A-to-I RNA editing in honeybees shows signals of adaptation and convergent evolution

Yuange Duan, Shengqian Dou, Hagit T. Porath, Jiaxing Huang, Eli Eisenberg, Jian Lu
elieis@post.tau.ac.il (E.E.)
luj@pku.edu.cn (J.L.)

Highlights
Nonsynonymous editing sites in honeybees were under positive selection
Differential editing may contribute to the phenotypic diversity between sub-castes
Target genes acquire editing in different clades, suggesting convergent evolution

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A-to-I RNA editing in honeybees shows signals of adaptation and convergent evolution

Yuange Duan,1,2 Shengqian Dou,1 Hagit T. Porath,3 Jiaxing Huang,4 Eli Eisenberg,5,* and Jian Lu1,6,*

Summary
Social insects exhibit extensive phenotypic diversities among the genetically similar individuals, suggesting a role for the epigenetic regulations beyond the genome level. The ADAR-mediated adenosine-to-inosine (A-to-I) RNA editing, an evolutionarily conserved mechanism, facilitates adaptive evolution by expanding proteomic diversities. Here, we characterize the A-to-I RNA editome of honeybees (Apis mellifera), identifying 407 high-confidence A-to-I editing sites. Editing is most abundant in the heads and shows signatures for positive selection. Editing behavior differs between foragers and nurses, suggesting a role for editing in caste differentiation. Although only five sites are conserved between bees and flies, an unexpectedly large number of genes exhibit editing in both species, albeit at different locations, including the nonsynonymous auto-editing of Adar. This convergent evolution, where the same target genes independently acquire recoding events in distant diverged clades, together with the signals of adaptation observed in honeybees alone, further supports the notion of recoding being adaptive.

Introduction
Adenosine (A)-to-inosine (A-to-I) RNA editing, catalyzed by enzymes of the ADAR (adenosine deaminase acting on RNA) family (Savva et al., 2012b), is an evolutionarily conserved mechanism that expands RNA diversity at the co-transcriptional or post-transcriptional level in metazoans (Bass 2002; Nishikura 2010; Eisenberg and Levanon 2018). Two catalytically active Adar genes are encoded in most metazoans. However, insects have lost ADAR1 and encode only a single Adar gene (Keegan et al., 2011). Due to the structural similarity between inosine (I) and guanosine (G), I is generally believed to be recognized as G in many cellular processes such as mRNA splicing (Ruerter et al., 1999; Flomen et al., 2004; Jin et al., 2007; LevMaor et al., 2007), microRNA (miRNA) biogenesis or target recognition (Liang and Landweber 2007; Borchert et al., 2009; Alon et al., 2012), and mRNA translation (Basilo et al., 1962; Licht et al., 2019). Therefore, an A-to-I RNA editing usually has a similar effect as an A-to-G DNA substitution. A-to-I editing plays essential roles in many biological and physiological processes (Keegan et al., 2001; Nishikura 2010), and dysregulation of A-to-I editing might be associated with cancer, autoimmune disorders, or other human diseases (Gallo et al., 2017).

During the past two decades, A-to-I RNA editing sites have been systematically characterized in various metazoan species (Ramaswami and Li 2016). The majority of RNA editing sites are located in clusters in non-coding regions of humans (Athanasiadis et al., 2004; Blow et al., 2004; Kim et al., 2004; Levanon et al., 2004; Picardi et al., 2017), monkeys (Chen et al., 2004; Yang et al., 2015), mice (Neeman et al., 2006; Daneczek et al., 2012), worms (Morse and Bass 1999; Zhao et al., 2015; Goldstein et al., 2017), corals (Porath et al., 2017b), and many other species (Porath et al., 2017a). In most species, only a minute fraction of the edits resides within the coding sequence. Notable exceptions are Drosophila (Graveley et al., 2011; Rodriguez et al., 2012; St Laurent et al., 2013; Mazloomian and Meyer 2015; Yu et al., 2016; Buchumenski et al., 2017; Duan et al., 2017; Zhang et al., 2017) and cephalopod species (Alon et al., 2015; Liscovitch-Brauer et al., 2017). Despite the deep conservation of A-to-I editing mechanism, the target landscapes of editing have considerably evolved during metazoan evolution. Only one editing site is known to be conserved across virtually all mammals, Drosophila, and cephalopods (Porath et al., 2019), and it thus seems that RNA editing modulates the diversity of the transcriptomes and proteomes in a lineage-specific manner.
The forces driving the evolution of A-to-I editing across species at the macro-evolutional scale are not well understood. RNA editing was hypothesized to facilitate adaptive evolution by increasing proteomic diversities temporally or spatially, in a manner more flexible than genomic mutations (Gommans et al., 2009; Nishikura 2010, 2016; Klironomos et al., 2013; Rosenthal 2015). However, in most species, the studied ratio of nonsynonymous (N) to synonymous (S) editing sites (N/S) is lower than expected for random sites, suggesting that recoding events may be overall non-adaptive (Xu and Zhang 2014). Here too Drosophila and cephalopods stand out as exceptions, exhibiting high N/S ratios indicating positive selection of recoding. Furthermore, hundreds of recoding sites were shown to be conserved across the Drosophila lineage (Gravely et al., 2011; Rodriguez et al., 2012; St Laurent et al., 2013; Mazloomian and Meyer 2015; Yu et al., 2016; Buchumenski et al., 2017; Duan et al., 2017; Zhang et al., 2017) and thousands in the behaviorally sophisticated cephalopods (Garrett and Rosenthal 2012; Alon et al., 2015; Liscovitch-Brauer et al., 2017), supporting the functional importance of these editing events. However, in only a few examples was the advantageous effect conferred by RNA editing explicitly demonstrated (see for example [Garrett and Rosenthal 2012]), and even it is not yet clear to what extent editing is indeed utilized for proteome diversification.

Social insects, including bees and ants, show extensive phenotypic plasticity. The morphologically and behaviorally differentiated social castes such as queens, workers, and drones have the same set of diploid or haploid genomes. The social insects provide us with model systems to study how phenotypic diversity is regulated (Page et al., 2012; Yan et al., 2014). RNA editing is well suited to contribute to the behavioral variation among genetically similar individuals. Indeed, differential editing was demonstrated for a few sites in leaf-cutting ants and in worker bumblebees (Li et al., 2014; Porath et al., 2019).

Here, we wish to study the contribution of RNA editing to caste differentiation in honeybee (Apis mellifera), an important pollinator and a model for complicated social behaviors in insects (Page et al., 2012). Unlike bumblebees, honeybees have a sharp task specialization, with distinct worker castes. Genetic mapping demonstrated that the phenotypic plasticity in honeybees is associated with complex epistatic and pleiotropic genetic networks that influence reproductive regulation and foraging behaviors (Page et al., 2012). We wish to check whether A-to-I RNA editing may contribute to the proteomic diversity underlying this phenotypic plasticity. We investigate the A-to-I RNA editome in different tissues of four honeybee drone individuals and detect over four hundred A-to-I sites in multiple tissues. Editing is enriched in the head and exhibits signs for positive selection and a particularly high N/S ratio. We show editing is elevated in foragers compared to nurses. Five editing sites are conserved between honeybee, bumblebee, and Drosophila. Interestingly, we find a significantly high number of cases where the same gene is edited in both bees and flies, even if not at the same position. One example is the auto-editing of Adar mRNA in bees and flies, which might play an auto-regulatory role in the two clades. This finding may further support possible convergent evolution and adaptation.

