1, Table S2*). Third, annotation of the 67 nuclear-encoded oxidative phosphorylation genes shows that the *L. humile* genome assembly is only missing *cox7a* (*SI Appendix S1, Table S2*). This gene is also absent from the *Apis mellifera* genome, suggesting that it has been lost from the *L. humile* genome.

We generated a de novo repeat library of 523 elements and performed a whole-genome repeat annotation (Table 2). We found that 41 predictions were classified as retroint transposable elements (TEs) and 42 as DNA transposons, and 424 could not be classified. Although it has been hypothesized that the paucity of TEs in the *A. mellifera* genome is a product of eusociality, their abundance in the *L. humile* genome suggests otherwise. The *L. humile* genome also possesses remnants of viruses and viroids from a variety of families (*SI Appendix S1, Table S4*). The total repeat content was 28.7 Mb (13%), but without de novo repeat libraries, only 3.78% (8.3 Mb) of the *L. humile* assembly was identified as repetitive, with only 1.4% (3 Mb) identified as TEs. The *L. humile* genome was 62% AT, which is intermediate between *A. mellifera* (67%) and *Nasonia vitripennis* (58%). When combined with the 37.7 Mb of missing and ambiguous nucleotides, the total repetitive fraction (~ 66.4 Mb; 30.8% of the genome) was significantly higher than *A. mellifera* (10%), but lower than *N. vitripennis* (40%).

We described 16,123 genes (16,177 transcripts) in the *L. humile* official gene set (OGS1.1); 8,303 (51%) were supported by EST evidence, which confirmed 70% and 76% of the splice sites and exons for these genes, respectively. Of these genes, 1,364 were shared with *A. mellifera* and *N. vitripennis*, but not

Drosophila melanogaster (Fig. 1), and 589 were shared by the two aculeate Hymenoptera, *L. humile* and *A. mellifera*, but not with *Nasonia* (the outgroup genome for the Aculeata) or *Drosophila*. Although *L. humile* and *A. mellifera* are both social, Aculeata also contains solitary species. A total of 7,184 genes (45%) were unique to *L. humile* relative to these three other species.

We used Interproscan (18) and KEGG (19) (*SI Appendix S1, Table S5 and Fig. S2*) to identify putative functional domains and compared Gene Ontology (GO; *SI Appendix S1, Fig. S3*) terms for all *L. humile* genes relative to *D. melanogaster*, *A. mellifera*, and *N. vitripennis* (*SI Appendix S1, Tables S6 and S7*) and for the *L. humile*-specific genes. Of 16,123 genes in the OGS1.1, a total of 7,514 (44%) were annotated with at least one GO term (average = 3). Of the 7,184 (16%) genes unique to *L. humile*, 1,174 (Fig. 1 and *SI Appendix S1, Fig. S3*) had ontology terms. In the subset of genes found only in *L. humile*, 99 terms were enriched ($P < 0.05$) relative to all *L. humile* genes (*SI Appendix S1, Fig. S3*). These included odorant receptors, peptidases that may play a role in venom production, genes associated with lipid activities that could be involved in cuticular hydrocarbon (CHC) synthesis or catabolism, and DNA methylation genes, which may play a role in caste development.

Relative to other species, six cellular location terms were enriched, with most associated with the synapse ($P < 1.96E-04$) or postsynaptic membrane ($P < 4.74E-04$; *SI Appendix S1, Table S7*). Of the 17 genes associated with enriched molecular functions, 6 (all $P < 1.87E-08$ for GO categories) included cation binding and may be involved in neurological or other signal transduction ($P < 1.36E-04$) processes associated with odorant binding and olfaction, learning, memory, and behavior. Lipid catabolism ($P < 1.59E-05$), lipase ($P < 8.88E-03$), and phospholipid function (3; all $P < 0.04$) enrichments may include genes for the synthesis of CHCs involved in social communication. Electron transport, heme, and cation binding functions characteristic of cytochrome P450s were also enriched, consistent with the observed expansion of these genes.

