Comparative Genomics Reveals Recent Adaptive Evolution in Himalayan Giant Honeybee *Apis laboriosa*

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Accepted: 26 September 2021

Abstract

The Himalayan giant honeybee, *Apis laboriosa*, is the largest individual honeybee with major ecological and economic importance in high-latitude environments. However, our understanding of its environmental adaptations is circumscribed by the paucity of genomic data for this species. Here, we provide a draft genome of wild *A. laboriosa*, along with a comparison to its closely related species, *Apis dorsata*. The draft genome of *A. laboriosa* based on the de novo assembly is 226.1 Mbp in length with a scaffold N50 size of 3.34 Mbp, a GC content of 32.2%, a repeat content of 6.86%, and a gene family number of 8,404. Comparative genomics analysis revealed that the genes in *A. laboriosa* genome have undergone stronger positive selection (2.5 times more genes) and more recent duplication/loss events (6.1 times more events) than those in the *A. dorsata* genome. Our study implies the potential molecular mechanisms underlying the high-altitude adaptation of *A. laboriosa* and will catalyze future comparative studies to understand the environmental adaptation of modern honeybees.

Key words: comparative genomics, honeybee, *Apis laboriosa*, whole genome sequencing, evolution.

Significance

The accumulation of knowledge of adaptively evolutionary mechanisms is of great significance for developing strategies for protecting Himalayan giant honeybee, *Apis laboriosa*. We presented high-quality draft genome sequences of *A. laboriosa* for the first time, along with a comparative genomics analysis with its closely related species, *Apis dorsata*. Our study may advance understanding of the molecular basis of environmental adaptation in *A. laboriosa* and provide valuable resources for future comparative/population studies of honeybee evolution.

Introduction

The Himalayan giant honeybee *Apis laboriosa*, the largest individual bee in the genus *Apis*, is a key pollinator and important honey producer in Himalayan regions (Joshi et al. 2004). *A. laboriosa* lives in mountainous areas from Nepal to southwest China and has evolved adaptive behaviors to cope with harsh environments: nesting on inaccessible cliffs above 1,200 m, migrating seasonally and foraging at low temperatures (Batra 1996). To comprehend its evolutionary dynamics and adaptations to the local environment, *A. laboriosa* is usually compared with the closely related species *Apis dorsata*, a lowland giant honeybee that is widespread throughout tropical and subtropical Asia. These two species were grouped into a “giant honeybees” clade in previous phylogenetic studies (Willis et al. 1992; Engel and Schultz 1997; Arias and Sheppard 2005; Chhakchhuak et al. 2016). Although *A. dorsata* genome has been sequenced recently (Oppenheim et al. 2020), whole-
genomic data for *A. laboriosa* are still lacking. An in-depth understanding of the evolution process of *A. laboriosa* necessitates a high-quality reference genome.

In this study, we presented high-quality draft genome sequences of *A. laboriosa*. We compared the draft genome of *A. laboriosa* and *A. dorsata* to describe the recent evolutionary trend because their divergence. The genome sequence and analyses will pave the way for future comparative studies and a mechanistic understanding of honeybee evolution.

**Results**

*Apis laboriosa* and *A. dorsata* have Similar Genome Compositions

A total of 74.16 gigabases (Gbp) of paired-end DNA reads were generated from whole genome shotgun sequencing of *A. laboriosa* worker bees. The short reads were de novo assembled into 4,376 scaffolds (total size: 226.1 Mbp, N50: 3,339,770 bp; fig. 1A). The GC content of the *A. laboriosa* assembly was 32.2% (fig. 1A), similar to that of *A. dorsata* (31.9%). A genome completeness assessment (fig. 1A) showed that the assembly of *A. laboriosa* was 99.2% complete, similar to the *A. dorsata* assembly (98.9%). The predicted gene number in *A. laboriosa* genome (11,466 genes) is higher than that in the *A. dorsata* genome (9,910 genes) (fig. 1A). Additionally, an analysis of repeat elements (fig. 1B) showed that the genome of *A. laboriosa* contained a higher proportion of transposable elements than that of *A. dorsata* (*A. laboriosa*: 15,518,024 bp, 6.86% of the genome; *A. dorsata*: 14,367,696 bp, 6.24% of the genome).

