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A Bioinformatics Approach for Evaluating Evolutionary Convergence of Gene Family Size in Hematophagous Insects

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Abstract

The act of blood-feeding can be nutritionally rewarding for blood-feeding arthropods. However, blood digestion can release pro-oxidant molecules such as heme and iron at potentially harmful levels. If left uncontrolled, this heme/iron can cause oxidative damage and eventually cell death. This has led to the evolution of various adaptations that protect blood-feeding arthropods against iron- and heme-associated damage. Here I postulate that the signature of this adaptation can be observed in patterns of gene family size. To test this hypothesis, I explore convergent evolutionary expansions and contractions of gene families in distinct lineages of hematophagous insects. Specifically, I compare the gene content present in available genomes from bloodfeeding and non-blood feeding arthropods (including outgroup taxa in the Lepidoptera [moths & butterflies]), to identify possible changes in gene family size in the blood-feeding taxa. Of the 206 heme/iron-associated genes identified from the model insect, Drosophila melanogaster, five were overrepresented (potentially duplicated) in the blood-feeding taxa: spook (cyp307A1), spookier (cyp307A2), cytochrome P450 12e1 (cyp12e1), hormone receptor and 51 (Hr51), NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6), and seven were underrepresented (potentially lost). However, when only Dipteran (fly) and Siphonaptera (flea) genomes were included in the analysis, just one iron gene and one heme gene (NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW) were overrepresented in the bloodfeeding taxa. Interestingly, the expanded cytochrome genes are known detoxifiers of many compounds, including heme and iron. More broadly, the analytical approach I employee here could be used to evaluate functional convergence for other phenotypic traits, conditional on the availability of annotated genomic data

Keywords: evolutionary convergence, hematophagy, Diptera, Lepidoptera, Siphonaptera, gene ontology, gene family expansions, cytochrome P450, NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW)

MONTCLAIR STATE UNIVERSITY

A Bioinformatics Approach for Evaluating Evolutionary Convergence of Gene Family size

Expansions in Hematophagous Insects

by

Mbemba Ceesay

A Master's Thesis Submitted to the Faculty of

Montclair State University

In Partial Fulfillment of the Requirements

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A BIOINFORMATICS APPROACH FOR EVALUATING EVOLUTIONARY CONVERGENCE OF GENE FAMILY SIZE IN HEMATOPHAGOUS INSECTS

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1. Introduction

1.1 Evolutionary Convergence

Evolutionary convergence is the independent evolution of related traits in distinct taxa. Organisms with varied phylogenetic evolutionary histories may nonetheless exhibit similar phenotypes due to comparable selective pressures (Losos 2011; Wheeler and Carstens 2018). These selective pressures lead to the evolution of the same adaptive syndromes that allow the performance of a particular task or the adaptation to a new lifestyle (Reich et al. 2003). The independent evolution of eyes in octopi, vertebrates, and spiders is a classic example of convergent evolution. Additionally, flight in insects, birds, and bats also attests to the same phenomenon (Xu et al. 2020; Losos 2011).

Convergence observations may support a hypothesis that natural selection was a driving mechanism of specific traits (Stern 2013, Wheeler and Carstens 2018). However, evolutionary convergence is not limited to adaptation due to similar selective conditions related to natural selection (Losos 2011). Species can also evolve similar phenotypes due to coincidences unrelated to natural selection, thus excluding the occurrence of a particular trait for selective environmental reasons. Secondly, apparent convergence could be due to exaptation. In this case, a change in the function of a trait during evolution may occur if a species adopts a beneficial feature to a particular environment for a different reason, resulting in species sharing similar phenotypes in comparable selective environments but which evolved as an adaptation to different selective pressures (Losos 2011). Finally, constraints that bias the number of possible changes could push distantly related species to evolve convergent traits through nonadaptive processes (Losos 2011; Sackton and Clark 2019). For instance, the evolution of small size might be due to

a particular environment through paedomorphosis and consequently, give rise to additional paedomorphic features such as fewer digit numbers or webbed feet (Losos 2011). Salamanders are an example of this phenomenon. The number of cells in the salamander's limb bud partially determines the number of digits on the limb. Accordingly, evolutionary changes corresponding to a fewer number of cells can lead to decreased body size resulting in digit numbers reducing from five to four (Losos 2011; Alberch and Gale 1985).

1.2 Measuring Convergent Evolution

Convergence at the genetic level frequently occurs due to similar genetic modifications. It does so through parallel and collateral genetic evolution. Parallel genetic evolution is the independent evolutionary occurrence of mutations in different species or populations. In contrast, collateral genetic evolution occurs firstly through the evolution of polymorphic alleles within populations with shared ancestry and secondly through introgression, a process in which an allele from one population is introduced into another population through hybridization (Stern 2013). Genetic approaches for examining convergent evolution tend to focus on convergent patterns in biological pathways, gene expression, gene copy number, transposable element content (TE), amino acid (AA) substitutions in specific genes, genome-wide AA substitutions, and/or the percentage of guanines and cytosines (GC content) (Xu et al. 2017).

An example of parallel evolution between populations within the same species: The mouse-ear cress (*Arabidopsis thaliana*), which, during the spring, goes through cold temperature exposure, a requirement to induce flowering in a process called vernalization. However, many populations amongst this species can flower without exposure to cold temperatures and thus forego vernalization. This subpopulation of *Arabidopsis thaliana* has 20 instances of independent

mutations that deactivate the gene responsible for vernalization, namely the FRIGIDA gene. Consequently, 70% of the noticed differences in flowering times are attributable to changes in the FRIGIDA gene (Shindo et al. 2005; Stern 2013).

