Reviewers’ comments:

Reviewer #1 (Remarks to the Author):

This manuscript studied the natural selection patterns of mites with different feeding habits through comparative genomics method. They identified the positively selected genes and gene family expansion related with herbivorous, blood-feeding, and fat-feeding habits. These potential candidate genes provide insights into the dietary evolution of mites under different selection pressures. I have some comments for improving the manuscript.

1. Line 98: how many genes were used for phylogenetic tree construction?
2. Line 99: should be “relaxed”.
3. Line 114: “detected to be common in the groups of ...”.
4. Line 136: should be “correlated with”.
5. Line 138: “the number of PSGs (more than 20) was much higher...”.
6. Lines 140-141: I doubt whether there were more than 200 sites under positive selection pressure for one gene. So, please check the DNA sequence alignments of this gene among these species, to avoid the sequence misalignment.
7. Lines 163 and 189: “owing to” should be “belonging to”?
8. Lines 167 and 209: please change as “manual filtration”.
9. Line 193: please delete “cleverly”.
10. Lines 205 and 252: When using the PZ and AA firstly, please give its full name.
11. Lines 249-251: please give related reference to support the absence of FAAH2 in rats and mice.
12. Line 262: “any expansion gene families”??
13. Line 263: What do you mean about “genetic expansion”? 
14. Lines 274-275: What do your mean about “Different levels of genetic convergence under the same diet was identified”? 
15. Figure 1: the species Latin names should be italic.
16. Figure 3A and 3B. It seemed that corrected P-values were not insignificant. How do you think about the results?
17. Figure 3D: the figure is unclear.

Reviewer #2 (Remarks to the Author):

This study provides the genetic evidence of mites and ticks adaptation to different dietary habits based on the comparative analysis of genomes of 16 arachnids. Authors analyzed mites and ticks of adaptations to Herbivory, blood-sucking and Fat-feeding by using the methods such as gene family expansion and gene selection pressure.

The finding of this working is interesting, but I got some comments as below:

Line 89. The genome assembles in Table 1 used in this analysis should be cited clearly and separately, so that the other researcher can find correct paper resources to repeat this work.

Line 90. Authors should submit the gene annotations for the 5 reannotated genomes to public database.

Line 93. In this manuscript, the authors expected to use the annotated gene sets, especially the protein-coding gene sets, to understand the mite evolution. However, only the genome assembly qualities of the selected mite and tick genomes were tested with BUSCO. To exam the qualities of annotated gene sets in this mite and tick genomes apparently more important. Authors should also use the BUSCO to evaluate the qualities of annotated gene sets.
There was a long-time debate for the Acari is diphyletic or monophyletic. It would be good to discuss this based on figure 1.

Authors should provide the source of RNAseq data for the four spider mite populations.

How were the top 200 genes found based on Sixteen samples from four population of spider mite? Please the authors provide the method and the gene list. Which method did authors used for GO and KEGG enrichment for the expanded genes? Authors should clarify in the method part.

How were the top 50 genes found based on Sixteen samples from four population of spider mite? Please the authors provide the method and the gene list.

Please provide high resolution Supplementary Fig. 4. It is not readable. Authors should label the bootstrap numbers on the tree, and also provide gene alignment file and the tree construction method in the method part.

For the KEGG and GO enrichment analysis, please authors explain why they selected the far related Drosophila melanogaster as background set. The Kobas used in this enrichment analysis supports a wide range of species, including mite and tick models species, such as Ixodes scapularis and Tetranychus urticae. The insects are innately very different from the mites and ticks in some pathways. Please authors explain how the gene enriched with the D. melanogaster gene sets could reflect the mite and tick biology? What is the p-value or q-value cutoff for Kobas result?

