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Sandra M. Rehan

University of New Hampshire, Durham, sandra.rehan@unh.edu

Karl M. Glastad

Georgia Institute of Technology

Sarah P. Lawson

University of New Hampshire, Durham

Brendan G. Hunt

University of Georgia

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The Genome and Methylome of a Subsocial Small Carpenter Bee, *Ceratina calcarata*

Sandra M. Rehan^{1,*†}, Karl M. Glastad^{2,†}, Sarah P. Lawson¹, and Brendan G. Hunt³

¹Department of Biological Sciences, University of New Hampshire, Durham

²School of Biology, Georgia Institute of Technology

³Department of Entomology, University of Georgia

[†]These authors contributed equally to this work.

*Corresponding author: E-mail: sandra.rehan@unh.edu.

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Abstract

Understanding the evolution of animal societies, considered to be a major transition in evolution, is a key topic in evolutionary biology. Recently, new gateways for understanding social evolution have opened up due to advances in genomics, allowing for unprecedented opportunities in studying social behavior on a molecular level. In particular, highly eusocial insect species (caste-containing societies with nonreproductives that care for siblings) have taken center stage in studies of the molecular evolution of sociality. Despite advances in genomic studies of both solitary and eusocial insects, we still lack genomic resources for early insect societies. To study the genetic basis of social traits requires comparison of genomes from a diversity of organisms ranging from solitary to complex social forms. Here we present the genome of a subsocial bee, *Ceratina calcarata*. This study begins to address the types of genomic changes associated with the earliest origins of simple sociality using the small carpenter bee. Genes associated with lipid transport and DNA recombination have undergone positive selection in *C. calcarata* relative to other bee lineages. Furthermore, we provide the first methylome of a noneusocial bee. *Ceratina calcarata* contains the complete enzymatic toolkit for DNA methylation. As in the honey bee and many other holometabolous insects, DNA methylation is targeted to exons. The addition of this genome allows for new lines of research into the genetic and epigenetic precursors to complex social behaviors.

Key words: small carpenter bee, epigenetics, maternal care, social evolution, insect genome.

Introduction

The goal of sociogenomics is to explore the molecular genetic basis of social life (Robinson et al. 2005). This field was spearheaded by sequencing the first genome of an eusocial species, the honey bee, *Apis mellifera* (Honeybee Genome Sequencing Consortium 2006). Following the publication of the honey bee genome, researchers sequenced the genomes of many other eusocial insect species and their solitary relatives, including seven ants (Bonasio et al. 2010; Nygaard et al. 2011; Smith, Smith, et al. 2011; Smith, Zimin, et al. 2011; Suen et al. 2011; Wurm et al. 2011) and nine additional bees (Kocher et al. 2013; Kapheim et al. 2015; Sadd et al. 2015). These genomes revealed much about the importance of gene family expansions, chemical communication, and epigenetics to caste determination (Gadau et al. 2012; Simola et al. 2013; Yan et al. 2014). Although we are beginning to elucidate the genetic basis of highly eusocial behaviors, hallmarked by overlapping

generations, reproductive division of labor, and cooperative brood care, we still understand very little about the genetic architecture of incipient sociality. Indeed, identification of the genetic precursors to advanced eusociality necessitates a comparison of highly social species with those species possibly transitioning to highly social behaviors.

A prominent theory in the transition to sociality is that incipient groups consisted of a mother and her offspring (Alexander et al. 1991). Maternal care is thus a key step in the evolution of social behavior (Michener 1974; Anderson 1984; Gadagkar 1990). Subsocial species often display behaviors thought to be necessary precursors to highly social behavior, such as maternal care, nest loyalty, cohabitation, and adult longevity (Sakagami and Maeta 1977; Tallamy and Wood 1986; Crespi 1994; Costa 2006; Wilson 2008). To date, there are published genomes of ten bee species (Kocher et al. 2013; Kapheim et al. 2015; Sadd et al. 2015), but there are no sequenced genomes of a subsocial bee species.

Because the evolutionary path from solitary and eusocial behavior likely involves a subsocial intermediate, comparisons between solitary and subsocial species closely related to eusocial taxa may provide evidence of the genetic scaffolding for more complex social behaviors (West-Eberhard 1967; Rehan and Toth 2015). *Ceratina* small carpenter bees (Hymenoptera: Apidae) are a diverse and cosmopolitan genus (Michener 2007). Most species of this genus are considered solitary, but sociality occurs naturally in some species (Sakagami and Laroca 1971; Sakagami and Maeta 1984; Rehan et al. 2009; Rehan et al. 2014) and can be induced artificially in others (Chandler 1975; Sakagami and Maeta 1987, 1989, 1995). This natural variation provides a unique opportunity to explore the transition from solitary to social life (Michener 1985). *Ceratina* are members of the subfamily Xylocopinae in the bee family Apidae and provide a phylogenetic contrast and independent origin of eusociality to honey bees and bumble bees, members of the subfamily Apinae (Cardinal and Danforth 2011; Rehan et al. 2012).