Mullerian mimicry complexes in *Heliconius* butterflies exemplify collateral evolution through hybridization. In Mullerian mimicry, *Heliconius* butterflies have evolved vibrant wing color patterns of red, orange, black, and white. These color combinations warn predators of the association between the manifested color combinations and unpalatability. (Heliconius Genome Consortium 2012; Stern 2013). Evidence suggests that gene duplication and reductions could arise convergently with far-reaching consequences. Pathways of genes essential to frigid responses and shade avoidance convergently retain duplicated gene copy numbers in various plant lineages. Possessing additional copies of these genes enhances plant survival in cold environments devoid of access to light (Wu et al. 2019). Conversely, convergent reductions in gene copy numbers could attenuate certain features, as is the case for seagrasses that have independently returned to an aquatic environment, consequently losing features such as complex floral structures and stomata (Kuo and Hartog 2007; Xu et al. 2020) Gene duplication and variations in gene copy numbers actively participate in adaptations to environmental challenges (Xu et al., 2017).

1.3 Blood Feeders

Estimates suggests there may be more than 1 million contemporary insect and arachnid species on Earth. Of these, ~14,000 have the capability to feed on vertebrate blood (Graça-Souza et al. 2006). For instance, blood-feeding ticks can ingest a volume of blood that is 100 times their premeal weight. (Reuben Kaufman 2007). An ability to feed on blood, generally known as

hematophagy, has occurred independently several times during insect evolution, (Graça-Souza et al. 2006), especially among dipterans (flies) which make up the most abundant and diverse order of blood feeders (Dashti et al. 2016). The evolution of blood-feeding may have arisen from an ability to feed on the skin or wastes of dead animals, and/or through pre-adapted piercing mouthparts that then allowed these species to penetrate animal skin to access blood (Graça-Souza et al. 2006).

Hematophagous insects rely on blood-feeding to acquire vital nutrients for development and reproduction (Dalton et al. 2004). While rewarding, the blood-feeding process can be dangerous for hematophagous invertebrates as vertebrates have defensive tendencies against blood-feeders. Accordingly, limiting contact with vertebrates is an essential survival trait for blood-feeders and forces them to ingest large amounts of blood in a single meal (Sterkel et al. 2017). This may expose the blood feeder to high concentrations of specific blood constituents. The chemical composition of dry vertebrate blood is almost 90% proteins with hemoglobin being the most abundant protein (~150mg/ml), then albumin (~50mg/ml) (Sterkel et al. 2017). Bloodfeeding insects generally ingest 2 to 100 times their body weight during a single blood meal, and the rich protein meal digestion releases amino acids essential for egg protein, carbohydrate, and lipid synthesis (Whiten et al. 2017). Compared to proteins, carbohydrates, for instance, only account for 0.4% of dry vertebrate blood composition (Sterkel et al. 2017). Although nutritionally beneficial, the blood-feeding process releases heme and iron at potentially harmful levels, requiring hematophagous arthropods to evolve means to limit oxidative damage during digestion (Whiten et al. 2017).

Interestingly, the lamprey demonstrates convergent evolution due to possessing five unique globin genes (Adgb, Gbx, Cygb, Hbs, and Mbs) that convergently evolved from an

identical globin predecessor (Schwarze 2014). Vertebrate hemoglobin comprises two identical alpha subunits (141 residues) and two identical beta subunits (146 residues), each possessing a single heme group (Perutz 1970). The interactive hetero-tetrameric hemoglobin, the most abundant protein in the blood, promotes the transport of O₂ in the circulatory system (Brunori 1999; Rohlfing et.al 2016). During the formation of the heme-protein (hemoglobin), its cytotoxic and unstable constituents, free alpha and beta subunits, free-heme, and uncomplexed iron ions, promote free radical aggregation, thus requiring coordination to reduce harmful effects (Baker et al. 2003). In addition to oxygen transport, the globin proteins, including hemoglobin (Hgb), myoglobin (Mgb), neuroglobin (Ngb), cytoglobin (Cygb), globin's E, X, Y, and androglobin (Adgb), are known detoxifiers of reactive oxygen and nitrogen species. They do so through their conserved heme prosthetic group fold that binds reversibly to O₂ (Rohlfing et al. 2016).

Heme is composed of an iron atom bonded to four nitrogen atoms belonging to the highly reactive pyrrole ring of protoporphyrin IX (Toh et al. 2010). Metabolic heme and iron are sourced from dietary heme in some hematophagous parasites. Heme binds directly to important molecules by transitory interactions, which generate free radicals. Due to its destabilizing effects, heme promotes cell lysis as it binds to macromolecules. By binding to enzymes and transcriptional factors, heme modulates the transcription and translation of protein synthesis, affecting the regulation of cell development, apoptosis, and signal transduction (Hou et al. 2007; Toh et al. 2010). Heme generates hydroxyl radicals and reactive oxygen species due to the presence of iron via the Fenton reaction or the heme-H₂O₂ reaction. The presence of ROS degrades and destroys proteins, lipids, and DNA molecules. Yet, low amounts of ROS are vital for critical biological functions such as the up-regulation of heme synthesis, signal transduction, and enzyme activation in cells (Tsiftsoglou et al. 2016; Toh et al. 2010).

The breakdown of hemoglobin releases large amounts of free heme and free iron. Under favorable conditions, these compounds enable damaging reactions (Everse and Hsia 1997). Although iron is essential for biological redox reactions, the easy conversion between Fe(II) and Fe(III) enhances iron toxicity by producing oxygen-derived radicals and other damaging molecules (Pierre et al. 2002). Knowledge of iron metabolism in humans is extensive compared to insects and less still for blood-feeding insects. Iron metabolism in humans requires multiple functional genes such as iron regulatory proteins, transferrin receptors, transferrin, ferritin, and many more genes, with the majority absent of human homologs in insects (Dashti et al. 2016). Understanding iron metabolism can explain why hematophagous arthropods survive and flourish, given their potentially harmful lifestyle (Galay et al. 2015).