Subsequently, we examined the shared gene families between these two species. A total of shared 8,404 gene families were identified based on the gene sets from *A. laboriosa* (9,857 of 11,466 genes), *A. dorsata* (9,001 of 9,910 genes), and seven other Hymenoptera species (fig. 1C). All these gene families were further classified into homologous groups. **Table 1** lists the summary statistics of the genome assembly, GC%, completeness and gene prediction of two species genomes.

**Table 1** Summary statistics of genome assembly, GC%, completeness and gene prediction

| Statistics                              | *Apis laboriosa* | *Apis dorsata* |
|-----------------------------------------|------------------|----------------|
| Size, bp                                | 226,033,288      | 230,340,171    |
| Scaffold, num                           | 4,376            | 4,040          |
| N50 scaffold, bp                       | 3,339,770        | 732,052        |
| Min scaffold, bp                       | 200              | 200            |
| Median scaffold, bp                    | 441              | 356            |
| Mean scaffold, bp                      | 51,509           | 57,614         |
| Max scaffold, bp                       | 9,783,586        | 3,998,413      |
| GC composition                         | 32.2%            | 31.9%          |
| Genome Completeness                    | 99.2%            | 98.9%          |
| Gene prediction                         |                   |                |
| Genes, num                              | 11,466           | 9,910          |
| Min genes, bp                          | 63               | 67             |
| Median genes, bp                       | 1,116            | 1,236          |
| Mean genes, bp                         | 1,663            | 1,667          |
| Max genes, bp                          | 59,058           | 57,621         |

**Fig. 1.**—The summary for genome assembly, GC%, genes, gene families, repeats, homologous groups, nucleotide divergence and phylogeny relationship of *Apis laboriosa* and *A. dorsata* genome. (A) Summary statistics of the genome assembly, GC%, completeness and gene prediction of two species genomes; (B) Bar charts describing the repeat elements of two genomes. ** indicates a statistical significance (Chi-squared test, *P* < 0.05) between two genomes; (C) Venn diagram describing the number of gene families and genes in two genomes; (D) Bar charts and pie charts describing homologous groups of two genomes in different categories. ** indicates a statistical significance (Chi-squared test, *P* < 0.05) between two genomes; (E) Summary statistics (mean ± standard deviation/SD) of genetic distances between single-copy genes from *Apis laboriosa* genome and *Apis dorsata* genome. The distances were calculated using R package ‘ape’; (F) Phylogenetic tree of the nine Hymenoptera species. The tree was reconstructed using maximum-likelihood method under the GTR model based on the concatenated alignments of the single-copy genes and the dating time with 95% Confidence Interval was assessed using Markov chain Monte Carlo analysis under the GTR model based on the concatenated alignments of fourfold degenerate sites from single-copy genes. The bootstrap values of all nodes were 100%. *Apis laboriosa* was indicated by pink label, and *A. dorsata* was denoted by blue label. Branch length represented time of divergence in million years.
families were further classified into different categories based on the phylogenetic composition and the copy number of the genes in each species (see Materials and Methods). We found that *A. laboriosa* and *A. dorsata* presented similar frequency distribution of gene family categories (*χ²* test, *P* > 0.05; fig. 1D), indicating that the genomes of these two species shared a high amount of core gene families.

The nucleotide divergency in single-copy genes between *A. laboriosa* and *A. dorsata* was around 0.03 with relatively large standard deviations (P-distance: 0.028 ± 0.061; JCG: 0.030 ± 0.020; K80: 0.031 ± 0.022; F84: 0.031 ± 0.021) (fig. 1E), demonstrating a close evolutionary relationship of these two species. The estimated divergence time between *A. laboriosa* and *A. dorsata* was 2.37 million years (Myr) (95% CI 1.9–4.3 Myr) (fig. 1F).