Reviewer #3 (Remarks to the Author):

In this study, the authors compared genomes of 16 Arachnida species, and showed different patterns of diet adaptations. I have a few concerns:
1. The manuscript was mainly based on published genome data. Expression data of spider mites were also used. It is not clear why only expression data of herbivory species were used. It seems the authors are choosing data arbitrary. I think they should provide more detailed and scientific explanations about data selection.
2. The authors showed that Arachnida species with different diet showed different evolutionary patterns based on some regular analyses. These results are expectable. It is not clear what is the key scientific problem the authors want to solve based on their analyses. What is the purpose of doing these analyses as well as publishing a paper, and how these results will serve the purpose? In the abstract and the conclusions, the authors mentioned twice: “These different genetic bases provide a new perspective for the study of the evolution and diversification of this group, and offers potential drug targets for pest control.” These statements are repetitive, and are very vague. What is the “new perspective” and how to offer potential drug targets? Details should be discussed at least. In addition, not all mite species analyzed in this study are pests. For example, Metaseiulus occidentalis is a predatory mite natural enemy.
3. There are many typos and formatting errors in manuscript (including the references). I think the authors should pay more attentions to fix these problems.
Dear Reviewers,

Many thanks for your important and helpful suggestions for our manuscript entitled “Genomic implications in diet evolution: Comparative analysis of mite genomes reveals positive selection for diet adaptation” (COMMSBIO-20-3585). In the following responses, we have carefully addressed all the issues, and we have revised our manuscript accordingly. Our references to line numbers refer to the no markup view that we have uploaded as a ‘Acari_Diet_main_text_changes_tracked.docx’ file. All changes have been accepted in the clean revised manuscript uploaded as a ‘Acari_Diet_supplementary_main_text_clean.docx’ file. We hope you find that we have adequately addressed all of the suggestions and that our manuscript is now suitable for publication. Please let us know if you have any further questions or suggestions.

Our point-by-point responses are as follows:

Reviewer #1 (Remarks to the Author):
This manuscript studied the natural selection patterns of mites with different feeding habits through comparative genomics method. They identified the positively selected genes and gene family expansion related with herbivorous, blood-feeding, and fat-feeding habits. These potential candidate genes provide insights into the dietary evolution of mites under different selection pressures. I have some comments for improving the manuscript.

Thank you for your recognition! We appreciate your positive comments and we made corresponding revisions to your suggestions.

Comment (1) Line 98: how many genes were used for phylogenetic tree construction

A total of 17,910 homologous genes were identified. Due to the different techniques of sequencing and assembly of the 16 genomes, critical filtration has been carried out in the species selection (lines
285-291) and reannotation was conducted for 6 genomes. However, the discrepancy of the amount of gene information cannot be avoided and the number of single-copy homologous genes annotated is limited. Finally, 65 single-copy homologous genes with 48,831 nucleotides were annotated by all 16 species and applied to construct the species tree.

Comment (2) Line 99: should be “relaxed”.
Thank you for your suggestion. We have corrected it in line 101.

Comment (3) Line 114: “detected to be common in the groups of …”.
Thank you for your suggestion. We have corrected it in line 115.

Comment (4) Line 136: should be “correlated with”.
Thank you for your suggestion. We have corrected it in line 130.

Comment (5) Line 138: “the number of PSGs (more than 20) was much higher….”.
Thank you for your suggestion. We have corrected it in line 140.

Comment (6) Lines 140-141: I doubt whether there were more than 200 sites under positive selection pressure for one gene. So, please check the DNA sequence alignments of this gene among these species, to avoid the sequence misalignment.

Gene CES2 (ID: OG0000064) is an outlier in blood-feeding group shown in Fig1D with a total of 1,128 amino acid sites after deleting aligned gaps. 285 positively selected sites were found, among which 197 sites with a posterior probability of over 95% were detected. We haven't found any errors in its result of sequence alignment (See following figure 1).
Figure 1. Protein sequence alignments of carboxylesterase (After deleting the gaps)

Comment (7) Lines 163 and 189: “owing to” should be “belonging to”?
Thank you for your suggestion. We have revised it to “Cyp18a1, a cytochrome P450 enzyme” in line 165.

Comment (8) Lines 167 and 209: please change as “manual filtration”.
Thank you for your suggestion. We have corrected it in line 168 and line 212.

Comment (9) Line 193: please delete “cleverly”.
Thank you for your suggestion. We have corrected it.

Comment (10) Lines 205 and 252: When using the PZ and AA firstly, please give its full name.
Thank you for your suggestion. Arachidonate acid (AA) was first mentioned in line 248. Protein Z (PZ) was supplemented in the revised manuscript, see line 209.

Comment (11) Lines 249-251: please give related reference to support the absence of FAAH2 in rats and mice.
Thank you for your suggestion! We have added this reference no.62 in the revised manuscript.
Comment (12) Line 262: “any expansion gene families”?
We are sorry for the confusing expression. We have revised it to “no common expanded gene family was found in this group” (see line 151).