Hematophagous arthropods, over time, have developed immune antioxidant responses to tackle the customary oxidative stress post-blood-feeding. As the first line of defense against heme toxicity after a blood meal, hematophagous insects generate insoluble heme products for excretion with wastes, thus quantitatively expelling most of the heme generated from the blood meal (Sterkel et al. 2017). Additional responses are protein, lipid, and nucleic acid oxidation. Intracellular antioxidants such as Cu, Zn, and Mn superoxide dismutase (SOD) and extracellular antioxidants such as the heme and iron chelator xanthurenic acid relieve oxidative stress after a blood meal (Lima et al. 2012; Whiten et al. 2017) Blood-feeding parasites, for instance, utilize enhanced hemolytic and proteolytic enzymes to lyse and catabolize red blood cells, blood's most significant cellular component (Brindley 1998). They have also developed defenses in the gut lumen, cytoplasm, and midgut epithelial cells (Whiten et al. 2017). Most of the iron absorbed from the blood meal is sourced from heme iron. However, mosquitoes, for instance, prevent iron overload by excreting excess iron, blocking iron absorption, storing absorbed iron in ferritin, and

circulating iron to eggs and other tissues (Zhou et al. 2007). Parasite survival, symbiotic fitness, and successful reproduction all require rigid control of iron metabolism (Dashti et al. 2016).

1.4 Genome Evolution and Gene Ontology (GO) in Blood-Feeders

Examining genome sequences of blood-feeding insects may allow a better understanding of the physiological adaptations undergone by hematophagous insects. In the case of disease vector insects, genomic analyses may also allow the identification of novel targets in disease transmission. Additionally, comparing genomic data of hematophagous insects and their nonblood-feeding relatives can offer convincing evidence regarding the physiological consequences of blood-feeding (Dashti et al. 2016).

Using genomic data, it can be challenging to infer a common molecular role based on nucleotide similarity, as orthologous proteins may have very different functions, while evolutionary divergent proteins may be important in the production of similar phenotypic effects (Doolittle 1994). If defined based on function rather than nucleotide sequences, evolutionary syndromes could be a possible solution to the mentioned challenge because they allow genetic convergence to be defined by function. With this approach, convergence can now be determined by functional similarity (Wheeler and Carstens 2018).

The use of gene ontology (GO) coding can effectively measure functional convergence because it allows the pairing of distinct genes to a specific numeric code relying on available experimental data that has determined sequence similarity to gene function (Wheeler and Carstens 2018). GO terms are assigned numerical codes to potential biological activities and ranked based on specificity. Consequently, search terms that are synonymous with a specific function can be used to measure evolutionary convergence between different species.

Furthermore, GO codes allow the selection of relevant functions before comparing genes between species with similar convergent functions (Wheeler and Carstens 2018).

Gene family expansions in insects may be linked to hematophagy, as the expression of additional copies of genes associated with blood-feeding could assist in blood digestion (Freitas and Nery 2020). The evidence gathered from the literature suggests that the blood feast generates toxic amounts of heme and iron, which require proper and timely disposal to avoid harming host cells, proteins, and DNA. This study aims to identify evolutionary convergence within potentially associated gene families that may have expanded during the evolution of the bloodfeeding process. Therefore, in this study, I predict that gene families related to the detoxification of heme/iron and implicated in the blood-feeding process may have expanded.

2. Materials & Methods

Arthropods that feed on vertebrate blood poses the need to detoxify the excess heme and iron present in the excessive blood feast. The need to detoxify heme and iron served as the basis for this study.

2.1 Species list

I generated a varied species list (Table 1) consisting of blood-feeding Diptera (flies) and Siphonaptera (fleas) representing six independent evolutionary lineages of hematophagy and a comparable number of non-blood-feeding Diptera (Figure 1). I also included several species of non-blood-feeding Lepidoptera (an outgroup sister Order to the Diptera/Siphonaptera clade) as a reference group to partially control for evolutionary relatedness within the Diptera and Siphonaptera. Annotated genomes of the dipterans were downloaded from <u>https://www.ncbi.nlm.nih.gov/assembly/?term=diptera</u>, while the lepidopteran genomes were downloaded from <u>https://www.ncbi.nlm.nih.gov/assembly/?term=lepidoptera</u>. The non-blood-feeding Diptera and Lepidoptera species served as additional controls to account for phylogenetic effects, as closely related species may resemble each other compared to species selected randomly.

2.2 Step 1: Get Genes Associated with Iron and Heme

For my study, the model organism *Drosophila melanogaster* served as the reference. This was done because gene functions are well characterized in this species. I used the custom Perl script "Select_word-associated_GO_terms.pl" to locate any GO term that contains the words' heme' or 'iron' from the *D. melanogaster* go-basic.obo database downloaded from http://purl.obolibrary.org/obo/go/go-basic.obo to obtain the *D. melanogaster* associated heme/iron genes. These terms were than manually examined to assess their potential association with heme and/or iron molecular processes in insects.

2.3 Step 2: Make Query File Once Per Search Term

Once extracted, I created a query file (a searchable sequence against the blast database) for all the *D. melanogaster*-related heme and iron genes using the heme and iron GO terms generated from the previous step (go_terms_heme.txt and go_terms_iron.txt). The Perl script, "perl get_drosophila_exons_with_GO_term_list.pl" executed with the command line using the command: (perl get_drosophila_exons_with_GO_term_list.pl iron_go_terms.txt gene_association_v2.1.fb all_dmel_exons.fa > dros_iron_query_seqs.fa) and (perl get_drosophila_exons_with_GO_term_list.pl heme_go_terms.txt gene_association_v2.1.fb

all_dmel_exons.fa > dros_heme_query_seqs.fa) created the query sequences file, once for each search term. This step associated the heme and iron genes from the previous step with their reference *D. melanogaster* genes. I used the program samtools-1.14

https://www.htslib.org/download/) to process the sequence alignments between the heme and iron genes and the reference *D. melanogaster* genes within the Perl script. This step gathered all of the iron and heme genes that match their orthologs in the *D. melanogaster* exons as query files (dros_heme_query_seqs.fa, dros_iron_query_seqs.fa), for subsequent use with the Basic Local Alignment Search Tool ('BLAST).