Ceratina calcarata (Hymenoptera: Apidae) is a small carpenter bee native to eastern North America (Rehan and Sheffield 2011; Shell and Rehan 2016). Unlike solitary species, such as the leaf cutter bee *Megachile rotundata*, *C. calcarata* displays prolonged maternal care, mother–adult offspring interactions, and mutual tolerance (Rehan et al. 2014). *Ceratina* females make their nests in the pith of dead, broken twigs (Daly 1973). The mother lays eggs within the stem and provides each offspring with its own nutrition-rich pollen ball (McFrederick and Rehan 2016). As the offspring grow, the mother will protect and groom her offspring (Michener 1985; Rehan and Richards 2010a). *Ceratina calcarata* females are long-lived and nest loyal, and provide care to their offspring throughout development and even into adulthood (Rehan and Richards 2010b). In spring, overwintered females emerge from their hibernacula, dispersing to construct new nests in twigs and stems. In summer, mothers remain in the nest to guard against predators and parasites. In autumn, offspring emerge and are primarily fed by the mother or female siblings prior to overwintering in the natal nest (Rehan et al. 2014). Male and female offspring overwinter together as adults and all individuals disperse and mate in the spring.

Ceratina calcarata is of special interest to the study of social evolution because in addition to prolonged mother–offspring interactions and overlapping generations this species also exhibits reproductive division of labor (Rehan et al. 2014). Mothers have complete control over the size and sex of their offspring and produce a special class of dwarf eldest daughters as well as regular daughters (Rehan and Richards 2010b). Regular daughters emerge in the natal nest, do not forage, and wait until spring to disperse and establish their own nests solitarily. Conversely, dwarf eldest daughters are first to emerge in autumn, forage and feed siblings prior to overwintering, and do not survive the winter (Rehan and Richards 2010a). The sibling care without reproduction

exhibited in *C. calcarata* is comparable in some ways with worker behavior exhibited in eusocial species such as the honey bee. However, *C. calcarata* exhibits alloparental care but not cooperative brood care in the strict sense (Michener 1974). The distinction here is that dwarf eldest daughters feed adult siblings but do not contribute to the next generation by provisioning immature offspring. Thus, *C. calcarata* exhibits two of the three hallmarks of eusociality: Reproductive division of labor and overlapping generations, yet this species does not exhibit cooperative brood care and therefore remains a bee on the brink of eusociality (Rehan et al. 2014). This system provides an excellent model for studying the genetic and epigenetic factors associated with maternal care, sibling care, and transitional stages of social behavior.

Here we present the genome of the small carpenter bee, *C. calcarata*. We also compare these data with the previously published bee genomes to identify genomic regions under selection at the brink of sociality. Finally, we provide the DNA methylome of *C. calcarata*. The hypothesis that DNA methylation in bee genomes may be correlated with social complexity has garnered recent support from an indirect measure of DNA methylation (Kapheim et al. 2015). However, empirical assessment of DNA methylation has remained limited among bees to *A. mellifera*. Our study provides the first direct comparison of DNA methylation in *A. mellifera* with a bee with markedly less complex social interactions.

Methods and Materials

Sample Preparation and Sequencing

DNA was isolated from the whole body of two haploid male *C. calcarata* collected in Durham, NH, in the winter of 2013. DNA extractions were performed using a Qiagen Genomic-tip 20/G kit (Valencia, CA) and standard protocol. Genomic DNA was used to generate nonamplified 150-bp DNA libraries for Illumina sequencing. In order to improve genome assembly, genomic DNA was extracted from the whole body of ten female *C. calcarata* as above, pooled, and used to construct 3–4 kb and 5–8 kb mate pair libraries. Four paired-end libraries with insert sizes ranging from 150 bp to 8 kb were constructed and sequenced on Illumina HiSeq 2500. Before filtering this produced 45 Gb of raw sequence data (150.5 million read pairs, 2×150 bp each). Low quality reads, reads with a high proportion of Ns, or poly-A structures were filtered and paired-end reads whose mates overlapped were merged prior to assembly. PRINSEQ (Schmeider and Edwards 2011) was used to filter out reads containing a high number of Ns or poly-A 5' or 3' runs (≥ 15 poly-A nts, $\geq 50\%$ Ns). FLASH (Magoč and Salzberg 2011) was used to merge overlapping reads of the same pair (due to fragment size $<$ read length $\times 2$ on many reads). All genomic data have been submitted to public repositories and can be found using the NCBI BioProject number PRJNA299559.

Genome Assembly

Reads were checked with FastQC (Patel and Jain 2012) for quality metrics (both before and after correction/adaptor removal). Trimmomatic (Bolger et al. 2014) was then used to remove adaptor contamination using Illumina nextera-related adaptor sequences. FastUniq was then used to remove duplicates from each library (Xu et al. 2012). SOAPdenovo's error correction module (Luo et al. 2012) was then used to correct overt errors in the reads using a kmer size of 23. Postfiltering, 32 Gb of sequence remained (107,384,903 read pairs; [supplementary table S1, Supplementary Material](#) online).