Comment (13) Line 263: What do you mean about “genetic expansion”?
We are sorry for the confusing expression. We have revised it to “expanded gene families” in line 150.

Comment (14) Lines 274-275: What do your mean about “Different levels of genetic convergence under the same diet was identified”?
We are sorry for the confusing expression. We have deleted the sentence.

Comment (15) Figure 1: the species Latin names should be italic.
Thank you for your suggestion! We have revised it in Figure 1.

Comment (16) Figure 3A and 3B. It seemed that corrected P-values were not insignificant. How do you think about the results?
Thanks for raising this issue! To find the significant pathways and gene ontology, we set the cutoff of P value < 0.05 in Fisher's exact test and FDR-corrected P < 0.1 according to the thresholds recommended in the previous studies (Liu, Yao-Zhong et al., 2017; Hulsegge, I et al., 2017). The KEGG pathway and Gene ontology related to detoxification function mentioned in the lines 165-181 (the following table 1) were considered statistically significant according to the thresholds.

| Term                          | Database         | P-Value    | Corrected P-Value |
|-------------------------------|------------------|------------|-------------------|
| Response to stimulus          | Gene Ontology    | 2.84E-05   | 0.0031299         |
| Response to external stimulus | Gene Ontology    | 3.93E-05   | 0.0037041         |
| ABC transporters              | KEGG PATHWAY     | 0.0054869  | 0.0910888         |

1. Liu, Yao-Zhong, Lei Zhang, Astrid M. Roy-Engel, Shigeki Saito, Joseph A. Lasky, Guangdi Wang, and He Wang. "Carcinogenic effects of oil dispersants: A KEGG pathway-based RNA-seq study of
human airway epithelial cells.” Gene 602 (2017): 16-23.

2. Hulsegge, I., A. Kommadath, and M.A. Smits. Globaltest and GOEAST: two different approaches for Gene Ontology analysis. BMC Proc, 2009. 3 Suppl 4: p. S10.

Comment (17) Figure 3D: the figure is unclear.
Sorry for the unclear version. We have split Figure 3 into new Figure 3 and 4 to show the information of figures better.

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Reviewer #2 (Remarks to the Author):

This study provides the genetic evidence of mites and ticks adaptation to different dietary habits based on the comparative analysis of genomes of 16 arachnids. Authors analyzed mites and ticks of adaptations to Herbivory, blood-sucking and Fat-feeding by using the methods such as gene family expansion and gene selection pressure.

We appreciate your positive comments and we have made corresponding revisions to your suggestions.

The finding of this working is interesting, but I got some comments as below:
Comment (1) Line 89. The genome assembles in Table 1 used in this analysis should be cited clearly and separately, so that the other researcher can find correct paper resources to repeat this work.

Thank you for your suggestion! We have supplemented separate citations for each genome in the revised manuscript (Table 1). Researchers can also obtain the genome data through the GenBank Assembly Accession ID in Table 1.

Comment (2) Line 90. Authors should submit the gene annotations for the 5 reannotated genomes to public database.

Thank you for your suggestion! We have uploaded the annotation files of the reannotated genomes to Mendeley Data (https://data.mendeley.com/datasets/xm23f9mkdx/1). This information is noted in the line 404-406 in the revised manuscript.
Comment (3) Line 93. In this manuscript, the authors expected to use the annotated gene sets, especially the protein-coding gene sets, to understand the mite evolution. However, only the genome assembly qualities of the selected mite and tick genomes were tested with BUSCO. To exam the qualities of annotated gene sets in this mite and tick genomes apparently more important. Authors should also use the BUSCO to evaluate the qualities of annotated gene sets.

Thank you for your suggestion! We have finished the analysis and provided the BUSCO results in Supplementary Table 1. We observed high BUSCO scores of genome completeness (average 90.9%) and gene completeness (average 79.9%). The proportion of complete gene is not very high in species Steganacarus magnus and Hypochthonius rufulus. However, the two species were set as background species when conducted the positive selection analysis and would not affect the identification of positive selection signals.

Comment (4) Line 100. There was a long-time debate for the Acari is diphyletic or monophyletic. It would be good to discuss this based on figure 1.