2.4 Step 3a: Make Blast Database (Once per Species)

BLAST aligns and compares a query DNA sequence with a sequence database. The matches generated from BLAST are ranked based on statistical significance (Altschul et al. 1990). Because BLAST requires a query sequence and a database to search against, the query sequences obtained from step 2 were searched against the databases generated in step 3a. Blast version: ncbi-blast-2.12.0+ was downloaded from

https://blast.ncbi.nlm.nih.gov/Blast.cgi?PAGE_TYPE=BlastDocs&DOC_TYPE=Download. Once downloaded, ncbi-blast-2.12.0+ created databases for each species with the command line using, the command: ~/bin/ncbi-blast-2.12.0+/bin/makeblastdb -in

GCA_020740445.1_IBAB_Cspp_1.0_genomic.fna -dbtype nucl. Each database created identifies with the appropriate species accession number. Step 3a generated databases for each species' genome, the second component required to conduct a blast search.

2.5 Step 3b: TBLASTN against each species' genome sequence, once per species

The BLAST tool 'tblastn' aligns protein sequences (query) to a translated nucleotide database (Gertz et al. 2006). Using tblastn, I searched amino acid sequences from D. melanogaster against the sequences of each taxon. The availability of species databases and search term query sequences from the previous steps made the blast searches possible. I searched D. *melanogaster* genes related to heme and iron query sequences against the genomic database of each species with the command line using the command: ~/bin/ncbi-blast-2.12.0+/bin/tblastn query dros iron query seqs.fa -db GCA 020740445.1 IBAB Cspp 1.0 genomic.fna -outfmt 6 -evalue 1e-10> blast out iron culicoides-tainanus.txt. And for heme, the command: ~/bin/ncbiblast-2.12.0+/bin/tblastn -query dros heme query seqs.fa -db GCA 020740445.1 IBAB Cspp 1.0 genomic.fna -outfmt 6 -evalue 1e-10 >blast out heme culicoides-tainanus.txt. The parameters are set to format tblastn outputs in a tabular format (-outfmt 6) and to filter out statistically significant tblastn matches (<1e-10). Individual blast outputs for each species (Diptera, Siphonaptera, and Lepidoptera) were generated from the above commands, highlighting the total number of D. melanogaster-related heme and iron genes and their total gene copy numbers. I assembled each species' blast results in a single tab-delimited excel file for further processing.

2.6 Step 4: Calculate the average number of blast hits per gene, once per species

The BLAST output with the *D. melanogaster-related* iron and heme genes and their copy numbers was further processed to obtain the average number of blast hits per gene (average gene copy numbers) using the Perl script "perl calculate_average_gene_blast_hit.pl". The Perl script, executed with the command line using the command: perl calculate_average_gene_blast_hit.pl blast_out_culicoides-tainanus.txt. 50 40 gene_list.txt > average_irongene_culicoides-

tainanus.txt. And perl calculate_average_gene_blast_hit.pl blast_out_culicoides-tainanus.txt. 50 40 gene_list.txt > average_hemegene_culicoides-tainanus.txt. I defined the blast search criteria as 50% minimum sequence similarity and 40 base pairs for sequence length to capture *D*. *melanogaster-related* heme and iron genes with 50% sequence similarity and at leaset 40 base pairs in length. This step gathered all the iron and heme genes within the study taxa genomes that satisfied these criteria.

2.7 Statistical Analysis

I assessed statically significant differences in gene family expansions or contractions between the blood-feeders Dipterans, non-blood-feeding dipterans, and the non-blood-feeding Lepidopterans. T-tests with the Perl script "calculate_t_test.pl" were conducted relying on the differences in "phenotypes," namely "Blood" and "No," to distinguish the blood-feeders from the non-blood-feeders. P-values below 0.05 were considered statically significant and representative of gene family expansions or contractions within the Diptera Siphonaptera and Lepidoptera taxa.

Additional analysis excluded the outgroup Lepidopterans and assessed gene family convergent expansions and contractions in the Dipteran blood and non-blood-feeders and the Siphonaptera blood-feeder. This analysis generated results that can be compared to the previous analysis that accounted for both the ingroups (Diptera & Siphonaptera) and the outgroup (Lepidoptera) to assess convergent gene family expansions and contractions within the ingroup. T-tests with the Perl script "calculate_t_test.pl" were conducted relying on the differences in "phenotypes," namely "Blood" and "No," to distinguish the blood-feeders from the non-bloodfeeders. P-values below 0.05 were considered statically significant and representative of gene family expansions or contractions within the Diptera taxa. I examined convergence using the average gene family copies of the blood-feeders and the non-blood feeders with p-values below 0.05. The selection of gene families based on appropriate p-value scores allows comparisons between average gene family copy numbers of the blood-feeding and non-blood-feeding taxa. The group with the higher average gene copy number is considered to have expanded or contracted convergently due to the blood-feeding lifestyle or the absence of blood-feeding.

3. Results

This study identified four hundred sixty-seven iron-associated GO terms and eighty-five hemeassociated GO terms. An assessment to identify all heme and iron genes in all species and their copy numbers yielded 206 annotated genes in *D. melanogaster* that have possible relations to iron and heme. The gene copy numbers varied amongst the species and ranged from 0 to 218 copies.