**Genes in *A. laboriosa* had Undergone More Frequently Positive Selection**

We performed a series of quality control procedures at gene/sites levels to achieve more accurately estimated selective pressure (fig. 2A, see Materials and Methods for details). The result showed that the synonymous substitution rate, *dS* values, in 6,718 single-copy gene families (core-gene ortholog families) were similar between these two species using PAML branch model (free-ratios) (*A. laboriosa*: 0.022 ± 0.026; *A. dorsata*: 0.019 ± 0.018; fig. 2B), indicating a similar neutral nucleotide substitution rate with a relatively large random fluctuations in these two lineages.

To test whether the evolutionary process of the coding regions in both species was deviated from the neutral evolution because the divergence from the latest common ancestor, we analyzed the selective pressure of single copy genes. The BUSTED branch-site models and likelihood ratio tests showed that compared with *A. dorsata*, a statistically significantly higher amount of genes in *A. laboriosa* have undergone positive selection at both gene families level (742 vs 291, *χ²* test: *P* < 0.001) (fig. 2C) and functional categories level (a total of nine enriched functional categories in the *A. laboriosa* genome, *χ²* test: adjusted *P* < 0.1; fig. 2D). Around 88.9% and 77.8% enriched gene ontology (GO) categories with positively selected genes are overlaid (697 and 507 genes).
Gene Families in *A. laboriosa* had Undergone More Frequent Duplications

We detected a total of 515 and 85 duplication or loss (DL) events in the *A. laboriosa* and *A. dorsata* genomes, respectively, by reconciling 8,404 gene trees with the reference species tree (fig. 1f) using RANGERDTL v2.0. We found that there was a significantly higher frequency of recent duplication events in *A. laboriosa* than in *A. dorsata* (438 vs 54 genes, \( \chi^2 \) test: \( P < 0.001 \); fig. 2E).

Furthermore, 39 positive selection events in these recently duplicated genes were identified in the *A. laboriosa* genome, whereas only four such events were identified in the *A. dorsata* genome (fig. 2F). For instance, key positively selected sites can be detected in the duplicated genes related to DNA binding or G protein-coupled receptor activity in *A. laboriosa* (fig. 2G).

We finally investigated the enriched/depleted functional categories of duplication/loss events in *A. laboriosa* compared with those in *A. dorsata* (fig. 2H). A total of 20 statistically significantly (\( \chi^2 \) test, adjusted \( P < 0.1 \)) GO terms with enriched duplications were detected. The recent evolution of these gene families in *A. laboriosa* potentially contributes to the adaptation of harsh living environment with strengthened pollination, enhanced learning, foraging, etc. (Honeybee Genome Sequencing Consortium 2006; Scheiner et al. 2006). The pattern of significantly higher proportion of recent duplication in *A. laboriosa* was confirmed by another duplication-loss reconciliation software, TREERECS (90% of enriched functional categories identified by RANGERDTL v2.0 were also identified by TREERECS; supplementary fig. S4, Supplementary Material online).

Several caveats are worth discussing. First, the small divergence between the genomes of *A. laboriosa* and *A. dorsata* (fig. 1E) could potentially introduce a high amount of random errors into the results. The genetic drift in a small population (e.g., giant bee clade) further complicated the interpretation of potential adaptive selection, with a high rate of false positives and false negatives in the positive selection and gene duplication/loss analyses. Also, the sequencing errors, alignment errors (Anisimova 2001), gene tree discordance (Mendes and Hahn 2016), polymorphism, and multinucleotide mutations (Venkat 2018) could introduce bias into positive selection analysis. Thus, we adopted several filtering procedures the minimize the bias and achieve more accurately estimated positive selection (fig. 2A). Besides, we found that the enrichment pattern at functional categories level were consistent with both PAML branch-site model and a branch-site model that incorporates multinucleotide mutations (HYPHY MNMs) (supplementary figs. S2 and S3A, B, Supplementary Material online), which further indicated the robustness of our positive selection analysis.

In conclusion, our comparative study has identified some genomic evolutionary processes that are possibly related to the local adaptation of wild *A. laboriosa*. In the future, more in-depth comparative and population-level studies will help to describe the landscape of local adaptation in this species, thereby facilitating the development of feasible strategies for protecting this important pollinator.