Contigs and initial scaffolds were constructed using SOAPdenovo (Luo et al. 2012) with a kmer size of 57. We then used REAPR to break putatively misassembled scaffolds at weak points (Hunt, Kikuchi, et al. 2013). SSPACE was then used to rescaffold broken scaffolds/contigs (Boetzer et al. 2011). GapFiller (Boetzer and Pirovano 2012) was then used to fill assembly gaps left after scaffolding. Finally, L_RNA_Scaffolder (Xue et al. 2013) was used along with independently assembled transcripts (Trinity; Haas et al. 2013) to further scaffold contigs and scaffolds that were broken in the middle of gene annotations, followed by a final round of gap filling with GapFiller (Boetzer and Pirovano 2012). VecScreen (NCBI) was used to scan the assembly for contamination from sequence adapters and vectors, and BLASTN was used to search the nucleotide database for foreign DNA (*Wolbachia*, *Escherichia coli*, bacterial). Scaffolds showing contamination were then removed or trimmed accordingly.

Genome Annotation

Gene annotation was performed with the MAKER2 pipeline (Holt and Yandell, 2011). Briefly, the gene prediction algorithms of SNAP (Korf 2004), Augustus (Stanke and Waack 2003), and GeneMark (Lukashin and Borodovsky 1998) were iteratively trained by *C. calcarata* RNA-seq data and homologous proteins in other bee taxa (see Methods in [supplementary material S1, Supplementary Material](#) online). Annotations with a MAKER AED score above 0.75 or no significant InterProScan domains were flagged as weakly supported. Our filtered gene set was produced requiring a MAKER2 AED of less than 0.75. AED corresponds to the level of external support (transcriptome coverage or homology to other bee gene sets) a given MAKER2 gene prediction received, with our cutoff (< 0.75) identified as reasonable in a previous publication (Holt and Yandell 2011).

RepeatMasker (Smit et al. 2015) was used with default parameters to identify repetitive elements based on the Repbase transposable element library ("species = all") and a custom species-specific repeat library generated by RepeatModeler (Jurka et al. 2005). Functional annotation of genes was performed with Blast2GO (Conesa et al. 2005), using default settings for GO assignment based on querying of the NR database (May 1, 2015) using BLASTP (E -value <

10^{-5}), along with all significant InterProScan hits. KEGG orthology was assigned online using the KAAS server (Moriya et al. 2007). Genome assembly and annotation completeness were assessed with Benchmarking Universal Single-Copy Orthologs (BUSCO; Simão et al. 2015). Specifically, the OrthoDB BUSCO pipeline (v.1.2; October 2013) was employed to test for the presence and completeness of orthologs to *A. mellifera* sequences from arthropod-level BUSCOs in *C. calcarata*.

Gene Family Expansion and Contraction

To identify orthologous groups for analysis of gene family expansion and contraction, we used OrthoMCL (Li et al. 2003) to assign proteins from our study species, *C. calcarata*, and ten other bee species: *Apis florea*, *A. mellifera*, *Bombus impatiens*, *Bombus terrestris*, *Dufourea novaeangliae*, *Eufriesea mexicana*, *Habropoda laboriosa*, *Lasioglossum albipes*, *Melipona quadrifasciata*, and *M. rotundata* into gene families. Briefly, the longest transcript isoform only for each gene was used and we only considered proteins larger than 50 amino acids. An all-versus-all sequence search was performed using BLASTP with default parameters and cut-off E -values of 10^{-7} . The output was uploaded to the OrthoMCL website and run using the default parameters (cutoff of 10^{-5} and 50% match).

To identify gene families that have undergone significant size changes, we used the program COUNT (Csűros 2010). COUNT uses phylogenetic birth-and-death models in probabilistic inference to calculate the probability of gene family evolution across the phylogeny. As an input phylogenetic tree (fig. 1), we used the following: (Hlab:91,(Ccal:65,(Mqua:68,(Bimp:13,Bter:13):55):10,(Emex:62,(Amel:19,Aflo:19):43):16):14):13):15,Mrot:106):9,(Dnov:85,Lalb:85):30) based on Kapheim et al. (2015).

The rate optimization was computed by COUNT using the default parameters and posterior probabilities for each gene family were given. These gene families were then described using Pfam (Finn et al. 2014) domains and keywords and subsequently sorted into gene ontology (GO) terms (Ashburner et al. 2000). By comparing the GO/IRP/KEGG enrichment results for *C. calcarata* with the previously published bee genomes, we identified genes specific to *C. calcarata*. To identify genes lost or absent from the *C. calcarata* genome, we identified genes found in the other ten bee species, but not in *C. calcarata* (see Methods in [supplementary material S1, Supplementary Material](#) online).