The mean GC content of the *L. humile* genome was 37.7%, and the ratio of observed to expected CpG nucleotides [CpG(O/E)] was 1.55 (Fig. 2); both values are within the ranges reported for other Hymenoptera (*A. mellifera* and *N. vitripennis*; refs. 20 and 21). A comparison of GC compositional-domain lengths among

Table 1. Genome and sequencing statistics for three successive assemblies

	Roche gsAssembler	Celera V0.3	Celera V0.4
Total scaffolded sequence, bp*	199,810,258 (80%)	218,892,451 (87%)	215,552,578 (86%)
No. of scaffolds	3,093	3,180	3,030
No. of contigs	48,934	22,086	18,227
N50 contig size, bp	18,503 [†]	28,104	35,858
N50 scaffold size, bp	453,083	1,427,074	1,386,360
Total contig coverage	25 \times	14.6 \times	23 \times
Total reads used in scaffolds	38,902,428	21,970,969	46,779,980
454			
3-kb paired-end	1,613,759	1,276,042	1,255,603
8-kb paired-end	1,789,614	298,804	292,277
Unpaired	5,040,847	4,440,267	4,346,871
Illumina			
3-kb paired-end	NA	12,088,916	1,226,551
8-kb paired-end	NA	3,866,940	4,047,251
Unpaired	31,222,237	0	24,572,427

NA, not applicable.

*Values in parentheses indicate percent of the total genome size (12).

[†]Contigs > 500 bp; genome size, 250.8 Mb (0.26 pg).

Caste differentiation and division of labor within colonies are key innovations underlying the proliferation of ants, both numerically and taxonomically. Although the processes underlying caste differentiation are not fully understood, CpG DNA methylation is an important process for transcriptional regulation in many animals (27–29) and plays a role in the development of honey bee queens vs. workers (30). *L. humile* possesses a fully intact methylation toolkit with all three *Dnmt* genes (*SI Appendix S1*, Table S2 and Fig. S4). The only other sequenced insect genomes with the de novo methyltransferase *Dnmt3* are the pea aphid, *A. mellifera*, and *N. vitripennis* (20, 31). Interestingly, all three genes occur as single copies in *L. humile*, whereas *A. mellifera* and *N. vitripennis* have two and three copies of *Dnmt1*, respectively. Although inherited and de novo methylation have not yet been tested in *L. humile*, active methylation occurs in at least two other ant subfamilies (32).

To test for a genomic signature of CG methylation, we performed two dinucleotide analyses (*SI Appendix S1*, Figs. S5–S10 and Table S8) that revealed overall CG bias similar to *A. mellifera*, but no indication of exon or intron-specific methylation as seen in *A. mellifera* or *N. vitripennis*, respectively. We also found that dinucleotide transition SNPs associated with DNA methylation (i.e., CG↔TG) were 10-fold more prevalent in the *L. humile* SNP data compared with either of the transversions at this site (i.e., CG↔AG, CG↔GG). We observed 21,623 SNPs at the C position of the CG/TG sites, which should yield an expected 17,428 CG↔TG transition SNPs (80.6%) based on the overall observed genome rate of 4.15 transition mutations per transversion. However, we measured 18,611 CG↔TG transitions (86%), which indicated significant bias of CG↔TG SNPs relative to all other mutations at this site ($\chi^2 = 80.3$, $P < 0.001$). This bias in dinucleotide CG↔TG mutation relative to all other dinucleotide mutations is consistent with the variable distribution seen for the CG dinucleotide in the genome and suggestive of in vivo methylation (*SI Appendix S1*, Figs. S5, S6, and S10). Genes ranked with the largest numbers of CG↔TG SNPs included *major facilitator superfamily transporters*, male sterility proteins, several classes of zinc finger transcription factors, and several Ig superfamily cell adhesion proteins implicated in neuronal development (*SI Appendix S1*, Table S8).

Wing polyphenism and reproductive division of labor between queens and workers are two important features associated with eusociality in ants. The gene networks that underlie wing and reproductive development have evolved to be differentially expressed between winged reproductive castes and wingless sterile worker castes in response to environmental factors (33, 34). We found that the mean CpG(O/E) (35) for genes involved in reproductive system development ($n = 38$; mean = 1.21; $P < 0.05$) and apoptosis ($n = 18$; mean = 1.22; $P < 0.05$) were significantly lower than the mean CpG(O/E) (mean = 1.63) for 50 random genes resampled 10,000 times. (Fig. 2 and *SI Appendix*). Genes in the wing polyphenism network were not significantly different from the average gene CpG(O/E) ($n = 37$; mean = 1.50; $P = 0.4865$). The mean CpG(O/E) of the *Drosophila* orthologs (coding regions) that underlie wing development (mean = 0.95; $P = 0.86$), reproduction (mean = 0.98; $P = 0.98$), and apoptosis (mean = 1.00; $P = 0.99$) were not significantly different (*SI Appendix S1*, Fig. S11) than average gene CpG(O/E) in *Drosophila*. These results indicate that developmental genes in the reproduction and apoptosis networks have distinct germ-line methylation signatures relative to the rest of the genome, whereas genes in the wing polyphenism network do not.