Materials and Methods

Sample Collection, DNA Extraction, Library Preparation, and Sequencing

Four *A. laboriosa* workers were collected from the same wild colony in Shangri-La, Yunnan Province, China in February 2019 (supplementary fig. S1A, Supplementary Material online). DNA was extracted from its thorax by CTAB immediately for library preparation. A total of 0.2 μg DNA was used for each sample as the input material for DNA library preparation. Sequencing library was generated using NEB Next Ultra DNA Library Prep Kit for Illumina (NEB) following the manufacturer’s recommendations (see supplementary data, Supplementary Material online for more details).

Whole genome shotgun sequencing was applied to generate short paired-end reads (150 bp) libraries with a series of short-insert length (11.05 and 35.79 G for 250 and 500 bp, respectively) and long-insert length (14.01 and 13.30 G for 2 and 5 kb, respectively). The sequencing was performed on a HiSeq platform (Illumina, San Diego, CA).

Quality Control and Genome Assembly

The quality control of raw reads (removed the adapters, low-quality bases/reads and PCR duplicates) was performed as previously described (Heshiki et al. 2017; Zheng et al. 2019). Afterwards, SOAPdenovo v2.04 (Li et al. 2010) was used to perform the de novo genome assembly by constructing and simplifying de Bruijn graph with default parameters (max_r-d_len [150]; avg_ins [350]; reverse_seq [0]; asm_flags [3]; pair_num_cutoff [3]; map_len [32]). BUSCO v4 (lineage_dataset [metazoa_odb9]; mode [genome]) was used to assess the genome assembly completeness (Simão et al. 2015).

Predictions of Gene and Repeats

Genes were predicted via a strategy combining ab initio gene prediction and homology-based gene prediction (Li et al. 2019). The coding sequences (CDS) of *A. laboriosa* were identified using EvidenceModeler v1.1.1 (weight of ab initio gene predictions: 8; weight of homology-based gene predictions: 10) (Haas et al. 2008) by incorporating gene predictions results (see supplementary data, Supplementary Material online for more details).
The transposons of *A. laboriosa* and *A. dorsata* genome were identified using RepeatMasker v4.1 (Tarailo-Graovac and Chen 2009) with the library “honeybee” and the default cutoff “225.” The identified transposons were grouped according to the annotated repeat class information.

Construction and Annotation of Gene Families

The gene families among *A. laboriosa* and eight *Hymenopterans* species were constructed using OrthoMCL v2.0.9 (Li et al. 2003) with default parameters (dbVendor [mysql]; percentMatchCutoff [50]; evalueExponentCutoff [−5]). Gene families were further classified into 11 categories based on the copy number and presence/absence information of genes in various bee lineages: 1) single-copy gene families among all nine *Hymenopterans*, among all seven bee species, among all five honeybee species, or among “giant honeybees” cluster; 2) nonsingle-copy gene families containing all nine *Hymenopterans*, all seven bee species, all five honeybee species, or “giant honeybees” cluster; 3) other shared gene families; 4) unshared gene families (*Apis laboriosa* and *A. dorsata* genes are not present simultaneously); and 5) singletons (fig. 1D).

GO annotations of *A. laboriosa* genes were performed using InterProscan v5.29 (Jones et al. 2014) with default parameters (appl [Pfam]; goterms). The GO annotation of the *A. laboriosa* genes within a gene family was used to represent the function of that gene family.

Construction of Species Tree and Estimation of Divergence Time

Phylogenetic trees of *A. laboriosa* and eight *Hymenopterans* species were reconstructed based on the single-copy genes using the maximum likelihood method via PhyML v3.0 package (Guindon et al. 2010), with the parameters of 1,000 bootstrap replicates, GTR model with gamma-distributed rate heterogeneity. The topology of the resulting species tree (fig. 1F) was consistent with the phylogenies in previous studies (Arias and Sheppard 2005; Kapheim et al. 2015; Chhakchhuak et al. 2016; Branstetter et al. 2017, 2018; Diao et al. 2018).

Genetic distance between *A. laboriosa* and *A. dorsata* was calculated based on the single-copy genes using R “ape” package. Divergence time among species was estimated based on the concatenated alignments of 4-fold degenerate sites from single-copy genes using MCMCtree in PAML v4.9 (Yang 2007) (see supplementary data, Supplementary Material online for more details).