Molecular Evolution

For alignments used in PAML analyses, we identified sequences with homology to putative 1-to-1 ortholog groups previously established among ten bee genomes by OrthoDB (Kriventseva et al. 2015). We first performed reciprocal BLASTP searches between the *C. calcarata* OGS and published

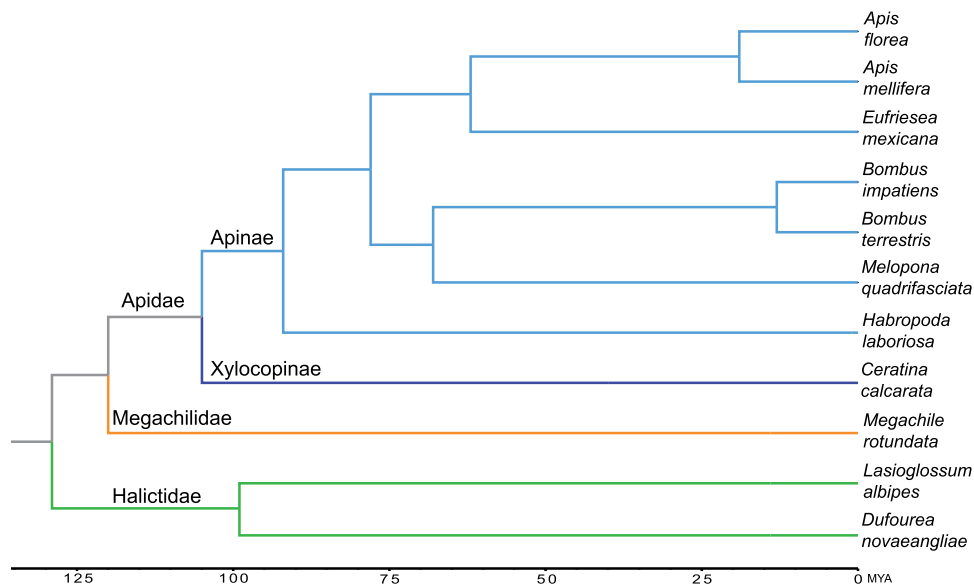


Fig. 1.—Phylogenetic placement of *Ceratina calcarata*. Divergence dates taken from Kapheim et al. (2015) and Rehan and Schwarz (2015). *Ceratina* and *Apis* diverged approximately 105 Ma. Tree topology was confirmed by an analysis of orthologous 4-fold degenerate sites (supplementary material, Supplementary Material online).

gene sets of each of the ten bee species. *Ceratina calcarata* genes were then assigned to a given ortholog group if the *C. calcarata* gene was a reciprocal best hit to the respective ortholog in at least 60% of the ten bees.

For multiple sequence alignment, PRANK was used to align amino acid sequences then back-translate to coding sequences (CDS) (with the option “+F”) from within the Guidance wrapper (v.2.0; Löytynoja and Goldman 2005; Sela et al. 2015). Initial gene trees produced by guidance were then improved using RAxML (Stamatakis 2014) with the GTRGAMMA model. Gblocks was then used to extract higher confidence alignment blocks, while maintaining reading frame, for downstream analysis (Talavera and Castresana 2007).

PAML (v.4.7) was then used to estimate synonymous and nonsynonymous substitution rates (dS and dN) and ω (dN/dS) with free ratios for each branch (Yang 2007). Values corresponding to the *C. calcarata* branch were used in analyses. To test for further evidence of positive selection, we used the branch-site test of positive selection (branch-site model A, test 2; Zhang et al. 2005), which uses likelihood ratio tests (LRTs) to detect positive selection on a small number of sites along a specific lineage. *Ceratina calcarata* was set as the foreground branch, with two dN/dS ratios for branches (model = 2). For the alternative model, ω_2 was estimated from the data (fix_omega = 0, omega = 1, NSsites = 2), while the null model fixes ω_2 at 1 (fix_omega = 1, omega = 1, NSsites = 2) as described in PAML documentation and sample files (Yang 2007). The log-likelihoods for the null and alternative models were used to calculate an LRT statistic, which was then compared against the $\chi^2_{df=1}$ distribution. P values

were then corrected for multiple testing using the method of Benjamini and Hochberg (1995), with a corrected P value threshold of 0.05 being used to assign significance.

Species Phylogeny

Four-fold degenerate sites were extracted from codon sequence alignments after treating with Gblocks. The `msa_view` utility (from the PhastCons program suite; Siepel and Haussler 2004) was used to extract and concatenate 4-fold degenerate sites from the alignments. PhyloFit (REV model) was then used to fit tree models to subwindows ($l = 20,000$; step = 10,000) of the concatenated 4-fold degenerate sites, which were then averaged using phyloBoot (supplementary material S1, Supplementary Material online).

Gene Expression

RNA-seq reads (Rehan et al. 2014; Durant et al. 2016) were aligned to the reference genome using Tophat2 with default options (bar limiting maximum intron length to -l 25000). Cuffnorm was used to produce normalized (geometric) FPKM values for genes (Trapnell et al. 2012). Raw data are available on the NCBI Sequence Read Archive with accession number SRX541305. Six classes of females were assayed and are represented in these studies: Spring mothers, summer mothers, autumn mothers, autumn workers, autumn gynes, and overwintering gynes. Each of the six RNA-seq libraries comprised whole brain tissue samples from 20-pooled females. Reads were filtered for quality (threshold greater than or equal to 20) with a length threshold of 50 bases. After adapter removal, approximately 20% of the reads

were removed from the libraries after quality filtering (98 million raw Illumina reads; 79 million reads after preprocessing).