Yellow genes occur in insects as well as some bacteria and fungi, but they are curiously absent in all noninsect metazoans (36, 37). The initially described *yellow-y* gene (*Y-y*) functions in cuticle pigmentation in *D. melanogaster* (38), but these genes have also been implicated in processes such as male courtship behavior (39), follicle cell function, and egg development (40). Proteins from a gene expansion of the ancestral *yellow-e3* gene in *A. mellifera* [the major royal jelly (*mrjpl*) subfamily] are involved in a regulating reproductive division of labor, but also have age-, sex-, caste-, and brain-specific expression (37, 41).

In the *L. humile* assembly we detected 10 *yellow* genes and 10 major royal jelly protein-like genes (Fig. 3C and *SI Appendix S1*, Fig. S14). The 8 *yellow* genes (*LhumY-y*, *-b*, *-c*, *-e*, *-e3*, *-g*, *-g2*, and *-h*) are similar to those of *D. melanogaster* and are likely orthologs. The remaining 2 are putative orthologs of the *yellow-x1* and *-x2* genes, which have been found only in the Hymenoptera. Interestingly, the *L. humile* genome contains an independent radiation of major royal jelly protein-like (*mrjpl*) genes similar to those in *A. mellifera* and *N. vitripennis*. The *mrjpl* and *mrjpl* gene sets of all three focal taxa each form their own strongly supported clade within the monophyletic *mrjpl* subfamily (Fig. 3C and *SI Appendix S1*, Fig. S14). These independent radiations in different hymenopteran lineages may indicate that the ancestral gene had a tendency to proliferate, allowing *mrjpls* to take on new functions and to respond quickly to new forms of selection (42).

Invasion Biology. Although Argentine ants are a widespread and familiar invasive species, many details of their invasion biology remain shrouded in mystery. To clarify some of the biological

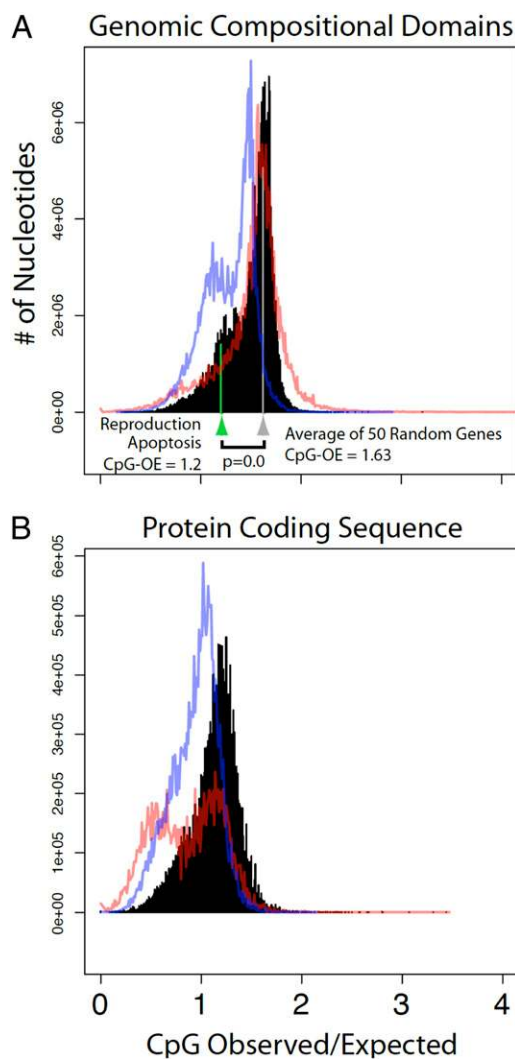


Fig. 2. Distribution of CpG(O/E) predictions in genomic compositional domains (A) and protein coding regions (B) for *L. humile* (black), *A. mellifera* (red line), and *N. vitripennis* (blue line). The combined average CpG(O/E) for all genes involved in either reproductive (mean = 1.21) or apoptosis processes (mean = 1.22) (green arrow and line) is significantly lower than the average CpG(O/E) for 50 random genes resampled 1,000 times as a control (gray arrow and line; $P < 0.05$, statistical randomization test).

processes that may be related to their invasive success, we annotated and analyzed genes that are likely to be involved in Argentine ant immune processes, insecticide resistance, and dietary detoxification. We also identified a large number of candidate SNP markers that will be valuable tools for identifying source populations in the Argentine ant's native range, reconstructing the history of introductions around the world, and identifying current routes of transport.