Results
Editome of honeybee

We deep sequenced genomic DNA and RNA from the head, thorax, and abdomen for each of four individual drones (12 RNA-seq samples) (Figure 1A). The transcriptome libraries of drones 1 and 2 were constructed by selecting polyA tailed mRNAs while the libraries of drones 3 and 4 were constructed by using the Ribo-Zero kit to deplete ribosomal RNAs (Transparent methods). The male drone is haploid, simplifying identification of its genomic single nucleotide polymorphisms (SNPs). DNA-Seq reads were mapped to the reference genome to identify 826,890-858,949 SNPs in each of the four drones and 1,303,225 unique sites combined (Transparent methods; Table S1). Median sequencing coverage at the SNP sites was 39, and in 96.6% of the sites, all DNA reads supported the SNP (Figure S1). The identified SNPs were then used to produce a masked genome version where the reference allele is substituted by the individual-specific alleles identified for each drone to facilitate accurate detection of RNA editing sites (Figure 1B; Transparent methods).

For each of the 12 RNA-Seq samples, 9.2-21.5 million reads were uniquely mapped to the reference genome, and similar numbers of reads were mapped to the masked genome sequences (Transparent methods; Table S2). Analyzing pooled data for each of the three tissues separately, we identified (false discovery rate [FDR] = 0.05) 376 (84.3%) A-to-G sites among 446 variations in heads, 106 (66.7%) A-to-G sites among 159 variations in thoraxes, and 137 (67.5%) A-to-G sites among 203 variations in abdomens (Figure S2). Combined, we obtained 407 (80.1%) unique A-to-G sites among 508 variations (Figure 1C). These
407 sites are regarded as A-to-I RNA editing sites (Table S3). The nucleotide context around these putative editing sites is consistent with the known ADAR binding motif (Figure S3). The median DNA coverage (over the 407 A-to-I editing sites) was 25 reads, sufficient to exclude possible SNPs (Figure S4).

To improve mapping accuracy, we discarded variants in repeat regions and required a variation site to be found by mapping to both the reference genome and the masked genome (Transparent methods). To test whether these criteria are too stringent, we checked the variation sites discarded by the two filtering steps. First, we looked at the variation sites located in the repeat regions. Without any filter, we found 19,573 unique variation sites that overlapped with repeat regions, only 3,694 (19%) of which were A-to-G variants. Following a binomial test and multiple testing correction (to remove random sequencing errors), 1,277 variation sites were maintained, only 324 (25%) of them A-to-G variations. Thus, filtering repetitive regions does contribute appreciably to the precision of our detection. Consistently, hyper-editing analysis did not reveal multiple sites in repetitive regions (Transparent methods).

As for the masked genome filter, only 38% (98/256) of the variation sites found by mapping to the masked genome but not to the reference genome are A-to-G mismatches. Following binomial test for sequencing error and multiple testing correction for p values, only 41 variation sites were maintained, none of them was an A-to-G variation. Conversely, only 31% (82/268) variation sites found by mapping to the reference genome but not to the masked genome are A-to-G mismatches. Of these 268 variation sites, 82 were maintained after multiple testing correction, 20 of them were A-to-G variations. Thus, most of the sites supported by only one method of mapping are likely not due to A-to-I RNA editing.

Among the 407 editing sites we identified, 199 sites are located in gene regions, including 111 nonsynonymous (N) sites, 9 synonymous (S) sites, and several sites in untranslated regions (UTRs) and introns (Figure 1D). The other 208 sites are annotated as intergenic. As expected from the different library construction strategies, the fraction of sites in coding regions is higher and the intergenic fraction is lower for sites
observed in drones 1 and 2 compared with drones 3 and 4 (23.2 ± 2.6% in coding sequence (CDS) in drones 1 and 2 versus 14.5 ± 5.1% in CDS in drones 3 and 4; 54.9 ± 2.2% in intergenic regions in drones 1 and 2 versus 70.5 ± 5.4% in intergenic regions in drones 3 and 4). The number of sites for which editing is observed in each sample (per-sample editing level >0) varies considerably (40–278 sites per sample; Table S4). The partial overlap between these sets of sites is mostly due to sites being undetected at a given sample due to low per-sample coverage (Figure S5).

Interestingly, editing seems to be enriched in heads. The number of detected sites (Figure S6), the editing levels (Figure 1E), and the fraction of coding sites (Figure S7) are all higher in heads. Clustering the samples by their editing profile, the three tissue types form distinct clusters (Figure 1F). Consistently, Adar expression is also higher in heads than in other tissues (Figure S8). Looking at specific sites, 78 sites (out of 378 sites with sufficient coverage, see Transparent methods) are differentially edited in the head, compared to non-head tissues. In all of these 78 sites, editing is higher in the head (Table S3. Information of the candidate editing sites identified in this study, Figure 1).

**Signals of adaptation**

The nonsynonymous to synonymous (N/S) ratio for sites in the coding region is 111/9 = 12.3 (Table S4), compared to a ratio of 2.26 observed for random adenosine to guanosine substitutions (see Transparent methods). This strongly suggests that the nonsynonymous editing events in honeybees are overall adaptive. Notably, the N/S is exceptionally high, even compared to the ratios we previously found in the brains of Drosophila (Duan et al., 2017) or in cephalopods (Alon et al., 2015; Liscovitch-Brauer et al., 2017). Looking at each sample separately, at pooled tissue data, or focusing on sites that appear in at least two individuals per tissue all show the N/S ratio to be especially high in heads (Figure 2A and Table S4). Furthermore, the editing level at nonsynonymous sites is significantly higher than synonymous sites editing levels in heads of honeybees (Figure 2B).

A previous study in cephalopods proposed that the coleoids massively edit their RNAs to diversify the transcriptome at the cost of constraining the evolution of genomic sequences (Liscovitch-Brauer et al., 2017). Maintaining beneficial editing requires genomic conservation of the genomic sequence encoding for the dsRNA structures that allow ADAR to bind and deaminate the adenosine. To test this, we first confirmed that the editing sites in honeybees are enriched in hairpin structures (Figure 2C). Then, we demonstrated that SNPs are depleted in the vicinity of nonsynonymous sites (Figure 2D), further supporting the notion of selective advantage for the nonsynonymous editing events that justifies the trade-off between transcriptome diversity and genome evolution.

**Evolutionarily conserved and non-conserved editing sites**

Editing at four nonsynonymous and one synonymous site is conserved between Drosophila and honeybees (Figure 3). Three of these sites are in the Shab transcript (shaker cognate b, two nonsynonymous and one synonymous) and the other two are in qvr (quiver). Editing at these sites is also observed in the bumblebee Bombus terrestris (Porath et al., 2019), and editing levels are high in all of these species (Figure 3). These findings suggest a potential functional role for these widely conserved editing sites. However, the majority of the editing sites are not shared between Drosophila and honeybees. For some sites, the editable adenosine is not conserved in the honeybee genome (Figure S9) and are thus clearly uneditable, while other adenosines are conserved but editing was not detected in RNA-Seq data (Figure S9). Overall, of the 639 Drosophila editing sites in CDS for which an orthologous site was identified in honeybees, 349 were conserved as A (55%), and 85, 127, and 78 (13%, 20%, and 12%) were mutated to C, G, and T, respectively.