Thank you for your suggestion! The debate about the monophyly of Acari has always been a hot topic. A recent study indicated that Acari constitutes a monophyletic group (Showed in the figure below) nested within a monophyletic Arachnida based on transcriptomic data from 95 species. (Jesus Lozano-Fernandez, et al., 2019). However, the study published at the same year recovered the major mite lineages by using ultraconserved genomic elements (UCEs) and found mites to be non-monophyletic (Van Dam, 2019). Our data was insufficient to address this problem, unless we add the groups that can represent all the spiders of Arachnida, such as whip spiders, llamshade spiders, mygalomorph spiders, hooded tickspiders, Sun Spiders, and so on. However, it is not the main focus of our study. It is a good idea for our follow-up research.

1. Lozano-Fernandez, Jesus, et al. "Increasing species sampling in chelicerate genomic-scale datasets provides support for monophyly of Acari and Arachnida." Nature communications 10.1 (2019): 1-8.

2. Van Dam, Matthew H., et al. "Advancing mite phylogenomics: designing ultraconserved elements for Acari phylogeny." Molecular ecology resources 19.2 (2019): 465-475.
(Jesus Lozano-Fernandez, et al., 2019)
Comment (5) Line 166 and 367. Authors should provide the source of RNAseq data for the four spider mite populations.

Thank you for your suggestion! RNAseq data was from the SRA database (Bioproject: PRJNA610897; Xue, Wenxin, et al., 2020). We have supplemented this information in the Method in the revised manuscript (see line 375).

1. Xue W et al., "Geographical distribution and molecular insights into abamectin and milbemectin cross-resistance in European field populations of Tetranychus urticae.", Pest Manag Sci, 2020 Aug;76(8):2569-2581

Comment (6) Line 167,168 and 172. How were the top 200 genes found based on Sixteen samples from four population of spider mite? Please the authors provide the method and the gene list. Which method did authors used for GO and KEGG enrichment for the expanded genes? Authors should
clarify in the method part.

Thanks for raising this issue! First, we conducted short sequence mapping using HISAT2 and calculated Fragments per kilobase per million mapped reads (FPKM) values for genes of each sample by Cufflinks (Version 2.2.1). Then, genes present in all sixteen samples were sorted by the median FPKM value. Then the 200 most highly expressed genes were selected and analyzed. KEGG and Gene ontology enrichment analysis were performed by KOBAS 3.0 with the cutoff of P value < 0.05 in Fisher's exact test and FDR-corrected P < 0.1 according to the thresholds recommended in the previous studies (Liu, Yao-Zhong et al., 2017; Hulsegge, I et al., 2017). The bubble diagrams of significantly enriched KEGG and Gene ontology were generated using R ggplot package. We have supplemented details in the Methods section (lines 374-387) and provided the gene list in the supplementary Table 8 in the revised manuscript.

1. Liu, Yao-Zhong, Lei Zhang, Astrid M. Roy-Engel, Shigeki Saito, Joseph A. Lasky, Guangdi Wang, and He Wang. "Carcinogenic effects of oil dispersants: A KEGG pathway-based RNA-seq study of human airway epithelial cells." Gene 602 (2017): 16-23.
2. Hulsegge, I., A. Kommadath, and M.A. Smits, Globaltest and GOEAST: two different approaches for Gene Ontology analysis. BMC Proc. 2009. 3 Suppl 4: p. S10.

Comment (7) Line 174. How were the top 50 genes found based on Sixteen samples from four populations of spider mite? Please the authors provide the method and the gene list.

The methods of selecting the 50 most highly expressed genes were consistent with those of selecting the 200 most highly expressed genes. We have supplemented details in the Methods section (lines 374-387) and provided the gene list in the supplementary Table 8 in the revised manuscript.

Comment (8) Line 211. Please provide high resolution Supplementary Fig. 4. It is not readable. Authors should label the bootstrap numbers on the tree, and also provide gene alignment file and the tree construction method in the method part.

Sorry for the unclear image and statement. We have replaced Supplementary Fig. 4 with a version of higher resolution and provided the detailed information of tree construction in the Method section.
in the revised manuscript. Supplementary Fig. 4 has been changed to Supplementary Fig. 3 in the revised manuscript. Since the tree figure focuses on displaying the expanded gene number and clades, we have provided the bootstrap value in a separate tree file instead of the tree figure to avoid shading the 212 branches. We have submitted this tree file with bootstrap value and gene alignment file to the Mendeley Data (https://data.mendeley.com/datasets/xm23f9mkdx/1) due to the big size of the files. We have provided a “Data availability” section in the revised manuscript (lines 404-406).