The Argentine ant's unicolonial social structure allows introduced populations to attain enormous population densities in its introduced range. However, it is not clear how Argentine ants control the proliferation of pathogens and parasites in these high-density populations, particularly in light of the extremely low levels of genetic diversity that typify introduced populations (5, 8). We annotated immune genes primarily from the *Toll*, *Imd*, *JNK*, and *JAK/STAT* signaling pathways. In total, 90 of 202 immune genes had reciprocal best matches (SI Appendix S1, Tables S2 and S16). Of these 202 genes, 152 are present in the *Drosophila* genome and 78 in *Apis*. Thus, *L. humile* appears similar to *Apis* and *Nasonia* in having markedly fewer immune genes than *Drosophila*. Moreover, the many hygienic behaviors and chemical secretions described for the social Hymenoptera may play key roles in controlling pathogens.

Introduced populations of Argentine ants are also frequently targets of heavy insecticidal control, suggesting that genes conferring insecticide resistance may be under strong positive selection. Like many invasive ants, Argentine ants are also extreme dietary generalists, and this plasticity may require an underlying ability to detoxify many different components of their food sources. Cytochrome P450s are a family of heme-thiolate enzymes that catalyze a diverse array of chemical reactions in nearly all organisms (43), and they have been implicated in both insecticide resistance and dietary detoxification.

The *L. humile* genome encodes 111 cytochrome P450s (SI Appendix S1, Table S2 and SI Appendix S2, Fig. S13), substantially more than *A. mellifera* (46 genes) (44) and *N. vitripennis* (92 genes) (45). It has been hypothesized that the paucity of P450 genes in the genome of *A. mellifera* is a product of its eusocial life history (44), but their abundance in *L. humile* suggests that eusociality alone cannot explain their scarcity in *Apis*. The *L. humile* genome encodes 69 CYP3 clan P450s (SI Appendix S2, Fig. S13), the most in any sequenced insect genome. CYP3 P450s are associated with oxidative detoxification of xenobiotics (46), and their abundance

in *L. humile* may be an adaptation to a variety of toxins encountered in the diet of this generalist ant, compared with the rather specialized diet of the honey bee.

To facilitate the development of population genetic markers for future analyses of the origin, history, and movement of Argentine ants, we scanned the genome for SNPs. Because the source material contained a single queen pupa and ~100 workers from a single nest (SI Appendix S1, Table S1), we were able to identify predicted polymorphic sites. We discovered 231,248 SNPs (0.9 SNP/kb) that occurred in at least three reads and at least 10% of overlapping reads. A total of 5,734 genes (36%) had at least 1 SNP, with 14,136 SNPs (6%, 0.7 SNP/kb) in exons, 26,720 (12%, 0.9 SNP/kb) in introns, 66,830 (29%, 2.8 SNP/kb) in annotated repeats, and 123,492 (53%, 1.1 SNP/kb) in nonrepeat intergenic regions. Although 77% of exonic SNPs would be nonsynonymous (23% synonymous) if all substitutions were equally likely, we observed that only 54% of SNPs were nonsynonymous and 46% were synonymous ($P < 0.001$). When each type of SNP was normalized to the total number of possible sites for that type, we observed 4.9×10^{-4} SNPs per nonsynonymous site and 1.9×10^{-3} SNPs per synonymous site. As expected, we saw a 3.87-fold excess of synonymous SNPs in the OGS1.1 gene set.

We also manually investigated the genes with the highest number of SNPs per kb of exon sequence (SI Appendix S1, Table S3). Interestingly, six *L. humile* genes with Interproscan-predicted functions similar to male sterility and gametogenesis genes in *Drosophila* showed a high degree of polymorphism in *L. humile* (24–38 SNPs/kb). These polymorphic genes may be under sexual selection or diversifying selection and could be useful for studying different patriline lineages in native vs. invasive ants. Cytochrome P450s, lipid metabolism genes, and transcription factors were also ranked highly, with >75 SNPs in the exon and intronic regions.