Similarly, most sites are not even conserved across the two bee species. The bumblebee study has reported 219 editing sites in CDS, 164 of which are nonsynonymous (Porath et al., 2019). Of these, we found 9 editing sites in coding regions conserved between honeybee and bumblebee (Table S5).

One of these conserved sites is a Ser > Gly site within the tipE (temperature-induced paralytic E; GB47375) transcript, editing levels of which correlate with task performance in bumblebees (nurses versus foragers) (Porath et al., 2019). Editing levels at this site in honeybee heads are ~0.7, similar to those observed in bumblebee brains. In the two other tissues, this gene is lowly expressed and poorly edited. The orthologous site in Drosophila genomes encodes the post-edit Gly codon GGC (Figure S10). Thus, tipE recoding seems to be bee specific, possibly related to social behavior and task performance.
Sex-dependent and caste-dependent editing

We further looked for potential sex-dependence and caste-dependent editing, comparing our honeybee drone head data with previously published brain RNA-seq data studying two sub-castes of female workers, foragers and reverted nurses (Herb et al., 2012) (Table S6). First, we compared the pooled editing levels between drones and workers for each well-covered (>10 reads in each pool) editing site separately. Twenty six sites exhibit differential editing (Wilcoxon rank-sum test; FDR = 0.05) between workers (females) and drones (males), including 4 recoding sites, 2 sites in 3′UTRs, 3 intronic sites, and 17 intergenic sites (Figure 4A and Table S7). In most cases, editing in drones is higher, but this may result from looking at sites identified in drone data to begin with. The difference in *Adar* expression was not statistically significant (Figure S11).

Comparing the pooled editing levels between the two sub-castes (foragers and nurses), one finds 230 out of 407 sites with lower editing levels in nurses, compared to only 91 exhibiting higher levels in nurses (equal level, mostly zero, was observed in 84 sites), suggesting a globally higher editing activity (at the detected sites) in foragers (P = 1.3 × 10\(^{-14}\), proportion test). The surcharge of forager higher sites is maintained for various coverage and editing difference cutoffs (Figure S12). Furthermore, *Adar* expression (Figure 4B) and the editing index (Figure 4C) are both higher in foragers, and the editing index correlates with *Adar* expression. However, reliable detection of specific differentially edited sites with the available sample size (6 foragers
and 6 nurses) is challenging, and not even a single site was identified with FDR = 0.05 (Wilcoxon rank-sum test). We have thus relaxed the statistical test, allowing for FDR = 0.3, and found 61 candidate sites, most of which are differentially edited (Figure 4D and Table S8). These include 10 nonsynonymous, 16 3' UTR, and 35 intergenic sites. Consistently, 55 of these sites exhibit higher levels of editing in foragers. Finally, PCA analysis of the editing profile across sites results in distinct clusters of drones, nurses, and foragers, a classification that is not achieved by the expression profile (Figures 4E and 4F). Taken together, these results show a global increase in editing in foragers compared to nurses.

Convergent adaptation of A-to-I editing?

One of the recoding sites conserved between the two bee species resides within Adar transcript (Table S5 and Figure 5A). In bumblebee, the recoding level of the conserved site, I482M (ATA to ATG), positively

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A

**Shab (shaker cognate b)**

|                | Editing level |
|----------------|---------------|
|                | Site1 | Site2 | Site3 |
| D. mel         | 0.85  | 0.77  | 1     |
| D. sim         | 0.88  | 0.67  | 0.96  |
| D. sec         | 0.93  | 0.63  | 1     |
| D. ere         | 0.98  | 0.59  | 0.97  |
| D. pse         |       |       |       |
| D. wil         | 0.92  | 0.45  | 0.96  |
| B. ter         | 0.93  | 0.97  | 0.98  |
| A. mel         | 0.31  | 0.79  | 0.81  |
| A. ech         | 0     | 0     | 0     |

B

**qvr (quiver)**

|                | Editing level |
|----------------|---------------|
|                | Site1 | Site2 |
| D. mel         | 0.88  | 0.61  |
| D. sim         | 0.96  | 0.60  |
| D. sec         | 0.99  | 0.84  |
| D. ere         | 0.98  | 0.80  |
| D. pse         | 0.96  | 0.86  |
| D. wil         | 0.98  | 0.79  |
| B. ter         | 0.50  | 0.34  |
| A. mel         | 0.87  | 0.67  |
| A. ech         | 0     | 0     |

Figure 3. Editing sites conserved between bees and flies

(A) One synonymous and two nonsynonymous editing sites in Shab transcripts are edited in heads of two bee and four Drosophila species.

(B) Two nonsynonymous editing sites in qvr transcripts are edited in heads of two bee and four Drosophila species. The editing sites are colored red. Editing levels measured in male heads of flies (unpublished), pooled heads of honeybee drones, and brains of bumblebees (Porath et al., 2019) are shown. D. mel, Drosophila melanogaster; D. sim, Drosophila simulans; D. sec, Drosophila sechellia; D. ere, Drosophila erecta; D. pse, Drosophila pseudoobscura; D. wil, Drosophila willistoni; D. moj, Drosophila mojavensis; A. mel, Apis mellifera; B. ter, Bombus terrestris.
correlates with the global editing activity (Porath et al., 2019), suggesting a possible auto-regulation mechanism. Interestingly, Drosophila Adar is also auto-edited at a different position, where a Ser (AGT) to Gly (GGT) substitution leads to a less active ADAR protein, resulting in a negative feedback loop of editing activity (Palladino et al., 2000; Savva et al., 2012a). Intriguingly, fly-edited Ser amino acid is conserved in honeybee and bumblebee, but a different codon (TCA, uneditable at the first codon position) is used. Similarly, the bee-edited Ile is conserved in flies, but the edited adenosines (at the third position of ATA) is synonymously mutated to T, and the editable ATA bee codon is substituted by ATT codon in flies, which could not be edited at the third position (Figure 5A). Thus, while auto-editing S430G in flies is abolished in bees due to an uneditable Ser codon, and the auto-editing I482M in bees is abolished in flies due to an uneditable Ile codon. However, the mechanism of auto-editing, possibly used for global ADAR regulation, is shared by the two lineages, albeit at different positions and possibly with different effects on the protein. This might hint at the possibility of convergent evolution of ADAR auto-regulation strategy.