Analysis of Positive Selection

Positive selection drives the adaptive evolution of the species through promoting the spread of beneficial alleles in the population (Sabeti et al. 2006). In this study, we assessed substitution rates using branch model (seqtype [1]; clock [0]; model [1]; NSsites [0]; fix_omega [0]; fix_blength [0]) in PAML v4.9 package (Yang 2007) and gene-wide positive selection signals using BUSTED (Branch-site Unrestricted Statistical Test for Episodic Diversification) (Murrell et al. 2015) in HyPhy (Hypothesis testing using Phylogenies) (Pond et al. 2005) in both *A. laboriosa* and *A. dorsata* branches after the divergence from their latest common ancestor.

To identify a more robust pattern of positive selection, two preprocessing and one postprocessing procedures were adopted for the calculation of selective pressure using the PAML branch models and HYPHY BUSTED branch-site models (fig. 2A): 1) masked the sites (without consideration of substitutions between two giant honeybees) with minor allele frequencies ≥0.125 (1/8) for *A. laboriosa* and ≥0.05 (1/19) for *A. dorsata* to avoid the false substitution induced by polymorphisms; 2) masked the substitution sites with coverage <20× of uniquely and accurately mapped paired-end reads in *A. laboriosa* genome to avoid substitutions resulted from low quality or misassembled sites/regions.

To determine the influence of discordance between species tree and gene tree on the positive selection, we compared the results of positive selection with a phylogeny either consisting of all nine *Hymenopterans* species, or only seven *Hymenopterans* species (two closely related species, *Apis mellifera* and *Apis cerana* were removed to avoid unresolved phylogeny). Only nine (1.4%) positively selected genes in the *A. laboriosa* genome were uniquely detected by using the phylogenetic trees consisting of all nine *Hymenopterans* species, suggesting a very low false-positive rate produced due to the discordance between species and gene trees.

Detection of Gene Duplications and Losses

Inferring the DL (D: duplication; L: loss) events within a given gene family is important to understand the macro evolution of a gene family (Bansal et al. 2018). In this study, RANGER-DTL v2.0 (Bansal et al. 2018) was employed to deduce all putative DL events related to *A. laboriosa* and *A. dorsata* by reconciling the reconstructed gene trees with the species tree (fig. 1F) using the cost parameters which only considers gene duplication/loss (duplication cost [2]; transfer cost [1,000]; loss cost [1]). TREERECS (Comte et al. 2020) was also adopted to confirm the robustness of enrichment/depletion patterns for the duplication/loss events (duplication cost [2]; loss cost [1]) (supplementary fig. S4, Supplementary Material online). For each gene family, after the multiple alignment of CDS using MUSCLE (Edgar 2004), a gene tree was constructed using PhyML v3.0 (Guindon et al. 2010) with bootstrap number of 100 and substitution model of “GTR” (see supplementary data, Supplementary Material online for more details).
Statistical Analysis
Chi-squared tests were used to detect the significant differences regarding repeats, homologous groups, and evolutionary events between A. laboriosa and A. dorsata. Categories with a false discovery rate (Benjamini–Hochberg procedures) below 0.1 were deemed to be significant.

Supplementary Material
Supplementary data are available at Genome Biology and Evolution online.

Data Availability
The drafted genome of the wild A. laboriosa is available at NCBI BioProject PRJNA647849. All command lines and configuration files for programs used in this study are available at https://github.com/lindan1128/Apis-Laboriosa-Project.

Acknowledgments
This project was supported by the APRC grant (9676008) from the City University of Hong Kong, and the earmarked fund for China Agriculture Research System (No. CARS-44-KX21). We thank Prof. Haiou Kuang from Yunnan Agricultural University for giving the help to collect samples.

Author Contributions
J.X. and J.L. designed this study. P.S. performed the experiments. D.L. and L.L. analyzed the data. D.L. drafted the manuscript. All authors commented on the manuscript and agreed to the submission.

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Associate editor: Dennis Lavrov