

# Evolutionary innovation in the long-term absence of sex in the oribatid mite *Platynothrus peltifer*

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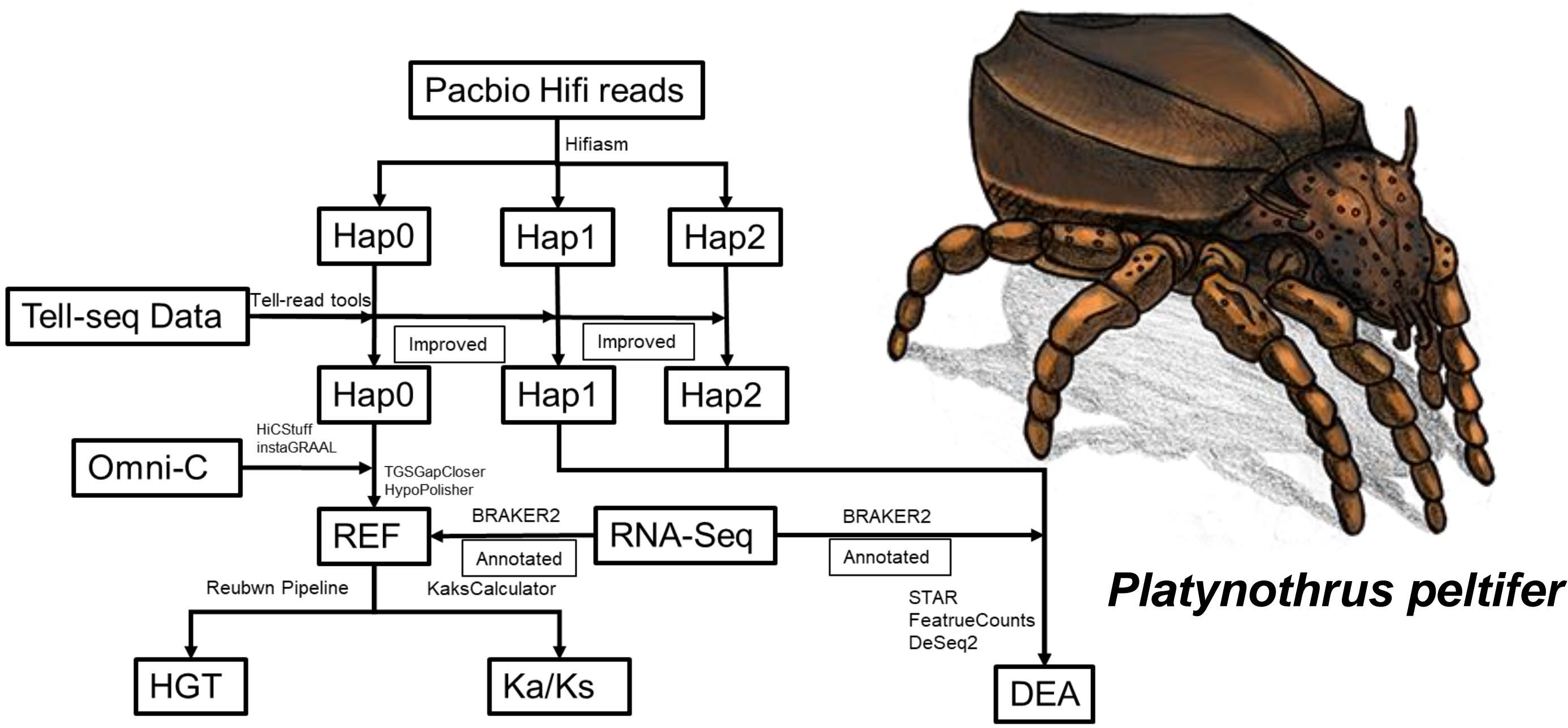
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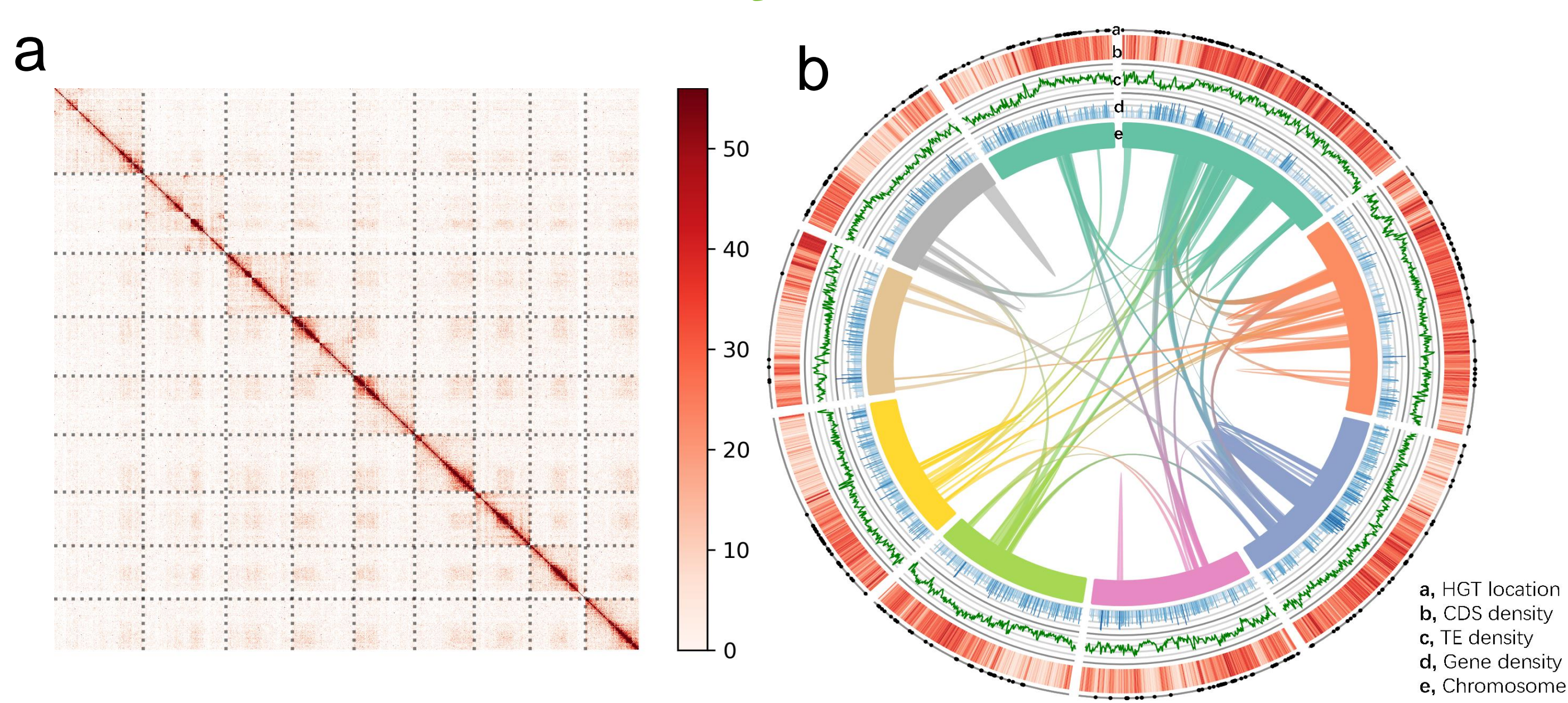
## Introduce