3.1 Iron gene family Expansions and Contractions in the Dipteran, Siphonapteran and Lepidopteran Taxa

Regarding gene family expansions and contractions in the Dipteran, Siphonaptera and Lepidopteran taxa, a t-test (Table 2) comparison of hematophagous Diptera vs. nonhematophagous (Diptera & Lepidoptera) taxa revealed five iron gene families (p < 0.05) that were expanded in the hematophagous taxa and seven (p > 0.05) that contracted in the hematophagous taxa. The iron gene families with statically significant p-values favoring the hematophagous taxa and indicating gene family expansions (Table 3) are spook (Spo) (p =0.01233), Cytochrome P450 12e1 (Cyp12e1) (q = 0.01364), hormone receptor 51 (Hr51) (p =

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0.0205), spookier (Spok) (p = 0.02546), and the NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6) (p = 0.03154).

The same t-test (Table 3) highlighted cardinal (Cd) (p = 0.003787), Curly su (Cysu) (p = 0.008056), Heavy metal tolerance factor 1(Hmt-1) (p = 0.01853), Peroxinectin-like (Pxt) (p = 0.03137), NADH dehydrogenase (ubiquinone) 13 kDa A subunit (ND-13A) (p = 0.03277), Nitric oxide synthase (Nos) (p = 0.04328), and Cytochrome P450 4d1 (Cyp 4d1) (p = 0.04479) as gene families that favored the non-hematophagous taxa thus indicative of contraction.

3.2 Heme gene family Expansions and Contractions in the Dipteran, Siphonapteran and Lepidopteran Taxa

The t-test analysis for heme gene family expansions and contractions in blood-feeding (Diptera and Siphonaptera) and non-blood-feeding (Diptera and Lepidoptera) yielded (table 4) spook (p=0.01233), Cytochrome P450 12e1 (p=0.01364), hormone receptor 51 (p=0.0205), spookier(p=0.02546), and the NADH dehydrogenase (ubiquinone) B16.6 subunit(p=0.03154) as expanded (Table 5).

The same t-test found cardinal (p=0.003787), Curly su (p=0.008056), Heavy metal tolerance factor 1(q=0.01853), Peroxinectin-like (q=0.03137), NADH dehydrogenase (ubiquinone) 13 kDa A subunit(q=0.03277), Nitric oxide synthase(q=0.04328), and Cytochrome P450 4d1 (q=0.04479) as heme gene families that contracted (Table 5).

3.3 Iron and Heme Gene Family Expansions in Diptera and Siphonaptera

To further investigate the iron related gene family contractions and expansions in Dipteran and Siphonaptera hematophagous, and non-hematophagous Dipteran taxa, a t-test was conducted,

(Table 6) and it yielded ND-PDSW (q = 0.04776) as the only gene family that expanded (Table 7). The remaining six gene families namely, Cyp4d1(q = 0.008542), Cyp6a16 (q=0.01446), Cyp6a17 (q = 0.03699), Cyp6a23 (q = 0.03714), ND-B14.5A (q = 0.04678), and Cyp6t3 (q = 0.04681) contracted.

Averages copies of each gene family were noted for the hematophagous and nonhematophagous Diptera taxa (table 7), and these averages were used to establish gene family contractions and expansions.

In this analysis (Table 8), a t-test comparison of hematophagous Diptera and Siphonaptera and non-hematophagous Diptera taxa as a group, ND-PDSW (q = 0.04776), was noted as the only heme gene family that expanded (Table 9). The remaining six gene families namely, Cyp4d1(q = 0.008561), Cyp6a16 (q = 0.01446), Cyp6a17 (q = 0.03699), Cyp6a23 (q = 0.03714), ND-B14.5A (q = 0.04678), and Cyp6t3 (q = 0.04681) contracted (Table 9).

4. Discussion

This study was designed to identify convergent evolutionary expansions within gene families that may have occurred during the evolution of hematophagy in dipterans. Accordingly, it was hypothesized that genes implicated in heme/iron detoxification during blood digestion may have expanded to limit oxidative damage during digestion, and additional copies of genes associated with hematophagy could assist in blood digestion.

Notably, in the analysis of heme and iron-related gene family expansions in the Diptera, Siphonaptera and Lepidopteran taxa, the results identified potential expansions in the following protein-coding genes: spook (cyp307A1), spookier (cyp307A2), cytochrome P450 12e1 (cyp12e1), hormone receptor and 51 (Hr51), NADH dehydrogenase (ubiquinone) B16.6 subunit

(ND-B16.6). When assessing iron and heme gene family expansions in the Diptera and Siphonaptera taxa alone, NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW) expanded.

The spook and spookier genes code for cytochrome P450 enzymes and are part of the more prominent cytochrome P450 gene family. Spook precisely retains distinct sequences found in P450 proteins, including the heme-binding domain (PFxxGXRxCxG), and it also plays a role in insect molting (Ono et al. 2006). The presence of the conserved heme-binding domain suggests a possible role of spook and other cytochrome P450s in detoxifying heme. Additionally, cytochrome enzymes are known detoxifiers of endogenous and exogenous compounds through oxidation reactions (Li et al. 2007). Experimental data are needed to verify the role of cytochrome P450 proteins in detoxifying excess heme after a blood meal in the taxa represented in this study. However, it is reasonable to suppose a detoxification role since these proteins are known to carry out a diverse and often surprising number of oxidation reactions to expunge many compounds (Werck-Reichhart and Feyereisen 2000). Furthermore, due to the presence of the conserved heme-binding domain, it possibly expels excess heme after a blood meal in the hematophagous taxa.