DNA Methylation

In order to assess DNA methylation at a single-nucleotide resolution, genomic DNA was extracted from ten whole body females using a standard phenol–chloroform protocol and then pooled into one sample for bisulfite conversion and sequencing. Unmethylated enterobacteria phage lambda DNA (GenBank accession: J02459.1) was added to *C. calcarata* genomic DNA as a control for bisulfite conversion efficiency. Bisulfite conversion and sequencing on the Illumina HiSeq platform was performed by Beijing Genomics Institute (Shenzhen, China). We obtained 37.9× mean coverage of genomic CpG sites. We used the program Bismark (Krueger and Andrews 2011) to align raw sequencing reads to the *C. calcarata* reference genome. *Apis mellifera* bisulfite-sequencing data from a worker whole body (Zemach et al. 2010) and from queen and worker brains (Lyko et al. 2010) were previously published, but raw reads were mapped and processed in the same manner as *C. calcarata*. For comparison with *C. calcarata* DNA methylation, we first examined data from an *A. mellifera* worker whole body library (Zemach et al. 2010) to achieve comparability in tissues assayed. Second, we examined data produced by merging worker whole body, worker brain, and queen brain libraries (Lyko et al. 2010; Zemach et al. 2010) to achieve comparability in read depth between species. Results from each comparison were qualitatively similar.

Significantly methylated CpG sites were assessed using a binomial test, which incorporated deamination rate (from our unmethylated control) as the probability of success, and assigned a significance value to each CpG site related to the number of unconverted reads (putatively methylated Cs) as they compare with the expected number from control (Lyko et al. 2010). Resulting *P* values were then adjusted for multiple testing using the method of Benjamini and Hochberg (1995). Only sites with false discovery rate (FDR) corrected binomial *P* values <0.05 and >3 reads sequence coverage were considered “methylated.”

Fractional methylation values were calculated for each CpG site as mCG/CG, where mCG is the number of reads with a methylated cytosine at a CpG site and CG is the total number of reads mapped to the site. Fractional methylation was calculated for specific genomic features (e.g., exons and introns) as the mean of all CpG fractional methylation values within that feature.

Results and Discussion

Genome and Annotation

The *C. calcarata* genome size is estimated at 364 Mb after using kmer-based error correction in QUAKE (Kelley et al. 2010). The final assembly has an N50 of 74 kb and a total

Table 1

Genome Assembly Statistics

Contigs (<i>n</i>)	46,222
Largest contig (bp)	52,361
Scaffolds > 1 kb (<i>n</i>)	3,577
N50 scaffolds (bp)	73,643
Scaffolds > N50 (<i>n</i>)	83
Largest scaffold (bp)	3,675,446
Predicted genes	Filtered: 11,310; unfiltered: 20,693
Ultraconserved core eukaryotic genes (complete/partial, %)	99.6/100
Arthropod BUSCOs in genome assembly	99.3% (97.0% single copy, 2.3% multicopy)
BUSCOs in predicted genes	96.6% (94.2% single copy, 2.4% multicopy; same results for filtered and unfiltered gene predictions)

length of 261.9 Mb (table 1). Completeness of the assembly was first assessed using the CEGMA pipeline (Parra et al. 2007). Of the 248 core eukaryotic genes (CEGs), 247 CEGs were completely assembled in the *C. calcarata* genome. A combination of RNA-sequencing, de novo, and homology-based predictions generated the unfiltered gene set including 20,693 predicted genes. A more conservative “filtered” official gene set was produced including 11,310 predicted genes. The closest relative to *C. calcarata* with a genome sequence is the honey bee, *A. mellifera*, and our results of genome size and gene count are comparable with that of the other ten sequenced bee species (supplementary table S2, Supplementary Material online).

The 11,310 predicted genes comprise 7,888 gene families (supplementary table S3, Supplementary Material online). A total of 5,307 gene families are shared among all bee species and there are 82 predicted orthogroups unique to *C. calcarata* (fig. 2). The orthogroup and Pfam protein family results for *C. calcarata*-specific genes are listed in supplementary table S4, Supplementary Material online. Among these 82 unique orthogroups many are associated with zinc finger protein binding and have overrepresentation of protein domain-specific binding (OG5_158738).