Asexuality is considered an evolutionary dead-end, but some ‘**ancient asexual organisms**’, like oribatid mites, have persisted and diversified over time. Very little is known about the processes that could generate evolutionary novelty and adaptability in the absence of sex. We analyzed potential genomic footprints of innovation in the haplotype-resolved, chromosome-scale genome of the ancient asexual oribatid mite *Platynothrus peltifer*. Large-scale structural variants might not play a major role, as there are few rearrangements within and between chromosomes as well as between haplotypes. However, heterozygosity is maintained and divergent haplotypes, as well as **differential allele expression (DEA)** with an functional enrichment in e.g. metabolic pathways, resource uptake and immune system. We further analyzed the evolutionary trajectories of these alleles, as these processes might contribute to adaptation and evolutionary novelty in the absence of sex. Moreover, **Horizontal Gene Transfer (HGT)** could supply the substrate for novelty via modulating gene regulation of these mites. Overall, identifying such signatures of evolutionary innovation will help to understand why some asexual can escape the dead-end fate and to identify the benefits of sex vice-versa.

## Workflow



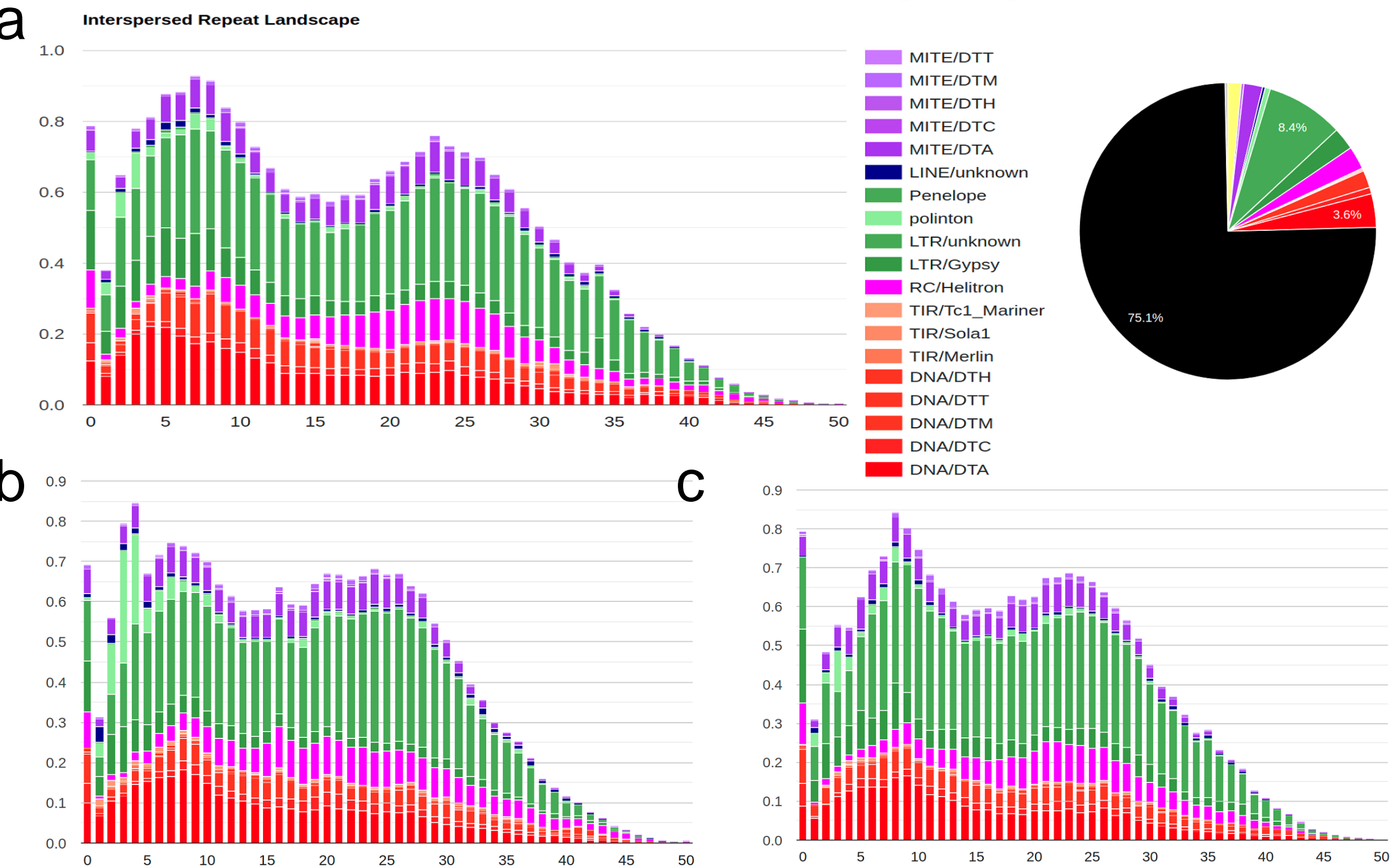
## 1. Genome assembly



Features	Hap0	Hap1	Hap2
total length [Mb]	220	208	200
N50 [Mb]	23	14	14
Transposable elements	52Mb(25%)	48Mb(23%)	45Mb(23%)
BUSCO [%] (assembly)	96.7	96.5	96.3
BUSCO [%] (assembly)	97.6	97.9	97.3
Nr annotated genes	24139	24357	22553