The expansion and diversity of detoxification genes probably enhanced the adaptation of insects to their environments and, in recent evolutionary times, has helped them endure numerous artificial chemicals (Strode et al. 2007). This study found the NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6) gene family to have potentially expanded. These proteins comprise the biggest enzyme complex in the mitochondrial electron transfer chain and facilitate electron transfers from NADH to ubiquinone. Its two functional arms, NADH and ubiquinone, are involved in oxidation and reduction reactions (Agip et al. 2019). There seems to

be a link between cytochrome P450s and NADH dehydrogenase enzymes in targeted defenses against oxidative agents. Cytochrome P450 proteins are known to utilize NADPH-linked electrons to produce molecular oxygen, which activates specific oxidative attacks against many substrates (Werck-Reichhart and Feyereisen 2000). A negative consequence of the blood meal is the generation of reactive oxygen species and general oxidative stress upon the host, thus requiring the elimination of these substances. Interactions between cytochrome P450s and NADPH dehydrogenases could offer a possible means of eliminating deleterious oxidative stress post blood meals in hematophagous arthropods.

Further evidence suggests the interactions between heme iron, cytochrome P450s, and NADPH in lowering oxidative stress. The convergence of cytochrome P450 genes and NADPH dehydrogenase genes amongst the hematophagous taxa indicates the functional interdependence of these genes to oppose oxidative stress resulting from the blood meal. In the presence of cytochrome P450, methemoglobin can be reduced by NADPH (Faassen et al. 1988). Methemoglobin is the oxidized form of hemoglobin through a spontaneous oxidation reaction that produces reactive oxygen species such as superoxides and hydrogen peroxide (Everse and Hsia 1997). Hematophagous arthropods may have retained these genes collectively due to the increased need to mitigate the harm resulting from the blood meal, given the large amounts of hemoglobin ingested. Consistent with the findings of this study, heme, although known for promoting free radicals, is an integral feature of proteins such as cytochromes, electron transport enzymes, cellular respiration, signal transduction, and detoxification (Hamza and Dailey 2012). The expanded Hr51 gene is a heme-binding protein (de Rosny et al. 2008).

This study found three proteins within the larger cytochrome P450 family that expanded. However, in the Diptera, Siphonaptera, and Lepidopteran taxa analysis, cytochrome P450 4d1

(Cyp4d1) contracted and had higher gene copies in the non-blood-feeding taxa than in bloodfeeders. Gene contractions and expansions may suggest adapting to a new lifestyle (Feritas & Nery 2020). Gene contractions were noted in cardinal, curly su (Cysu), heavy metal tolerance factor 1 (Hmt-1), peroxinectin-like (Pxt), NADH dehydrogenase (ubiquinone) 13 kDa A subunit (ND-13A), nitric oxide synthase (Nos), and Cy4d1.

In the analysis of the hematophagous and non-hematophagous (Diptera) taxa and the hematophagous (Siphonaptera) taxa, five genes contracted (Cytochrome P450 4d1, Cytochrome P450 6a16, Cytochrome P450 6a17, Cytochrome P450 6a23, NADH dehydrogenase (ubiquinone) B14.5 A subunit and Cytochrome P450 6t3). These gene contractions may be due to adaptations to new lifestyles that find these genes energetically costly to retain, and therefore mutations that inhibit gene transcription may be advantageous, resulting in gene losses.

The methods of this paper are not limited to the represented study, extending to other comparable analyses, given the availability of genomic data and a "functional syndrome" that differs between closely related species. Such an approach would yield results that segregate and quantify genes implicated in the determined "functional syndrome."

The current use of pesticides to target disease vector insects threatens insects such as bees and other pollinators essential for global food production. Even at sub-lethal doses, pesticide exposure could make bees susceptible to pathogens and parasites and possibly a significant contributor to the global mortality of honeybee colonies (Pettis et al. 2012). Future conservation strategies could determine the genes involved in detoxifying the blood meal, such as cytochrome P450s, as these genes are known detoxifiers of many substances, including pesticides.

Interestingly the *A. mellifera* (honeybee) has a ten-fold decrease in xenobiotic detoxification genes such as cytochrome P450s compared to the hematophagous *A. gambiae*

(malaria mosquito). The significant decrease in these detoxification genes amongst bees could explain their sensitivity to pesticides (Claudianos et al. 2006) Approaches that block transcription of the expanded genes in this study offer a viable and environmentally friendly alternative to fight disease vector insects. A thorough study of these genes could help develop a class of pesticides specific to disease-causing insects, thus limiting the indiscriminate attack posed by pesticides on beneficial insects while also fighting against pesticide resistance.

Additionally, selective inhibition of heme and iron degradation pathways would neutralize disease-transmitting insects, allowing for the invention of novel compounds capable of targeted neutralization of disease vector insects. Indeed (Sterkel, Perdomo, et al. 2016) demonstrated that inhibition of the first and second enzymes in the phenylalanine/tyrosine degradation pathway, namely 4-hydroxyphenylpyruvate dioxygenase (HPPD) and tyrosine aminotransferase (TAT) with mesotrione an HPPD inhibitor caused the death of hematophagous insets after a blood meal, however, the non-hematophagous arthropods remained unaffected. Compared to the neurotoxic insecticides in current use, this less-toxic approach could bring about diseases vector insect controls that are less toxic to humans and the environment.

Predictably as taxa converge on similar phenotypes, functional genomic convergence transpires, as the results of this study indicate. This study discovered genes that expanded due to the blood-feeding lifestyle amongst dipterans. However, not all of the statistically significant genes expanded. There were also observed contractions in some gene families.

This study was limited by the absence of available annotated GO terms for hematophagous arthropods. *D. melanogaster* served as the reference species. However, a practical model organism, *D. melanogaster*, does not feed on blood, stressing the need for experimental assembly of GO terms for hematophagous arthropods. As GO terms of hematophagous arthropods become available, greater accuracy in assessing the genes implicated in the blood-feeding process would result in studies that follow this methodology. The expanded gene sequences in this study could be compared amongst the hematophagous and nonhematophagous taxa to observe possible convergent mutation patterns. Future studies evaluating the role of iron and heme in hematophagous arthropods could generate knockouts of genes such as cytochrome P450s in hematophagous arthropods and compare their ability to detoxify excess heme and iron to their wild-type counterparts.