Comment (9) Line 332. For the KEGG and GO enrichment analysis, please authors explain why they selected the far related Drosophila melanogaster as background set. The Kobas used in this enrichment analysis supports a wide range of species, including mite and tick models species, such as Ixodes scapularis and Tetranychus urticae. The insects are innately very different from the mites and ticks in some pathways. Please authors explain how the gene enriched with the D. melanogaster gene sets could reflect the mite and tick biology? What is the p-value or q-value cutoff for Kobas result?

Thanks for raising this issue! There are several considerations: (1) D. melanogaster is a well-studied specie in KEGG and Gene ontology (GO) in Arthropod, sharing a common ancestor with ticks and mites. (2) KEGG database is available for all species while Gene ontology database is only available for I. scapularis, and D. melanogaster, not for T. urticae. (3) When D. melanogaster was the background specie, more gene data of the 200 most highly expressed genes could be obtained from KEGG and GO annotations to retain more information for enrichment analysis. When D. melanogaster was set as background set, 60 genes could be obtained from KEGG pathway while 147 genes could be obtained from GO annotation. When I. scapularis was set as background set, 53 genes could be obtained from KEGG pathway while 110 genes could be obtained from GO annotation. When T. urticae was set as background set, 57 genes could be obtained from KEGG pathway while no information was obtained from GO annotation. Among these genes obtained from KEGG annotation, more than 90 percent genes of I. scapularis and of T. urticae were annotated in D. melanogaster. Among the genes obtained from GO annotation, 88 percent genes of I. scapularis were annotated in D. melanogaster. Hence, we chose D. melanogaster as the background set to generate more information. We set a threshold of P value < 0.05 in Fisher's exact test and FDR-
corrected $P < 0.1$ for KOBAS result.

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Reviewer #3 (Remarks to the Author):

In this study, the authors compared genomes of 16 Arachnida species, and showed different patterns of diet adaptations. I have a few concerns:

Comment (1) The manuscript was mainly based on published genome data. Expression data of spider mites were also used. It is not clear why only expression data of herbivory species were used. It seems the authors are choosing data arbitrary. I think they should provide more detailed and scientific explanations about data selection.

Thanks for raising this issue! Our study aimed to explore the dietary adaptations of mites based on released genomes. We have supplemented the details of data selection in the revised manuscript (lines 285-291). “(1) All Arachnida species in the database (before 2020.06) were searched as candidates; (2) Genome with the best completeness score was selected as the representative if there were two or more genomes for one species; (3) Species with genome completeness less than 80% were eliminated; (4) Gene prediction was conducted for the genomes lack of gene annotation information; (5) Contigs less than 1kb were excluded from the whole analysis”. Finally, we chose one tick and fourteen mites, and the velvet spider as the background from 26 candidate species (see the following Table 1).

To see if the results of positive selection and gene expansion have been reflected in gene expression, we supplied transcriptome analysis for the different dietary groups. Sampling of mites is a big challenge, so we conducted database search for mite transcriptome of lipid-feeding, blood-feeding and plant-feeding species. The samples taken from whole blood or whole tissue transcriptome from natural individuals were included. After the database search, two samples of blood-feeding were insufficient to do biological repetition and excluded; four samples of lipid-feeding were included. Only “Ribosome” was statistically significant with correction Fisher test $P < 0.05$ and FDR corrected $P < 0.1$ (see the following Table 2). Finally, we have shown details for gene expression of
the herbivorous group in our manuscript (see line 170-185).
Table 1. Genomes feature of candidates