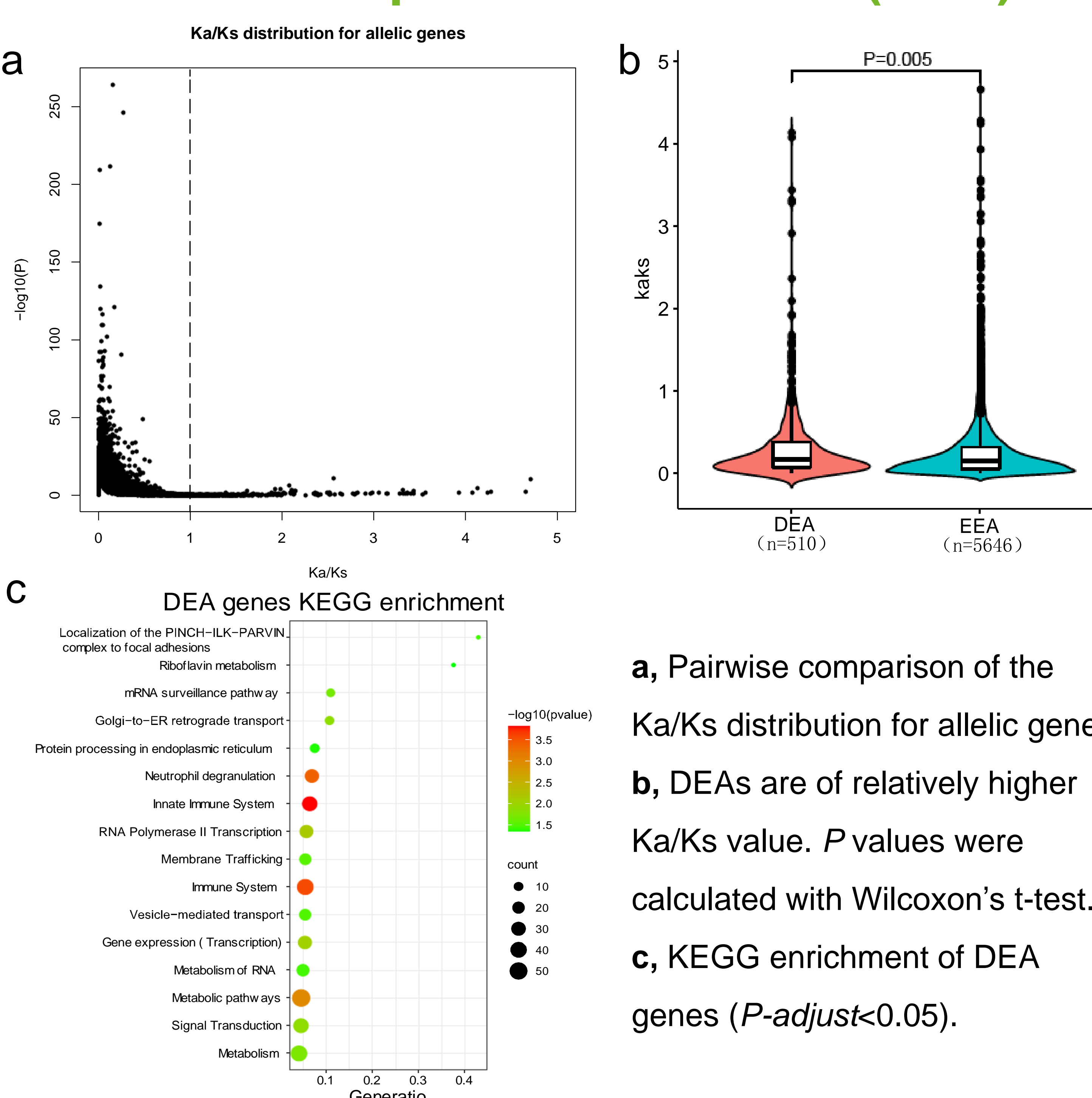
a, Contact map of Hi-C links among 9 chromosomes. b, Rearrangement of the *Platynothrus peltifer* and genomic features.