5. Conclusion

In conclusion, this study underlines the importance of genomic data in assessing evolutionary convergence within gene families that may have expanded resulting from the blood-feeding lifestyle amongst hematophagous insects. Notably, in the analysis of the Dipteran, Siphonapteran, and Lepidopteran taxa, spook (cyp307A1), spookier (cyp307A2), cytochrome P450 12e1 (cyp12e1), hormone receptor and 51 (Hr51), NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6) possibly expanded due to hematophagy. NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW) is the only potential gene expansion in the Diptera and Siphonaptera taxa analysis. The approach presented in this study represents a promising approach for genomic studies aimed at discovering convergent gene family expansions in taxa.

Scientific Name	Order	Family	Phenotype	Accession Number
Aedes aegypti	Diptera	Culicidae	Blood	GCA_002204515.1
Anopheles gambiae	Diptera	Culicidae	Blood	GCA_000005575.1
Culex quinquefaciatus	Diptera	Culicidae	Blood	GCA_015732765.1
Culicoides tainanus	Diptera	Ceratopogonida	Blood	GCA_020740445.1
Glossina fuscipes	Diptera	Glossinidae	Blood	GCA_014805625.1
Lutzomyia longipalpis	Diptera	Psychodidae	Blood	GCA_000265325.1
Phlebotomus papatasi	Diptera	Psychodidae	Blood	GCA_000262795.1
Sarcophaga bullata	Diptera	Sacrophagidae	Blood	GCA_005959815.1
Stomoxys calcitrans	Diptera	Muscidae	Blood	GCA_001015335.1
Bactrocera tryoni	Diptera	Tephritidae	No	GCA_016617805.2
Belgica antarctica	Diptera	Chironomidae	No	GCA_000775305.1
Bradysia coprophila	Diptera	Sciaridae	No	GCA_014529535.1
Ceratitis capitata	Diptera	Tephritidae	No	GCA_000347755.4
Clanio marinus	Diptera	Chironomidae	No	GCA_900005825.1
Clogmia albipunctata	Diptera	Psychodidae	No	GCA_001014945.1
Contarinia nasturtii	Diptera	Cecidomyiidae	No	GCA_009176525.2
Hermeta illucens	Diptera	Stratiomyidae	No	GCA_905115235.1
Lucilia sericata	Diptera	Calliphoridae	No	GCA_015586225.1
Musca domestica	Diptera	Muscidae	No	GCA_000371365.1
Polypedilum vanderplanki	Diptera	Chironomidae	No	GCA_018290095.1
Polypedilum pembai	Diptera	Chironomidae	No	GCA_014622435.1
Propsilocerus akamusi	Diptera	Chironomidae	No	GCA_018397935.1
Rhagoletis pomonella	Diptera	Tephritidae	No	GCA_013731165.1
Rhagoletis zephyria	Diptera	Tephritidae	No	GCA_001687245.1
Teleopsis dalmanni	Diptera	Diopsidae	No	GCA_002237135.2
Zeugodacus cucurbitae	Diptera	Tephritidae	No	GCA_000806345.1
Ctenocephalides felis	Siphonaptera	Pulicidae	Blood	GCA_003426905.1
Aricia agestis	Lepidoptera	Lycaenidae	No	GCA_905147365.1
Bombyx mori	Lepidoptera	Bombycidae	No	GCA_014905235.2
Danaus plexippus	Lepidoptera	Nymphalidae	No	GCA_009731565.1
Helicoverpa armigera	Lepidoptera	Noctuidae	No	GCA_002156985.1
Manduca sexta	Lepidoptera	Sphingidae	No	GCA_014839805.1
Parage aegeria	Lepidoptera	Nymphalidae	No	GCA_905163445.1
Plutella xylostella	Lepidoptera	Plutellidae	No	GCA_905116875.3
Spodoptera frugiperda	Lepidoptera	Noctuidae	No	GCA_011064685.1
Trichoplusia ni	Lepidoptera	Noctuidae	No	GCA_003590095.1
Zerene cesonia	Lepidoptera	Pieridae	No	GCA_012273895.2

 Table 1: Species list with unique sequence identifier (Accession Number).