| Species                      | GenBank Assembly | Number of contigs | Total size of contigs | Number of contigs > 500 nt | Number of contigs > 1K nt | Number of contigs > 10K nt | N50 contig | BUSCO completeness          |
|------------------------------|------------------|-------------------|-----------------------|-----------------------------|--------------------------|-----------------------------|------------|-----------------------------|
|                              | Accession        |                   |                       |                             |                          |                             |            |                             |
| Achipteria coleoptrata       | GCA_000988765.1  | 70,955            | 87,501,022            | 41.20%                      | 26.60%                   | 1.50%                       | 3,583      | C:82.5%[S:80.5%,D:2.0%],F:9.2%,M:8.3%,n:303 |
| Brevipalpus yothersi         | GCA_003956705.1  | 2,451             | 70,567,388            | 99.90%                      | 96.90%                   | 62.20%                      | 56,520     | C:83.9%[S:82.2%,D:1.7%],F:3.0%,M:13.1%,n:303 |
| Dermatophagoides farinae     | GCA_000767015.1  | 7,089             | 51,638,314            | 91.30%                      | 83.70%                   | 22.50%                      | 14,557     | C:90.4%[S:90.1%,D:0.3%],F:3.3%,M:6.3%,n:303 |
| Dermatophagoides farinae     | GCA_002085665.1  | 1,716             | 91,934,661            | 100.00%                     | 99.90%                   | 92.00%                      | 188,869    | C:95.4%[S:77.6%,D:17.8%],F:0.7%,M:3.9%,n:303 |
| Dermatophagoides pteronyssinus| GCA_003076615.1  | 1,390             | 66,623,663            | 100.00%                     | 100.00%                  | 90.20%                      | 80,070     | C:64.7%[S:54.1%,D:10.6%],F:2.6%,M:32.7%,n:303 |
| Dermatophagoides pteronyssinus| GCF_001901225.2  | 4,324             | 68,557,481            | 94.20%                      | 77.30%                   | 25.60%                      | 74,612     | C:70.3%[S:68.6%,D:1.7%],F:3.3%,M:26.4%,n:303 |
| Dinothrombium tinctorium     | GCA_003675995.1  | 25,507            | 180,156,552           | 92.80%                      | 88.30%                   | 21.50%                      | 16,116     | C:80.5%[S:41.9%,D:38.6%],F:4.3%,M:15.2%,n:303 |
| Euroglyphus maynei           | GCA_002135145.1  | 72,786            | 43,436,854            | 34.90%                      | 13.10%                   | 0.00%                       | 787        | C:66.0%[S:65.3%,D:0.7%],F:20.5%,M:13.5%,n:303 |
| Hypochthonius rufulus        | GCA_000988845.1  | 151,357           | 171,814,378           | 39.00%                      | 25.10%                   | 1.30%                       | 3,254      | C:89.1%[S:87.1%,D:2.0%],F:5.9%,M:5.0%,n:303 |
| Leptotrombidium deliense     | GCA_003675905.2  | 17,210            | 15,149,695            | 59.20%                      | 31.50%                   | 0.00%                       | 1,422      | C:84.5%[S:83.5%,D:1.0%],F:6.3%,M:9.2%,n:303 |
| Platynothrus peltifer        | GCA_000988905.1  | 118,520           | 100,021,536           | 48.80%                      | 23.10%                   | 0.10%                       | 1,326      | C:75.2%[S:74.9%,D:0.3%],F:11.9%,M:12.9%,n:303 |
| Species                      | Accession | Length (bp) | CNTF (bp) | N50 (bp) | GC (%) | Quality (%) | C:| S:| D:| F:| M:| n: |
|-----------------------------|-----------|-------------|-----------|----------|--------|-------------|---|---|---|---|---|----|
| *Psoroptes ovis*            | GCA_002943765.1 | 93 | 63,214,126 | 100.00% | 100.00% | 86.00% | 2,279,290 | C:93.7% | S:92.7%, D:1.0% | F:2.0% | M:4.3% | n:303 |
| *Sarcoptes scabiei*         | GCA_000828355.1 | 19,246 | 56,251,741 | 58.80% | 41.90% | 6.00% | 11,383 | C:91.1% | S:91.1%, D:0.0% | F:4.0% | M:4.9% | n:303 |
| *Steganacarus magnus*       | GCA_000988885.1 | 120,241 | 112,750,608 | 46.20% | 21.60% | 0.50% | 1,727 | C:85.5% | S:84.2%, D:1.3% | F:7.9% | M:6.6% | n:303 |
| *Tetranychus articae*       | GCF_000239435.1 | 2,036 | 89,613,205 | 99.40% | 93.30% | 6.00% | 11,383 | C:91.1% | S:91.1%, D:0.0% | F:4.0% | M:4.9% | n:303 |
| *Dermatophagoides pteronyssinus* | GCA_003439945.1 | 17,171 | 959,010,206 | 99.80% | 99.50% | 0.90% | 3,060 | C:26.4% | S:26.4%, D:0.0% | F:28.1% | M:45.5% | n:303 |
| *Ixodes ricinus*            | GCA_000973045.2 | 205,231 | 514,471,516 | 99.80% | 99.50% | 0.90% | 3,060 | C:26.4% | S:26.4%, D:0.0% | F:28.1% | M:45.5% | n:303 |
| *Ixodes scapularis*         | GCF_002892825.2 | 19,746 | 3,088,623,987 | 100.00% | 100.00% | 96.90% | 517,316 | C:91.4% | S:58.1%, D:37.3% | F:0.3% | M:4.3% | n:303 |
| *Ixodes scapularis*         | GCA_000208615.1 | 570,637 | 1,388,472,180 | 99.90% | 98.20% | 3.50% | 2,942 | C:85.1% | S:83.8%, D:1.3% | F:6.6% | M:8.3% | n:303 |
| *Metaseiulus occidentalis*  | GCF_000255335.1 | 3,993 | 151,323,873 | 99.70% | 98.30% | 1.00% | 3,060 | C:85.1% | S:83.8%, D:1.3% | F:6.6% | M:8.3% | n:303 |
| *Rhodeius microplus*        | GCA_000217655.1 | 251,890 | 1,946,541,351 | 97.40% | 94.00% | 20.50% | 18,585 | C:51.9% | S:40.3%, D:11.6% | F:10.2% | M:37.9% | n:303 |
| *Tropilaelaps mercedesae*   | GCA_000208610.1 | 74,567 | 326,213,305 | 64.40% | 53.30% | 12.80% | 13,741 | C:90.4% | S:90.1%, D:0.3% | F:5.3% | M:4.3% | n:303 |
| *Varroa destructor*         | GCA_000181155.2 | 52,152 | 329,105,442 | 76.40% | 62.10% | 21.90% | 15,568 | C:92.5% | S:90.8%, D:1.7% | F:2.3% | M:5.2% | n:303 |
| *Varroa destructor*         | GCF_000243255.1 | 4,498 | 368,670,960 | 100.00% | 99.90% | 64.30% | 201,886 | C:95.7% | S:94.4%, D:1.3% | F:1.7% | M:2.6% | n:303 |
| *Varroa jacobsoni*          | GCF_002532875.1 | 8,234 | 365,177,116 | 100.00% | 99.90% | 65.10% | 96,030 | C:96.7% | S:95.4%, D:1.3% | F:1.3% | M:2.0% | n:303 |
| *Stegodyphus mimosarum*     | GCA_00061955.2 | 159,639 | 2,694,371,924 | 72.00% | 64.60% | 41.40% | 46,340 | C:88.1% | S:85.1%, D:3.0% | F:5.3% | M:6.6% | n:303 |
Table 2. KEGG pathway enrichment of 200 most highly expressed genes in fat-feeding mites