Following this example, we wondered whether there are more genes for which editing is observed in both honeybees and Drosophila, even if the exact location of the editing site is not conserved. Excluding conserved sites, there are 53 genes exhibiting CDS editing in honeybee heads, 101 genes exhibiting CDS editing in bumblebee heads, and 312 genes edited in CDS in male brains of D. melanogaster. Of these, 21 genes exhibit editing in both bee species, 14 are edited in honeybee and Drosophila, 14 are edited in bumblebee and Drosophila, and six genes are found to be edited in all three species (Figure 5B and Table S9). These numbers are significantly higher than expected by random sampling, (p < 1 × 10⁻⁶),
A randomization test), even if one accounts for the expression profile (Transparent methods, Figures 5C and S13)( Table S9). We have repeated the analysis using stricter criteria to characterize sites, considering as edited only sites with an observed editing level >10% (all samples pooled), and a binomial P < 0.05

|    | site1 | site2 | Editing level |
|----|-------|-------|---------------|
| D. mel | TTT   | AGT   | CCT           |
| D. sim | TTT   | AGT   | CCT           |
| D. sec | TTT   | AGT   | CCT           |
| D. ere | TTT   | AGT   | CCT           |
| B. ter | TTT   | TCA   | CCT           |
| A. mel | TTT   | TCA   | CCA           |

Ser > Gly site is highly conserved across Drosophila species. The Drosophila-editable Ser codon AGT is changed to an uneditable Ser codon TCA in bees. On the other hand, the Ile > Met auto-editing site is conserved between honeybee and bumblebee. The bee-editable Ile codon ATA (edited at the third position) appears as an uneditable Ile codon ATT in flies. Editing sites are colored red. The editing levels from male heads of flies (unpublished), pooled heads of honeybee drones, and brains of bumblebee (Porath et al., 2019) are shown. D. mel, Drosophila melanogaster; D. sim, Drosophila simulans; D. sec, Drosophila sechellia; D. ere, Drosophila erecta; A. mel, Apis mellifera; B. ter, Bombus terrestris.

B Venn diagram demonstrating the overlap between orthologous genes with editing sites in coding regions. (A) Adar transcripts are auto-edited in Drosophila and bees. The Ser > Gly site is highly conserved across Drosophila species. The Drosophila-editable Ser codon AGT is changed to an uneditable Ser codon TCA in bees. On the other hand, the Ile > Met auto-editing site is conserved between honeybee and bumblebee. The bee-editable Ile codon ATA (edited at the third position) appears as an uneditable Ile codon ATT in flies. Editing sites are colored red. The editing levels from male heads of flies (unpublished), pooled heads of honeybee drones, and brains of bumblebee (Porath et al., 2019) are shown. D. mel, Drosophila melanogaster; D. sim, Drosophila simulans; D. sec, Drosophila sechellia; D. ere, Drosophila erecta; A. mel, Apis mellifera; B. ter, Bombus terrestris.

C The observed number of orthologous genes with editing in coding regions, compared to the expected distribution. Genes with RPKM >1 in both species were chosen to calculate the expected numbers (results are essentially the same for other expression cutoffs, Figure S13). p values calculated by randomization test.

D The nonsynonymous to synonymous (N/S) ratios are significantly higher than expected, for both shared and species-specific sites (except for species-specific sites in B. terrestris). Here, the shared and species-specific sites refer to the sites in shared genes and species-specific genes as defined in the above Venn diagram in (B). The expected ratio is evaluated by calculating the effect of putative editing on all adenosines in the coding region. p values are obtained from Fisher’s exact test.

Figure 5. Convergent evolution of editing
(A) Adar transcripts are auto-edited in Drosophila and bees. The Ser > Gly site is highly conserved across Drosophila species. The Drosophila-editable Ser codon AGT is changed to an uneditable Ser codon TCA in bees. On the other hand, the Ile > Met auto-editing site is conserved between honeybee and bumblebee. The bee-editable Ile codon ATA (edited at the third position) appears as an uneditable Ile codon ATT in flies. Editing sites are colored red. The editing levels from male heads of flies (unpublished), pooled heads of honeybee drones, and brains of bumblebee (Porath et al., 2019) are shown. D. mel, Drosophila melanogaster; D. sim, Drosophila simulans; D. sec, Drosophila sechellia; D. ere, Drosophila erecta; A. mel, Apis mellifera; B. ter, Bombus terrestris.

(B) Venn diagram demonstrating the overlap between orthologous genes with editing sites in coding regions. (C) The observed number of orthologous genes with editing in coding regions, compared to the expected distribution. Genes with RPKM >1 in both species were chosen to calculate the expected numbers (results are essentially the same for other expression cutoffs, Figure S13). p values calculated by randomization test.

(D) The nonsynonymous to synonymous (N/S) ratios are significantly higher than expected, for both shared and species-specific sites (except for species-specific sites in B. terrestris). Here, the shared and species-specific sites refer to the sites in shared genes and species-specific genes as defined in the above Venn diagram in (B). The expected ratio is evaluated by calculating the effect of putative editing on all adenosines in the coding region. p values are obtained from Fisher’s exact test.
(excluding the null hypothesis of no editing), and considering a site to be not edited if the binomial \( P < 0.05 \) for the null hypothesis that the site is edited at 10%. Reassuringly, the results are robust to this change of definitions (Figures S14 and S15). These results suggest an interesting convergent evolution of A-to-I editing. Recoding sites are rarely conserved across clades, but the same genes in different lineages tend to acquire a recoding event, apparently independently, indicating a functional importance for recoding of this gene.

**Discussion**

Despite over fifteen years of developments, reliable detection of CDS editing remains a challenge. Systematic searches often lead to high false-positive rates (reflected by the fraction of non-AG mismatches observed), especially for mammalian transcriptomes where the scope of recoding is rather low. Important exceptions are *Drosophila* and cephalopods, where the recoding signal is much more pronounced and easier to detect. Applying a novel strict alignment approach, we were able to achieve here a high accuracy in CDS editing detection (84.3% in heads) despite the overall modest scope of recoding sites. Another similarity between honeybees *Drosophila* and cephalopods is related to the important question regarding the extent to which recoding is adaptive. This is often estimated by the N/S ratio, where *Drosophila* and cephalopods exhibit a pattern markedly different from the one seen in other species studied (including human). In most species, the N/S ratio observed is lower or similar to the one expected under neutrality, and it is significantly higher in *Drosophila* and cephalopods. Again, we find here that honeybees show a strikingly high N/S ratio, similar to the above two clades. We suggest that the two last points are interconnected. The apparently low N/S ratio observed in many species (human included) may reflect the low accuracy of the lists of CDS editing sites and not the (lack of) adaptive potential for recoding in these species.

The importance of post-transcriptional and post-translational mechanisms in generating the proteomic complexity of higher organisms has been emphasized in recent decades. These epigenetic mechanisms allow for diversification of the proteome and functional heterogeneity across tissues, developmental stages, brain regions, or even among individual cells.

Recoding by A-to-I RNA editing is an epigenetic mechanism capable of diversifying the proteome, creating a range of proteins from a single genomically encoded gene in a temporally regulated, tissue-specific, condition-dependent way and providing the organism with a new means for acclimation and adaptation. Indeed, several studies have demonstrated how recoding levels at specific sites do change as a function of the organism’s condition (Garrett and Rosenthal 2012; Robinson et al., 2016; Gallo et al., 2017; Porath 2017; Terajima et al., 2017; Yablonovitch et al., 2017). Importantly, many studies have demonstrated altered editing of individual recoding targets in various disease states (Gallo et al., 2017). However, these interesting examples notwithstanding, the extent to which the recoding phenomenon is actually used as a means for proteome diversification is still under debate. In fact, several recent studies have raised the possibility that nonsynonymous editing may be used to compensate for otherwise deleterious G-to-A mutations, rather than allowing for the two alleles (A and G) to co-exist (Jiang and Zhang 2019; Mai and Chuang 2019; Popitsch et al., 2020). Thus, a high N/S ratio is not sufficient to prove that editing serves for adaptation through proteome diversification.