Conclusion. With the sequencing and annotation of the Argentine ant genome and the development of associated genomic resources, this species is well positioned to become a model organism in which powerful genetic approaches can be coupled with a wealth of natural history and behavioral and ecological knowledge. Given the immense financial and ecological costs associated with introductions of Argentine ants, these tools will likely find widespread application and produce tangible benefits for agriculture, societies, and ecosystems. Finally, the evolutionary forces driving some of the unusual and remarkable genomic patterns reported here

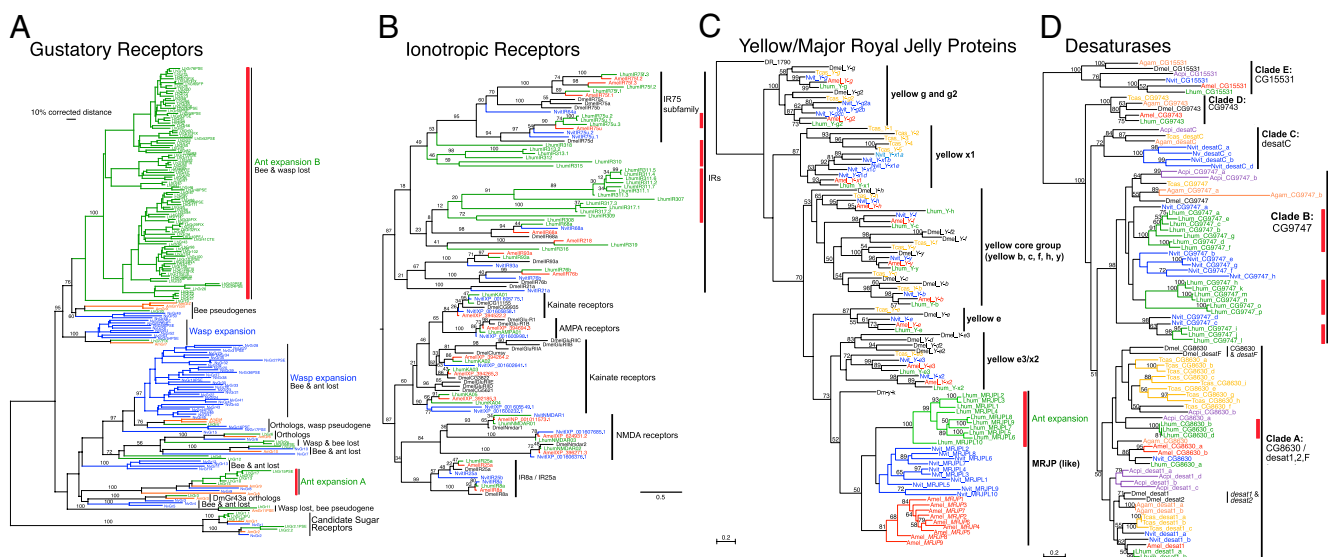


Fig. 3. Gene expansions in the Argentine ant genome. (A) Gustatory receptors. (B) Ionotropic receptors. (C) Yellow/major royal jelly proteins. (D) Desaturases. Vertical red bars indicate *L. humile* gene duplications and expansions. The *L. humile* genome also possesses an enormous expansion of odorant receptors (Ors; SI Appendix S2, Fig. S12). Green, *L. humile*; red, *A. mellifera*; blue, *N. vitripennis*; black, *D. melanogaster*.

remain unknown and will be productive avenues for future research that explores the basis of eusociality and the causes and consequences of biological invasions.

Materials and Methods

Source Material. We collected an Argentine ant colony fragment from a residential orchard in Santa Clara County, CA. We confirmed that these ants belong to the large supercolony that dominates the introduced range in California using behavioral assays, microsatellite genotyping, and analysis of CHC profiles (*SI Appendix S1, Table S14 and Fig. S17*).

Library Preparation and Sequencing. Transcriptome. cDNA was generated from mixed source material and sequenced using Roche 454 Genome Sequencer LR70 FLX technology (Table 1 and *SI Appendix S1, Table S1*). This process yielded ~128 Mb of DNA sequence, which was assembled into 20,070 contigs. **Genomic.** Genomic DNA was extracted and purified from a single queen pupa (Saratoga) and sequenced using seven runs of 454 FLX Titanium sequencing. Additional worker-derived genomic libraries were constructed and sequenced on the 454 and Illumina platforms.

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Assembly. We created several assemblies using Newbler and CABOG assembler (13), as described in *SI Appendix*.

Annotation. We first used the automatic annotation pipeline MAKER (14) to annotate the genome of *L. humile*. We also manually annotated ~1,000 genes (*SI Appendix S1, Table S2*), including a number that are related to fundamental biological processes (oxidative phosphorylation, ribosomal), complex behaviors (learning, memory and aggression), sensory biology (vision, chemoreception), insecticide resistance, immunity, and developmental networks (wings, reproduction).

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