| Term                                         | Database        | ID       | Input number | Background number | P-Value      | Corrected P-Value |
|----------------------------------------------|-----------------|----------|--------------|-------------------|--------------|------------------|
| Ribosome                                     | KEGG PATHWAY    | dme03010 | 15           | 240               | 2.31E-09     | 1.04E-07         |
| Spliceosome                                  | KEGG PATHWAY    | dme03040 | 4            | 128               | 0.022180427 | 0.375099591       |
| Protein processing in endoplasmic reticulum  | KEGG PATHWAY    | dme04141 | 4            | 133               | 0.025006639 | 0.375099591       |
| Phenylalanine metabolism                     | KEGG PATHWAY    | dme00360 | 1            | 8                 | 0.070305572 | 0.489286869       |
| Arginine and proline metabolism              | KEGG PATHWAY    | dme00330 | 2            | 53                | 0.072717795 | 0.489286869       |
| Longevity regulating pathway - multiple species | KEGG PATHWAY    | dme04213 | 2            | 56                | 0.079707921 | 0.489286869       |
| Endocytosis                                  | KEGG PATHWAY    | dme04144 | 3            | 122               | 0.080418185 | 0.489286869       |
| ECM-receptor interaction                     | KEGG PATHWAY    | dme04512 | 1            | 12                | 0.0999555241| 0.489286869       |
| Fatty acid biosynthesis                      | KEGG PATHWAY    | dme00061 | 1            | 13                | 0.107219727 | 0.489286869       |
| Oxidative phosphorylation                    | KEGG PATHWAY    | dme00190 | 3            | 144               | 0.115970198 | 0.489286869       |
Comment (2) The authors showed that Arachnida species with different diet showed different evolutionary patterns based on some regular analyses. These results are expectable. It is not clear what is the key scientific problem the authors want to solve based on their analyses. What is the purpose of doing these analyses as well as publishing a paper, and how these results will serve the purpose? In the abstract and the conclusions, the authors mentioned twice: “These different genetic bases provide a new perspective for the study of the evolution and diversification of this group, and offers potential drug targets for pest control.” These statements are repetitive, and are very vague. What is the “new perspective” and how to offer potential drug targets? Details should be discussed at least. In addition, not all mite species analyzed in this study are pests. For example, Metaseiulus occidentalis is a predatory mite natural enemy.