Gene Family Expansion

A total of 306 gene families had been significantly expanded in *C. calcarata* (supplementary table S5, Supplementary Material online). Two notable gene families appear to be expanded in the *C. calcarata* lineage: Zinc finger and odorant receptors proteins. Zinc finger proteins have been implicated in gene regulation in female reproduction and are reportedly expanded in the eusocial termite (*Zootermopsis nevadensis*; Terrapon et al. 2014). We found an expansion of zinc finger proteins in *C. calcarata* (OG5_126627) with 50 copies in their

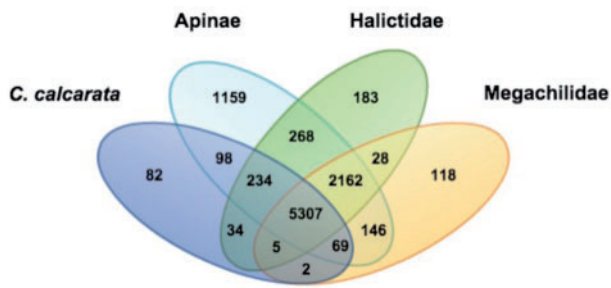


Fig. 2.—Gene family overlap among four bee lineages. The 11 species were sorted into 4 groups based on taxonomic classification: (1) *Ceratina calcarata* (subfamily Xylocopinae) versus (2) subfamily Apinae (*Apis florum*, *Apis mellifera*, *Bombus impatiens*, *Bombus terrestris*, *Eufriesea mexicana*, *Habropoda laboriosa*, and *Melipona quadrifasciata*) versus (3) family Halictidae (*Lasioglossum albipes* and *Dufourea novaeangliae*) versus (4) family Megachilidae (*Megachile rotundata*). Numbers indicate the gene families in each comparison. A total of 5,307 gene families are shared among all 11 bee species.

genome compared with 16 in *L. albipes* and only 2 copies in *A. mellifera*. Zinc finger proteins are known transcription factors and recent analyses of ten bee genomes revealed increasing number of zinc finger binding sites in eusocial versus solitary species (Kapheim et al. 2015). Detailed comparative analyses within and between lineages of bees will be required to reveal the role and regulation of transcription factors across the social spectrum from solitary to eusocial (Rehan and Toth 2015).

Odorant receptors are a large family of transmembrane chemosensory proteins that detect odor molecules and initiate a signal transduction cascade to neurons in the brain (Sanchez-Gracia et al. 2009). The *C. calcarata* genome contains 16 members of this gene family, in contrast to 4 in *M. rotundata* and 2 in *L. albipes* (OG5_168510). Recent analysis of ten bee genomes (Kapheim et al. 2015) found odorant receptors to be expanded in complex eusocial lineages (Apinae) compared with solitary (Megachilidae and Halictidae) species. However, we found expansion of odorant receptors in a subsocial apid bee, consistent with gene family expansion common to the Apidae and likely preceding the evolution of eusociality.

Gene Family Contraction

A total of 281 gene families have been significantly contracted in *C. calcarata* (supplementary table S6, Supplementary Material online). One underrepresented protein domain of interest is the Perilipin gene family. *Perilipin* is also known as lipid droplet-associated protein or *PLIN*, a protein that, in *Drosophila*, is encoded by the *PLIN* gene. The perilipins are a family of proteins that associate with the surface of lipid droplets. Phosphorylation of *perilipin* is essential for the mobilization of fats in adipose tissue (Beller et al. 2010). Past studies

have implicated lipid storage and metabolism as key to overwintering physiology in this and other Hymenopteran species (Strassmann et al. 1984; Toth et al. 2009; Wasielewski et al. 2014; Durant et al. 2016). Moreover, previous studies have linked honey bee immune status to lipid store gene expression profiling (Mao et al. 2013; Salmela et al. 2015). Future studies on lipid metabolism and the role of fat bodies on bee health are an important line of research and will benefit from additional genome-wide comparisons. *Ceratina calcarata* is an emerging model species to understand the lipid metabolism and overwintering diapause and nutritional regulation of caste determination (Rehan et al. 2014; Durant et al. 2016).

DNA Methylation

Epigenetic marks can be influenced by environmental variation, and are capable of influencing postembryonic development (Jaenisch and Bird 2003). Such a role for DNA methylation has been highlighted in the honey bee *A. mellifera*, where larval knockdown of *DNA methyltransferase 3* (*DNMT3*), an enzyme essential to de novo DNA methylation (Klose and Bird 2006), resulted in a developmental shift from the phenotype of workers to the phenotype of queens (Kucharski et al. 2008). Reversible changes in DNA methylation are also associated with changes in behavior linked to the division of labor among workers in *A. mellifera* (Herb et al. 2012). These findings, along with the analysis of mutational signatures of DNA methylation in ten bee genomes, have given rise to the hypothesis that increased gene regulatory capacity may contribute to the evolution of social complexity in bees (Kapheim et al. 2015). One potential mechanism implicating DNA methylation in such regulatory complexity is its role in the regulation of alternative mRNA splicing (Shukla et al. 2011; Li-Byarlay et al. 2013).