The behavioral, physiological, and morphological plasticity exhibited by social insects suggests an important role for epigenetic mechanisms. Differential RNA editing was previously shown for different castes of the highly social leaf-cutting ant *Acromyrmex echinatior* (Li et al., 2014), as well as for bumblebee workers (Porath et al., 2019). Our results expand these finding to another species, further supporting the notion of RNA editing as a source of proteomic complexity, that may be recruited to provide phenotypic variation among genetically identical individuals.

ADAR enzymes and the recoding phenomenon are widely conserved across metazoan. However, the repertoire of recoding sites varies considerably and seems to have developed almost independently in different clades. Here, we point out that the same gene targets are being edited in distant species, even if the exact locations of the editing sites within the transcript vary. A particularly interesting example is that of nonsynonymous auto-editing of *Adar*, shown to affect ADAR activity in bumblebees, flies, and even mammals (Rueter et al., 1999; Savva et al., 2012a; Porath et al., 2019), possibly serving as global editing autoregulation. It thus seems possible that the selective advantage in having multiple versions of the protein product is shared by the target genes, while there is more than one specific way to achieve this diversity.
through recoding. Clades developed along different evolutionary routes have converged on different implementations (in terms of the specific editing site) of the same diversifying solution. These ideas should be further tested in the future, using larger data sets and across additional species and clades. Moreover, as biochemical and functional understanding of the impact of the different edits are gained, it will be possible to look into a possible common effect for the different edits of the common targets.

**Limitations of the study**

The editomes of the bee species analyzed here are based on a limited number of samples. They probably represent a subset of the actual repertoire of editing sites. The results should be re-tested when more data are available. In particular, the honeybee editome was built using drone samples. Thus, some individual differential editing between the nurses and foragers sub-castes may have missed worker-specific sites. However, the global increased editing in foragers, supported by a higher Adar expression, is probably robust.

**Resource availability**

**Lead contact**
Further information and requests for resources should be directly and will be fulfilled by the Lead Contact, Jian Lu (luj@pku.edu.cn).

**Materials availability**
This study did not generate new unique materials.

**Data and code availability**
All deep-sequencing data generated in this study were deposited in the China National Genomics Data Center Genome Sequence Archive (GSA) under accession number CRA002262. Other deep-sequencing data analyzed were downloaded from SRA as follows. Brains of honeybee workers: accession numbers SRR445999 to SRR446004 (reverted nurse) and SRR446005 to SRR446010 (forager) (Herb et al., 2012). Bumblebee: SRP166322 (Porath et al., 2019). D. mel head: SRP067542 (Zhang et al., 2018). D. sim head: SRP074828 (Duan et al., 2017). D. pse: DRR055250 and DRR055251 (Nozawa et al., 2016). D. wil: SRR341127 and SRR341129 (Meisel et al., 2012). D. moj: SRR037508-SRR037518 (generated by the Drosophila modENCODE project). D. sec, and D. ere data are unpublished data generated in our own lab. The reference genomes versions used for Drosophila species are Dmel_r6.04 (http://flybase.org/) for D. mel, and versions droSim1, droSec1, droEre1, dp4, droWil1, and droMoj3 downloaded from UCSC Genome Browser (http://genome.ucsc.edu/) for the other fly species. For the Bombus terrestris, we used version Bter_1.0.

**Methods**
All methods can be found in the accompanying Transparent methods supplemental file.

**Supplemental information**
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101983.

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**Author contributions**
J.L designed the research; J.X.H. contributed the honeybee materials; S.Q.D. performed the research; Y.G.D., H.T.P., E.E., and J.L. performed the bioinformatics analyses; E.E. and J.L. wrote the paper.
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Declaration of interests

The authors declare no competing interests.

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Supplemental Information

A-to-I RNA editing in honeybees shows signals of adaptation and convergent evolution

Yuange Duan, Shengqian Dou, Hagit T. Porath, Jiaxing Huang, Eli Eisenberg, and Jian Lu
Supplemental Figures

Figure S1. Per-individual DNA-Seq coverage at detected SNPs sites (left) and the variation level (alternative reads count / coverage) of these SNPs (right), Related to Figure1.
Figure S2. Distribution of DNA-RNA mismatch types for sites detected in heads, thoraxes, or abdomens of honeybees, Related to Figure 1.
Figure S3. Local sequence motif around editing sites (left), SNPs (middle) or unedited adenosines in genome (right), Related to Figure 1.
Figure S4. Histogram of per-individual DNA-Seq coverage at candidate RNA editing sites and other variations in each sample, Related to Figure 1.
Figure S5. Venn diagram demonstrating the overlaps of editing sites in four honeybee drones, Related to Figure 1.
Figure S6. Functional annotation of editing sites detected in heads, thoraxes, or abdomens of honeybees, Related to Figure 1.
Figure S7. Venn diagram demonstrating the overlaps of editing sites in three honeybee tissues, Related to Figure 1.
Figure S8. *Adar* expression is highest in heads (left) and is positively correlated (Spearman’s correlation) to cumulative editing levels in each sample (right), Related to Figure 1. The cumulative editing level is the sum of the editing level over all sites. Error bars represent the standard error of mean. *Adar* expression values were calculated by the DESeq2 software.
**Figure S9. Non-conserved editing sites in gene NaCP60E and eag, Related to Figure 3.**

*Drosophila*-specific editing sites are colored in red. Many of these sites are genomically encoded as G in the honeybee genome.
**tipE** (temperature-induced paralytic E)

|          | site1 | site2 | Site1 level |
|----------|-------|-------|-------------|
| *D. mel* | CTG   | CTG   |             |
| *D. sim* | CTG   | CTG   |             |
| *D. sec* | CTG   | CTG   |             |
| *D. ere* | CTG   | CTG   |             |
| *B. ter* | CTC   | AGC   |             |
| *A. mel* | CTC   | AGC   |             |

|          | site1 | site2 | Site1 level |
|----------|-------|-------|-------------|
| *D. mel* | GGC   | ACC   |             |
| *D. sim* | GGC   | ACA   |             |
| *D. sec* | GGC   | ACA   |             |
| *D. ere* | GGC   | ACA   |             |
| *B. ter* | AGC   | ACC   | 0.63        |
| *A. mel* | AGC   | ACC   | 0.70        |