## 4. Transposable element(TE) landscape



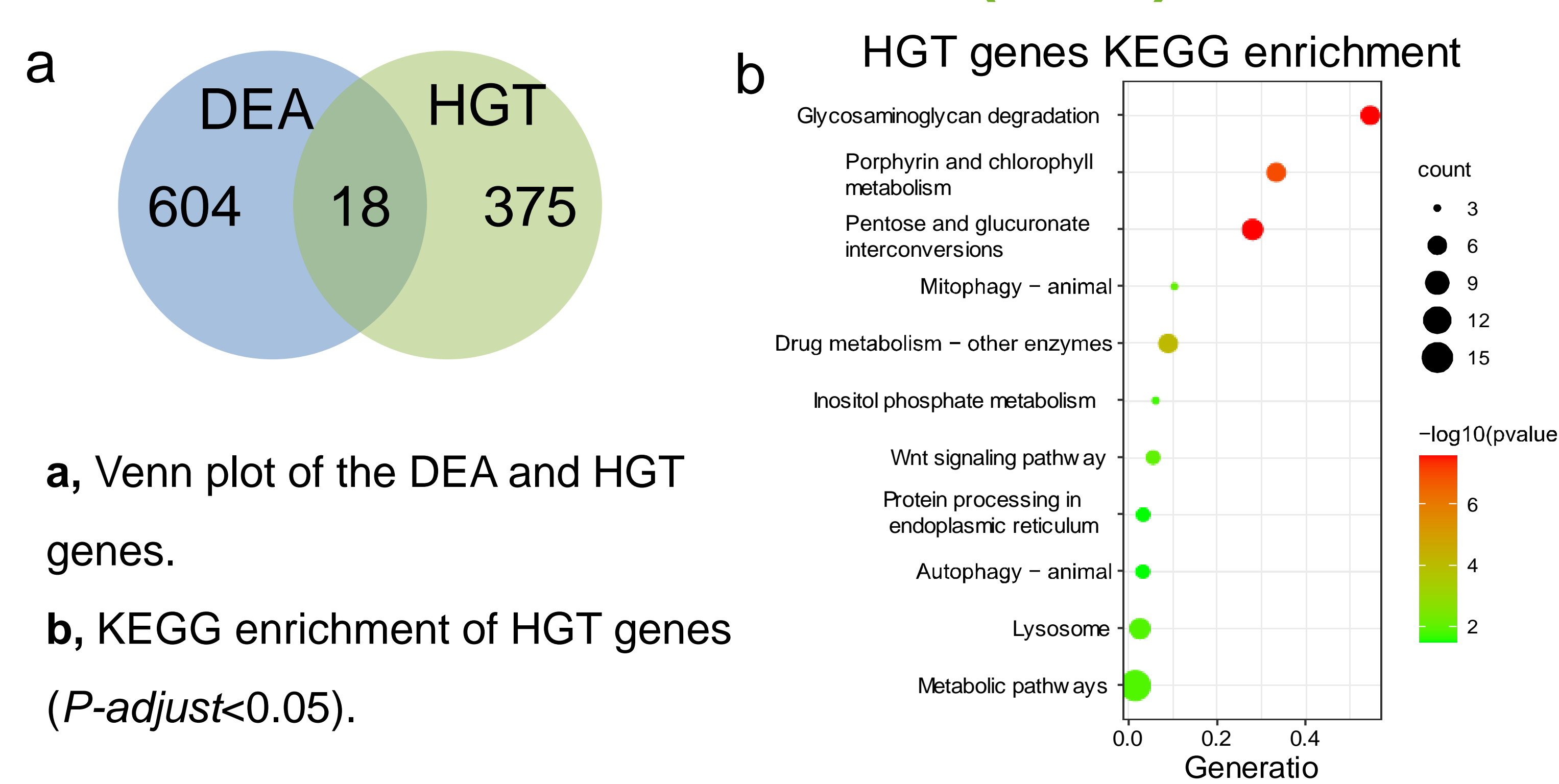
a, b, c are Transposable element(TE) landscapes of Hap0, Hap1 and Hap2

## 2. Differential expression of alleles(DEA)



a, Pairwise comparison of the Ka/Ks distribution for allelic genes. b, DEAs are of relatively higher Ka/Ks value. *P* values were calculated with Wilcoxon's t-test. c, KEGG enrichment of DEA genes (*P-adjust*<0.05).

## 3. Horizontal Gene Transfer (HGT)



a, Venn plot of the DEA and HGT genes. b, KEGG enrichment of HGT genes (*P-adjust*<0.05).

## 5. Conclusion and outlook

- Single individual chromosome-level scale reference genome and two high quality haplotypes assemblies.
- $K_a/K_s$  of DEA relatively higher EEA genes.
- 10% single copy genes between two haplotypes show DEA
- 77.5% HGT with intron and 1.8% HGT show DEA

Outlook: construct Pan-genome of the *Platynothrus peltifer*