Here, we present results of whole genome bisulfite sequencing (WGBS) for *C. calcarata* (supplementary table S7, Supplementary Material online). Like *A. mellifera* (Wang et al. 2006), the genome of *C. calcarata* contains a complete enzymatic toolkit for DNA methylation (fig. 3a; Klose and Bird 2006), comprised in *C. calcarata* of two copies of *DNMT1* (Ccalc.v2_000349 and Ccalc.v2_004812) and one copy of *DNMT3* (Ccalc.v2_012237); *DNMT2* is not known to methylate DNA. Moreover, genes targeted by DNA methylation in *C. calcarata* are enriched for GO biological processes associated with core cellular functions, including translation, and have detectable expression in our RNA-seq data (fractional CDS DNA methylation vs. gene expression level [FPKM], Spearman's rho = 0.58, $P < 0.0001$; same correlation when removing genes with mCG/CG < 0.01 and FPKM < 1, Spearman's rho = 0.14, $P < 0.0001$). These findings are consistent with those from diverse invertebrate taxa, including *A. mellifera* (Sarda et al. 2012; supplementary table S8, Supplementary Material online). As in other investigated holometabolous insects (Hunt, Glastad, et al. 2013), DNA

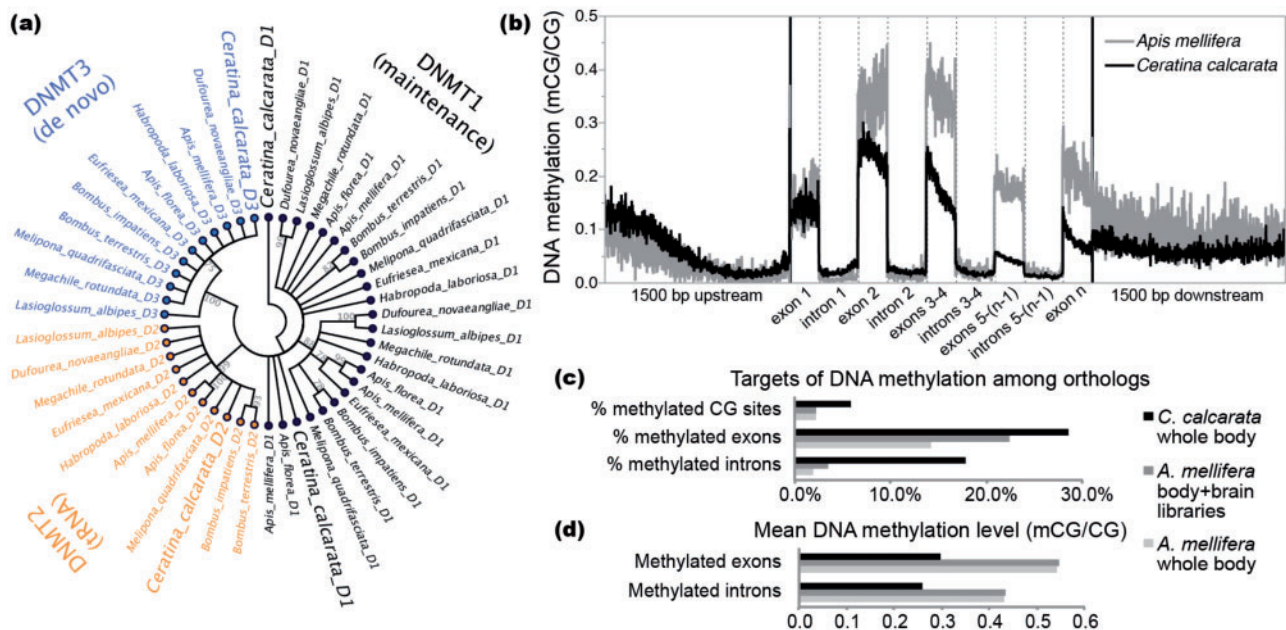


Fig. 3.—DNA methylation in *Ceratina calcarata*. (a) Unrooted neighbor-joining cladogram showing relationships among the DNA methyltransferase proteins DNMT1 (D1), DNMT2 (D2), and DNMT3 (D3) from *C. calcarata* and 10 other bee species, demonstrating the presence of a fully functional DNA methylation toolkit in *C. calcarata*. Polytomies are plotted for nodes with less than 75% support among 1,000 bootstrap replicates. (b) Spatial profiles of mean DNA methylation levels in genes with significant methylation from *C. calcarata* and the honey bee *Apis mellifera*. Exons are divided into 150 proportional bins and introns are divided into 200 proportional bins. DNA methylation is preferentially targeted to exons 1–3, and levels decay toward the 3' end of genes in each species (Wilcoxon rank-sum tests between exons 1–3 and 4–*n*, $P < 0.0001$). (c) Percent of CG dinucleotides, exons, and introns targeted by DNA methylation in 5,698 shared orthologs suggests more widespread targeting of DNA methylation in *C. calcarata* versus *A. mellifera*, whereas (d) mean methylation levels suggest that *C. calcarata* exhibits lower levels of DNA methylation, where targeted, compared with *A. mellifera*.

methylation is preferentially targeted to exons positioned proximal to the translation start site (TSS) in *C. calcarata* (fig. 3b; Wilcoxon rank-sum test between exons 1–3 and 4–*n*, $P < 0.0001$). We previously put forth the hypothesis that TSS-proximal DNA methylation plays a role in differentiating states of transcriptional initiation and elongation (Glastad et al. 2015).