**Figure S10. Bee-specific editing sites in **tipE**, Related to Figure 3.** Two *tipE* recoding sites are observed in honeybee (orthologous gene GB47375), one of which is also seen in bumblebee (orthologous gene XM_003393369) (Porath et al. 2019), colored in red. Editing is not seen at this site in *Drosophila*, as the genome encodes edited form of the protein.
Figure S11. *Adar* expression in drones and workers, Related to Figure 4. Reads count was normalized by DESeq2. Error bars represent standard error of mean. T-test was used to calculate the statistical significance. No significant difference was obtained (*P* = 0.23).
Figure S12. Comparison of editing levels in two sub-castes of workers, Related to Figure 4. The numbers of editing sites with levels higher in foragers (purple) and nurses (orange) are shown. Different cutoffs of sequencing coverages were used.
Figure S13. The observed and expected numbers of orthologous genes with editing in coding regions, Related to Figure 5. Genes with RPKM > 10 were chosen to calculate the expected numbers.
Figure S14. Venn diagram demonstrating the overlap between orthologous genes with editing sites in coding regions, Related to Figure 5. Strict criteria for determining sites as edited or not-edited were applied. The details are described in the main text.
Figure S15. The observed and expected numbers of orthologous genes with editing in coding regions, Related to Figure 5. Strict criteria for determining sites as edited or not-edited were applied. The details are described in the main text. Genes with RPKM > 1 or RPKM > 10 were chosen to calculate the expected numbers.
Figure S16. Histograms showing the cluster length of normal editing sites (left) and hyper-editing sites (right), Related to Figure 1. Editing sites within 100bps were clustered.
Figure S17. The nucleotide context of hyper-editing sites, Related to Figure 1. Pos -1, the upstream nucleotide. Pos +1, the downstream nucleotide.
Figure S18. Venn diagram demonstrating the overlaps of hyper-editing sites in three tissues of honeybee drones, Related to Figure 1.
Figure S19. Venn diagram demonstrating the overlaps of hyper-editing sites in four honeybee drones, Related to Figure 1.
Figure S20. Annotation of hyper-editing sites for repetitive and non-repetitive regions, separately, Related to Figure 1. The hyper-editing sites in repeats have lower fractions of nonsynonymous and synonymous sites but have higher fractions of intronic and intergenic sites.
Figure S21. Histograms of sequencing coverage of normal editing sites (left) and hyper-editing sites (right), Related to Figure 1. Editing sites in all samples were used.
## Supplemental Tables

**Table S1. Number of SNPs identified in each individual honeybee, Related to Figure 1.**

|          | D1    | D2    | D3    | D4    | All    |
|----------|-------|-------|-------|-------|--------|
| 5’UTR    | 2,573 | 2,495 | 2,480 | 2,445 | 3,769  |
| Nonsyn   | 8,344 | 8,032 | 8,090 | 8,296 | 12,494 |
| Syn      | 20,771| 19,509| 19,904| 19,814| 30,559 |
| 3’UTR    | 10,199| 9,518 | 9,640 | 9,688 | 14,779 |
| Intron   | 359,949| 347,684| 343,875| 352,415| 544,971|
| Intergenic| 456,956| 439,499| 440,563| 451,767| 696,412|
| Other    | 157   | 153   | 154   | 159   | 241    |
| Total    | 858,949| 826,890| 824,706| 844,584| 1,303,225|
| Drones | Tissue | Total reads (M) | Reference genome | Masked genome |
|--------|--------|----------------|-----------------|---------------|
|        |        |                | Uniquely mapped reads (M) | Uniquely mapped reads (%) | Uniquely mapped reads (M) | Uniquely mapped reads (%) |
| D1     | Head   | 25.0          | 17.9            | 71.6          | 17.8          | 71.1          |
|        | Thorax | 25.8          | 16.8            | 65.1          | 16.1          | 62.4          |
|        | Abdomen | 23.5      | 17.2            | 73.2          | 17.2          | 73.1          |
|        | Head   | 18.8          | 14.2            | 75.3          | 14.1          | 74.8          |
| D2     | Thorax | 30.8          | 18.4            | 59.6          | 17.1          | 55.6          |
|        | Abdomen | 28.5      | 21.5            | 75.3          | 21.4          | 75.2          |
|        | Head   | 37.2          | 12.4            | 33.4          | 13.0          | 34.9          |
| D3     | Thorax | 32.9          | 10.2            | 31.1          | 10.6          | 32.3          |
|        | Abdomen | 43.1      | 12.8            | 29.8          | 13.4          | 31.2          |
|        | Head   | 48.4          | 15.2            | 31.4          | 15.5          | 32.1          |
| D4     | Thorax | 31.9          | 10.4            | 32.6          | 10.7          | 33.4          |
|        | Abdomen | 32.9      | 9.2             | 28.1          | 9.5           | 28.8          |

Reference genome: the reads mapped to the reference honeybee genome.

Masked genome: the reads mapped to the genome sequences which have been replaced with the alternative alleles at SNP sites.
Table S4. Number of editing sites in each sample, Related to Figure 1 and Figure2.

| Drones | Tissue | Total sites | Nonsyn (N) | Syn (S) | N/S | P value |
|--------|--------|-------------|------------|---------|-----|---------|
|        | Head   | 278         | 72         | 8       | 9.0 | 1.59E-05 |
| D1     | Thorax | 71          | 13         | 4       | 3.3 | 0.610   |
|        | Abdomen| 94          | 14         | 1       | 14.0| 0.0491  |
|        | Head   | 262         | 73         | 7       | 10.4| 4.36E-06 |
| D2     | Thorax | 70          | 12         | 5       | 2.4 | 1 |
|        | Abdomen| 96          | 14         | 1       | 14.0| 0.0491  |
|        | Head   | 177         | 44         | 6       | 7.3 | 0.00318 |
| D3     | Thorax | 40          | 3          | 0       | Inf | 0.558   |
|        | Abdomen| 55          | 0          | 0       | 0.0 | 1 |
|        | Head   | 208         | 59         | 6       | 9.8 | 6.67E-05 |
| D4     | Thorax | 41          | 3          | 1       | 3.0 | 1 |
|        | Abdomen| 60          | 5          | 1       | 5.0 | 0.674   |
| All head|       | 376         | 108        | 8       | 13.5| 8.46E-10 |
| All thorax |       | 106         | 14         | 5       | 2.8 | 0.807   |
| All abdomen |     | 137         | 16         | 1       | 16.0| 0.0318  |
| All    |        | 407         | 111        | 9       | 12.3| 1.02E-09 |

Totally 407 unique editing sites were identified. The numbers of editing sites in each sample are listed respectively. “All head” is the number of sites that appear in at least one head sample among the four individuals. The same goes for “All thorax” and “All abdomen”.

P value: The P value of the observed N/S ratios compared to the expected N/S ratio under neutral evolution calculated from Fisher’s exact tests.
Table S5. Conserved coding editing sites between honeybee and bumblebee, Related to Figure 3.

| Site               | Gene   | Editing level | Site               | Gene               | Editing level |
|--------------------|--------|---------------|--------------------|--------------------|---------------|
| Group8.6:728981    | GB40519| 0.48          | Group16.3:2744556  | XM_003401834       | 0.87          |
| Group8.6:728990    | GB40519| 0.42          | Group16.3:2744565  | XM_003401834       | 0.41          |
| GroupUn131:18550   | GB46982| 0.31          | Group10.1:6435961  | XM_003398309       | 0.93          |
| GroupUn131:18489   | GB46982| 0.79          | Group10.1:6436022  | XM_003398309       | 0.97          |
| GroupUn131:18488   | GB46982| 0.81          | Group10.1:6436023  | XM_003398309       | 0.98          |
| Group1.29:1345934  | GB47508| 0.50          | Group1.8:1831958   | XM_003393355       | 0.88          |
| Group1.29:1345950  | GB47508| 0.34          | Group1.8:1831942   | XM_003393355       | 0.79          |
| Group15.19:753243  | GB50090| 0.42          | Group15.5:2877069  | XM_003401089       | 0.45          |
| Group1.29:1555391  | GB47375| 0.70          | Group1.8:1578895   | XM_003393369       | 0.63          |
Table S6. Mapping summary of each worker sample, Related to Figure 4.