Thank you for your constructive advice, we have added a description of the motivation of our research in the Abstract and Introduction section to help readers quickly understand the main motivation of our research. Here is a short summary “Diet is one of the most fundamental aspects of an animal’s biology and is a powerful evolutionary force for species adaptation and diversification. Acari (mites and ticks) is one of the most abundant clades of Arachnida, exhibiting extraordinarily diverse dietary types. While studies have focused on morphological and physiological adaptations to different dietary habits, the genetic mechanisms underlying these adaptations are not fully understood. Based on a comparative analysis of 15 Acari genomes and five dietary habits, we found different genetic bases for different diets, mainly related to the need to handle different food types, including increased abilities to find, prepare and digest food.”. Details are in the Introduction section.

Sorry for the confusing expression. We have changed the vague sentence “Based on comparative analyses of 15 Acari genomes, we found genetic bases for three specialized diets. Herbivores experienced stronger selection pressure than other groups; the olfactory genes and gene families involving metabolizing toxins showed strong adaptive signals. Genes and gene families related to anticoagulation, detoxification, and haemoglobin digestion were found to be under strong selection pressure or significantly expanded in the blood-feeding species. Lipid metabolism genes have a faster evolutionary rate and been subjected to greater selection pressures in fat-feeding species; one positively selected site in the fatty-acid amide hydrolases 2 gene was identified. Our research
provides a new perspective for the evolution of Acari and offers potential target loci for novel pesticide development.”. More information has been discussed in the specific dietary section in the revised manuscript.

Our study of evolutionary adaptation for different diets was not only based on pests but based on the mites with specialized dietary styles. However, the genetic adaptation of three specialized diets was implied and could consequently help with inhabitation of some pests such as spider mites, honey bee mites and so on. We could take efficient measures to weak the abilities for finding (olfaction), preparing (detoxification) and digesting (metabolism), which are found in the current study. For example, we have detected positive selection sites in HSP genes involving in olfactory pathways, which offered several drug targets for control the spider mites. Similarly, we have found that the evolutionary speed of lipid metabolism is significantly accelerated in honey bee mites, especially of arachidonic acid lipid metabolism pathway (Figure 6a and 6c). We could design drugs targeting at the candidate genes of arachidonic acid lipid metabolism such as FAAH2 to inhibit or kill honey bee mites in the following research.

Comment (3) There are many typos and formatting errors in manuscript (including the references). I think the authors should pay more attentions to fix these problems. Sorry for these mistakes. We have carefully modified our manuscript. And the revised manuscript was edited for proper English language, grammar, punctuation, spelling, and overall style by one or more of the highly qualified native English speaking editors at SNAS (verification code 30F9-7501-1FD4-89B3-2576).

Finally, thank you again for your great suggestions. We hope our manuscript can be of great interest to researchers of related fields now.

Best Regards.

Sincerely,
Dr. De Chen on behalf of all authors
REVIEWERS' COMMENTS:

Reviewer #2 (Remarks to the Author):

Thanks authors' Responses. I have no further comments. I am looking forward to the manuscript published on communication biology.

Reviewer #3 (Remarks to the Author):

I think this manuscript has been well updated, and can be published.