To gain insight into the potential relationship between social complexity and DNA methylation, we compared *C. calcarata* DNA methylation with that observed in orthologs of the advanced eusocial honey bee *A. mellifera* using WGBS data generated from female adults of each species (Lyko et al. 2010; Zemach et al. 2010; [supplementary table S9](#), [Supplementary Material](#) online). We observed that a greater proportion of CpG dinucleotides were methylated in *C. calcarata*, which is generally considered subsocial, compared with *A. mellifera*, which exhibits complex eusociality (fig. 3c). Moreover, among 3,841 ortholog pairs with significant DNA methylation detected in CDS of at least one species, 3,147 (81.9%) were targeted by DNA methylation in both species, 456 (11.9%) were targeted by DNA methylation in *C. calcarata* only, and 238 (6.2%) were targeted by DNA methylation in *A. mellifera* only. Thus, our results stand in contrast with the prediction that DNA methylation targets are more widespread

in species with complex social organization in bees (Kapheim et al. 2015). However, we did observe that methylated cytosines in *A. mellifera* exhibited higher DNA methylation levels (i.e., were markedly more frequently methylated across sequenced cells) than in *C. calcarata* (fig. 3b and d). Overall, our results highlight the multidimensionality of DNA methylation—the number of targets and the frequency with which cells are methylated each stand to influence gene regulation. Thus, detailed investigation of DNA methylation patterns in specific tissues from numerous taxa will be required to resolve whether generalities exist in the relationship between DNA methylation and the evolution of sociality.

Molecular Evolution

Ceratina calcarata is the first bee of the subfamily Xylocopinae and the first subsocial bee species with genome characterization. Thus, this species represents an important point of comparison for the molecular evolution of sociality (Rehan and Toth 2015). To identify genes showing signatures of positive selection in the *C. calcarata* lineage, we conducted branch-site tests in PAML. We included all currently published bee genomes to perform these analyses. These taxa include ten species (*A. mellifera*, *A. florea*, *E. mexicana*, *B. terrestris*,

Table 2

GO Enrichment for Genes Subject to Positive Selection according to a Branch-Site Test (FDR $P < 0.05$ vs. FDR $P > 0.05$). Gene ontology categories include molecular function (F) and biological processes (P)

Term	GO-ID	Category	No. of Genes	Fold Enrichment	P Value	FDR
Nuclease activity	GO:0004518	F	19	2.24	3.49E−03	1.00E+00
Lipid metabolic process	GO:0006629	P	35	1.45	3.32E−02	1.00E+00
DNA metabolic process	GO:0006259	P	34	1.43	4.11E−02	1.00E+00
Secondary metabolic process	GO:0019748	P	3	5.66	4.73E−02	1.00E+00

B. impatiens, *Mel. quadrifasciata*, *H. laboriosa*, *M. rotundata*, *L. albipes*, *D. novaeangliae*). We used the “branch-site A” model in PAML to search for sites showing signatures of positive selection in the *C. calcarata* lineage using a LRT to calculate significance. Following correction for multiple testing, 877 genes showed signatures of positive selection in the *C. calcarata* lineage (FDR $P < 0.05$; [supplementary table S10](#), [Supplementary Material](#) online). Among the noteworthy genes with signatures of positive selection were two genes associated with regulation of TOR functioning (Ccalc.v2_007251 and Ccalc.v2_008102; [supplementary table S11](#), [Supplementary Material](#) online). TOR genes are highly conserved among bees (Kapheim et al. 2015) and have been shown to play a functionally important role in honey bee caste development (Patel et al. 2007; Mutti et al. 2011).

In general, genes associated with lipid transport, protein binding, and ribonuclease activity appear to have undergone positive selection in *C. calcarata* relative to the other species examined (table 2). Genes associated with DNA recombination are also evolving more quickly in *C. calcarata* (Ccalc.v2_012690, Ccalc.v2_001846, Ccalc.v2_013055, Ccalc.v2_004327; [supplementary table S11](#), [Supplementary Material](#) online). High rates of DNA recombination are known in advanced eusocial insects (Wilfert et al. 2007; Sirviö et al. 2011; Kapheim et al. 2015) and could be linked to accelerated evolution of caste related traits (Kent et al. 2012). It is possible that the accelerated evolution of DNA recombination genes may reflect this group’s transitional form of sociality including the presence of an incipient worker caste (Rehan et al. 2014).

Conclusions

The maternal care and sibling care expressed in an incipiently social bee make *C. calcarata* ideal for developmental and sociodemographic studies to characterize the genomic effects of social experience on behavioral plasticity. Our results suggest that mechanisms associated with DNA methylation and nutrition are candidate targets of evolutionary change during the origin of more complex social forms. The addition of *C. calcarata* to the published bee genomes provides the first subsocial bee species to ask questions on the evolutionary origins of social behavior. Moreover, the characterization of

a fully functional *C. calcarata* methylome allows for new lines of research into the epigenetic regulation of maternal care, sibling care, and worker behavior.

Supplementary Material

Supplementary tables S1–S11 and [supplementary material S1](#) are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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