| Sample     | Total reads (M) | Uniquely mapped reads (M) | Uniquely mapped reads (%) |
|------------|-----------------|---------------------------|---------------------------|
| SRR445999  | 213.8           | 118.2                     | 55.3                      |
| SRR446000  | 295.4           | 203.8                     | 69.0                      |
| SRR446001  | 152.0           | 95.2                      | 62.6                      |
| SRR446002  | 154.5           | 87.9                      | 56.9                      |
| SRR446003  | 138.6           | 89.2                      | 64.4                      |
| SRR446004  | 226.0           | 149.9                     | 66.3                      |
| Nurse pool | 1180.3          | 744.2                     | 63.0                      |
| SRR446005  | 178.9           | 97.5                      | 54.5                      |
| SRR446006  | 193.1           | 130.3                     | 67.5                      |
| SRR446007  | 121.2           | 72.8                      | 60.1                      |
| SRR446008  | 164.6           | 100.2                     | 60.9                      |
| SRR446009  | 179.7           | 119.4                     | 66.5                      |
| SRR446010  | 139.4           | 84.5                      | 60.6                      |
| Forager pool | 976.8           | 604.6                     | 61.9                      |
Table S9. Shared genes with CDS editing across three species, Related to Figure 5.

| D. melanogaster | A. mellifera | B. terrestris |
|-----------------|--------------|--------------|
| FBgn0086372     | GB43906      | XM_003403010 |
| FBgn0263354     | GB48155      | XM_003396523 |
| FBgn0263111     | GB51897      | XM_003394356 |
| FBgn0035538     | GB54467      | XM_003395885 |
| FBgn0004242     | GB54827      | XM_003402654 |
| FBgn0262483     | GB55567      | XM_003402632 |

The gene IDs of *B. terrestris* are retrieved from the bumblebee study (Porath et al. 2019) according to the editing site.
**Table S10. Number of hyper-editing sites in each sample, Related to Figure 1.**

| Drones | Tissue | Total sites | 5’UTR | Nonsyn | Syn | 3’UTR | Intron | Intergenic | other |
|--------|--------|-------------|-------|--------|-----|-------|--------|------------|-------|
|        | Head   | 1,644       | 7     | 86     | 50  | 72    | 509    | 915        | 5     |
| D1     | Thorax | 1,576       | 32    | 182    | 100 | 58    | 387    | 801        | 16    |
|        | Abdomen| 2,015       | 29    | 215    | 114 | 65    | 607    | 977        | 8     |
|        | Head   | 2,416       | 110   | 290    | 170 | 37    | 660    | 1,110      | 39    |
| D2     | Thorax | 484         | 19    | 54     | 44  | 46    | 143    | 172        | 6     |
|        | Abdomen| 2,202       | 64    | 221    | 153 | 68    | 730    | 954        | 12    |
|        | Head   | 2,393       | 37    | 196    | 135 | 52    | 817    | 1,143      | 13    |
| D3     | Thorax | 571         | 0     | 44     | 32  | 9     | 168    | 318        | 0     |
|        | Abdomen| 1,019       | 6     | 57     | 49  | 17    | 291    | 598        | 1     |
|        | Head   | 1,756       | 11    | 145    | 92  | 29    | 495    | 977        | 7     |
| D4     | Thorax | 1,716       | 18    | 189    | 102 | 42    | 602    | 755        | 8     |
|        | Abdomen| 306         | 0     | 18     | 8   | 17    | 123    | 138        | 2     |
| Pool   |        | 12,436      | 276   | 1,355  | 845 | 488   | 3,784  | 5,591      | 97    |
**Transparent Methods**

**Honeybee collection**
Drones were raised in a colony of *Apis mellifera* by professional beekeepers in Jie Wu’s Lab, Chinese Academy of Agricultural Sciences. For each male drone adult (haploid), the head and thorax were separated with surgical scissors and the testis was dissected. Residual abdomen tissues were preserved for DNA extraction. All samples were flash-frozen in liquid nitrogen and stored at -80°C for further procedures.

**RNA extraction and mRNA-Seq of four drone individuals**
Four drone individuals were selected for mRNA-Seq. Total RNA was extracted from head, thorax, and testis from each individual, separately, using TRIzol reagent (Thermo Fisher). For two individuals (1 and 2), Poly(A)+ mRNAs were selected on oligo-dT25 DynaBeads (Thermo Fisher), while for the other two (3 and 4) RNAs were treated with Ribo-Zero Gold rRNA Removal Kit (Illumina) to remove the rRNA-derived fragments. All these purified RNAs were fragmented. The 40-80 nt fragments were purified from 15% TBE-Urea gels for deep sequencing, and subject to 3'-dephosphorylation with T4 Polynucleotide Kinase (NEB), 3'-ligation, 5'-phosphorylation with T4 Polynucleotide Kinase (NEB) and ATP, 5'-ligation, and reverse-transcription into cDNA with SuperScript™ III Reverse Transcriptase (Thermo Fisher). The cDNAs were PCR-amplified and size-selected in 20% TBE gels for fragments in correct ranges. Purified products were prepared for quality tests (Fragment Analyzer, Agilent Technologies) and sequencing (Illumina HiSeq-2500 sequencer; run type: single-end; read length: 50 nt).

**Genomic sequencing of four drone individuals**
To efficiently exclude SNPs from RNA editing sites, genomic DNA from abdomen tissue of each individual drone separately was extracted using the Genomic DNA Extraction Kit (TIANGEN) following manufacturer’s instructions. The library preparation and sequencing were performed in Biomedical Pioneering Innovation Center, Peking University (Illumina HiSeq-2500 sequencer; 100bp paired-end reads).

**Identification of SNPs in four drones**
Honeybee reference genome sequence (*A. mel 4.5*) was downloaded from BeeBase (http://hymenopteragenome.org/beebase/). For each of the four individual drones, we mapped
the DNA-Seq reads to the reference genome with BWA v0.7.4 (Li and Durbin 2009). PCR duplicates were removed using Picard v1.119, and SNPs were called using SAMtools mpileup (Li 2011) with default parameters.

**Identification of variation sites in four drones**

To identify reliable A-to-I RNA editing events, we employed two different aligners STAR (2.4.2a) (Dobin et al. 2013) and BWA (Li and Durbin 2009) to map the RNA-Seq reads to the reference genome (A. mel 4.5). For each BAM sequence alignment file, we extracted all the alignments with mapping quality \( \geq 10 \) using SAMtools 1.3.1 (Li 2011). The mismatches between RNA-Seq reads and reference genome were extracted using Sam2Tsv (Pierre 2015). Bases with mapping quality lower than 30, soft clipping bases, bases at the 10bp of reads’ ends and mismatches in repeat regions were discarded. Only mismatch sites supported by both STAR and BWA were retained.

Next, we produced for each individual drone its own version of the genome, by substituting the SNP sites found in this individual to the reference genome. For example, in individual drone1, 858,949 SNPs were identified, and the reference genome (A. mel 4.5) was modified by replacing the reference alleles with the alternative alleles at all these SNP sites of drone1. The modified genome was regarded as the masked genome for drone1. Note that drone individuals are haploids and there are no allele-specific SNPs. Thus, the masked genome is the entire haploid genome sequence of each individual. We then mapped again the RNA-Seq reads of drone 1 to its masked genome, using both STAR and BWA as before. The same pipeline was applied to drones 2, 3, and 4.

Ideally, one would need to map the reads only to the masked genome (presumably, the true genome of each individual drone). However, our SNP detection is imperfect, and false positive SNPs identification could lead to RNA-DNA variations against the masked genome. Thus, to be conservative, we retained only variation sites supported by mapping to both the reference genome and the masked genomes. These RNA-DNA variations are unlikely to include many variations due to genomic polymorphisms, and should be enriched in RNA level alterations. We further discarded variations occurring in less than three out of twelve samples (head, thorax, and abdomen of four honeybee individuals), resulting in 1,742 candidate variation sites. Of these, 1417 (81.3%) were A-to-G.
Defining A-to-I RNA editing sites

Finally, we pooled the reads of the four drones for each of the tissues, and calculated the probability $P_k(E_0)$ that the mismatches at position $E_0$ (each of the 1,742 sites found above) observed in tissue $k$ ($k =$ head, thorax, or abdomen) can be explained by sequencing error (binomial test, with A-to-G error rate $\varepsilon = 0.00167$ (Duan et al. 2017), followed by Benjamini-Hochberg multiple testing correction with FDR = 0.05 (Benjamini and Hochberg 1995)). The multiple testing correction took into account all $\sim 230$ Mbp of the honeybee genome (excluding “N”s), at which the above variations may have been detected. Pooled data was analyzed separately for each tissue.

Editing level was estimated as $\frac{G}{(G+A)}$, in which $G$ is the alternative allele count and $A$ is the reference allele count.

For the downloaded deep-sequencing data of brains of honeybee workers (Herb et al. 2012), we mapped the reads with STAR and called variants with SAMtools mpileup (Li 2011) with default parameters as we mentioned above. We directly retrieved the reads counts on the candidate editing sites and looked at the editing levels. Alignments with mapping quality $\geq 10$ were maintained and no other filters were applied. Editing levels was compared between different castes or sub-castes using Fisher’s exact test. Editing levels in other species (e.g. sites in genes Shab and qvr), were evaluated based on the datasets described below. We mapped the sequencing reads of each species to the reference genome using STAR (Dobin et al. 2013). The variants were called with SAMtools mpileup (Li 2011). The genomic coordinates were transferred from D. melanogaster to the other fly species by using liftOver chain downloaded from UCSC Genome Browser (http://genome.ucsc.edu/).

Annotation of A-to-I RNA editing sites

We used the software SnpEff version 4.3 (Cingolani et al. 2012b) for functional annotation of the editing sites. The canonical transcript of each gene, as defined by the software, was chosen for annotation.

Expected N/S ratio

To find the expected N/S ratio under neutrality, we replaced all adenosines in all coding regions with guanosines, one at a time, and used the software SnpEff (Cingolani et al. 2012a), (version 4.3, parameters “-ud -canon -v amel_OGSv3.2”) to determine how many of these
substitutions would cause a nonsynonymous or synonymous change. In total, 4,492,737 nonsynonymous and 1,986,130 synonymous substitutions were obtained, and therefore the expected N/S ratio was 2.26. To verify that this ratio is not biased by editing being limited to a small set of genes, we have repeated the calculation focusing only on adenosines in genes harboring editing sites. Here we have found 68,220 nonsynonymous and 30,150 synonymous substitutions were obtained, and therefore the expected N/S ratio was very similar, 2.26. We therefore used the ratio 2.26 is all further analyses.

**Conservation analysis**
Orthologous genes (between *D. melanogaster* and honeybee, or between honeybee and bumblebee) were defined as reciprocal best hits of pairwise BLASTP (Camacho et al. 2009). The protein sequences of the orthologous genes were then aligned with the clustalw program (Thompson et al. 1994), and the alignments of RNA CDS were achieved by the tranalign program (Rice et al. 2000) based on the corresponding protein alignments. We used *D. melanogaster* reference genome version FlyBase_r6.04. For the bumblebee genome we used NCBI reference sequence version Bter_1.0. NCBI RefSeq annotations for genes and coding regions were downloaded from the BeeBase site (http://hymenopteragenome.org/beebase) on 7 May, 2014.

To calculate the expected numbers of genes sharing CDS editing at different positions within the same gene we first determined the set of expressed genes in in each of the three species as those genes with RPKM (reads per kilobase per million mapped reads) values exceeding a given cutoff. Then, we have randomly chosen 53 honeybee genes, 101 bumblebee genes and 312 fly genes (the numbers of genes harboring editing sites in CDS per species, excluding sites conserved in any two or three species), and looked for the number of orthologous gene groups reoccurring in any two of these sets or in all three of them. For each expression cutoff, the distribution of the expected overlap was calculated using 1,000,000 such random choices, and the p-value was calculated comparing the actual overlap to this distribution.

**mRNA secondary structure prediction**
We folded the mRNAs of honeybee with RNALfold (Lorenz et al. 2011). Local structures with Z score < -3 were defined as stable hairpins.
PCA Analysis

The PCA analysis of editing levels and gene expression levels was performed by function "princomp()" in R. The first two principal components (PC1 and PC2) were used to plot the graph.

Supplemental Note

The hyper-editing pipeline detects many sites in lowly expressed regions

We used the hyper editing script (Porath et al. 2014) with parameters suitable for 50bp reads "0.05 0.8 30 0.6 0.1 0.8 0.2" (Porath et al. 2014) to identify densely edited clusters, with a total of 12,436 unique sites. Of these, only 2,056 (17%) are located in repetitive regions. This fraction is higher than the one found by our main procedure, but still much lower than the one observed in most other species analyzed. A partial explanation is the rather short reads, preventing reliable alignment of heavily edited reads.

    Hyper-editing sites are clustered. Clustering together editing sites (distance < 100 bps) we find that hyper-editing sites tend to cluster in big clusters (larger than the ones required for their detection) (Figure S16). Notably, half of regular editing sites are also clustered.

    The sequence context of these hyper-editing sites is similar to the pattern observed in bumblebees (Porath et al. 2019), where the upstream nucleotide shows a preference of A > T > C > G and the downstream nucleotide exhibits A > T > G > C (Figure S17). In each sample, 306–2,416 hyper editing sites were detected in different tissues of four honeybee individuals (Table S10). Heads bear the greatest numbers of hyper editing sites than other tissues (Figure S18), although the sites from heads of different individuals do not show much overlap (Figure S19).

    Compared to the editing sites identified by our regular pipeline, those hyper-editing sites have a remarkably higher fraction in intergenic and intron regions (Figure S20). In fact, only 52 of these 12,436 sites overlap with the 407 editing sites identified by traditional pipeline, and 45 of the overlapped sites are located in intergenic or intronic region. This result agrees with the finding that hyper-edited regions are often lowly expressed (Porath et al. 2017), and corroborated by the sequencing coverage spectrum which for hyper-editing sites is skewed to lower depths compared to normal editing sites (Figure S21). None of the hyper-editing sites overlap with conserved or task-related editing sites in the bumblebee, which further suggests...
that hyper-edited regions are weakly expressed and likely to be non-conserved.
Supplemental References (Related to Supplemental Figures, Supplemental Tables, and Transparent Methods)

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