Table 2: TBL is represented	ASTN results of avera with the corresponding	ge D.melanoga: 7 p-value score.	ster-related iron	gene copy nur	nbers for genes	with p < 0.05 i	n comparing he	ematophagous L	iptera and Siph	onaptera and no	on-hematophag	ous Diptera and	Lepidoptera ta	ıxa. Each gene
Phenotype	Species	Order	Spo (P =0.0123)	Cyp4d1 (P = 0.0448)	Nos (P =0.0433)	ND-B16.6 $(P = 0.032)$	ND-13A $(P = 0.033)$	Hr51 (P = 0.0205)	Cyp 12el (P = 0.0136)	Hmt-1 $(P = 0.019)$	Cysu (P = 0.0081)	Spok (P = 0.0255)	Pxt $(P = 0.031)$	Cd (P = 0.0038)
Blood	A. aegypti	Diptera	9	4.67	3.33	4	2	1	9	2.5	1.5	3	2.75	1.5
Blood	A. gambiae	Diptera	9	4.67	5.67	4	2	2	4	2.67	1.5	3	2.6	1.5
Blood	C. quinquefaciatus	Diptera	9	5.33	4.33	2	2	1	10	3.2	1	3	1.8	1.5
Blood	C. tainanus	Diptera	9	3.33	3.33	4	2	1	0	4	1.5	3	2	1
Blood	G. fuscipes	Diptera	6	4	3.27	4	2	1	2	2.8	1.4	6	2	2.33
Blood	L. longipalpis	Diptera	6	3.5	5	6	0	1	12	7	1.5	3	1.83	1.5
Blood	P. paptasi	Diptera	6	3.33	3.3	2	2	1	12	8.67	1.75	3	1.5	1.5
Blood	S. bullata	Diptera	6	4.5	3	4	2	1	10	3.2	1.2	4.5	1.67	1.33
Blood	S. calcitrans	Diptera	6	7.5	3.27	2	2	1	6	2.8	1.4	6	1.83	2
Blood	C. felis	Siphonaptera	9	0	7.5	8	2	1.33	10	7	3	3	2.67	1
No	B. tryoni	Diptera	3	4	6.75	4	4	1	14	3.6	1.4	3	1.83	1.5
No	B. antarctica	Diptera	6	8	5.4	2	2	1	0	2.5	1.6	0	1.67	1
No	B. coprophila	Diptera	6	9.5	7.67	4	2	1	2	5.5	1.75	3	2	1.5
No	C. capitata	Diptera	6	3.5	5.75	4	4	1.5	10	4	1.6	3	1.67	1.5
No	C. marinus	Diptera	3	9.33	3.43	2	2	1	0	3	1.75	3	1.8	-
No	C. albipunctata	Diptera	3	8.67	3	2	0	1	0	5.5	1.75	3	1.8	1
No	C. nasturtil	Diptera	6	5	3.82	2	2	2	0	7	2.25	0	1.83	2
No	C. sericata	Diptera	12	8	3.27	4	0	1.2	2	3.6	1.2	6	3.17	2
No	H. illucens	Diptera	12	10	3.3	2	2	1	0	7	1.25	6	2	1.5
No	M. domestica	Diptera	9	6	4.09	4	2	1	2	2.8	1.2	4.5	1.83	2
No	P. pembai	Diptera	3	3.5	3.6	0	2	1	0	4	1	0	2.5	-
No	P. vanderplanki	Diptera	0	5.5	3.6	0	2	1	0	5.5	1.25	0	3	-
No	P. akamusi	Diptera	6	2	3	2	2	1	0	2	1.75	33	2	
No	R. pomonella	Diptera	6	6	11.5	4	6	1	6	3.2	1.2	3	1.33	1.5
No	R. zephyria	Diptera	6	4.67	10.5	4	4	1	6	6.8	3.4	3	2.33	2.5
No	T. dalmanni	Diptera	6	11.5	3.27	4	2	1.75	18	8	2.6	3	2.5	5
No	Z. cucurbitae	Diptera	6	7.33	6	2	4	1	10	3.6	1.2	3	1.5	1.33
No	A. agestis	Lepidoptera	3	2	6	2	2	0	0	9.33	3.5	3	4	5
No	B. mori	Lepidoptera	3	2	6.38	2	2	0	0	7.5	3.5	0	3.33	4
No	D. plexippus	Lepidoptera	3	2	6.75	2	2	0	0	14.5	3.25	3	4	2.5
No	H. armigera	Lepidoptera	6	6	6	2	4	0	0	11.33	3	3	3.33	5
No	M. sexta	Lepidoptera	6	6	5.62	2	2	1	0	10.67	5	0	3.67	2.5
No	P. aegeria	Lepidoptera	0	6	5.62	2	2	0	0	10	3.25	3	3	3.5
No	P. xylostella	Lepidoptera	3	10	5.67	2	2	0	0	13.33	3.25	3	2.33	3.5
No	S. frugiperda	Lepidoptera	12	4	5.62	2	4	0	0	12.67	4.5	0	3.67	3.5
No	T. ni	Lepidoptera	6	2	7.88	2	2	0	4	10	3	3	3.67	3.5
No	Z. cesonia	Lepidoptera	0	4	5.62	2	4	0	0	9.5	3	ω	ы	6

Table 3: Average Iron gene copy numbers for contracted and expanded genes comparing the hematophagous and non-hematophagous taxa. Spook (Spo), Cytochrome P450 12e1 (Cyp12e1), Hormone receptor 51(Hr51), Spookier (Spok) and NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6) potentially expanded. Cardinal (Cd), Curly Su (Cysu), Heavy metal tolerance factor 1 (Hmt-1), Peroxinectin-like (Pxt), NADH dehydrogenase (ubiquinone) 13 kDa A subunit (ND-13A), Nitric oxide synthase (Nos), and Cytochrome P450 4d1 (Cyp4d1) potentially contracted. "Sdv"= standard deviation results.

Gene Family Name	Hematophagous average gene copy number	Non-hematophagous average gene copy number
Spo	7.5 (Sdv = 1.581)	5.3 ((Sdv = 3.363)
Cyp12e1	7.2 (Sdv = 4.237)	2.7 (Sdv = 4.872)
Hr51	1.1 (Sdv = 0.322)	0.8 (Sdv = 0.597)
Spok	3.8 (Sdv = 1.275)	2.5 (Sdv = 1.715)
ND-B16.6	4 (Sdv = 1.886)	2.4 (Sdv = 1.155)
Cd	1.5 (Sdv = 0.403)	2.5 (Sdv = 1.492)
Cysu	1.6 (Sdv = 0.539)	2.3 (Sdv = 1.112)
Hmt-1	4.4 (Sdv = 2.273)	6.9 (Sdv = 3.638)
Pxt	2.1 (Sdv = 0.445)	2.5 (Sdv = 0.832)
ND-13A	1.8 (Sdv = 0.632)	2.5 (Sdv = 1.312)
Nos	4.2 (Sdv = 1.461)	5.5 (Sdv = 2.146)
Cyp4d1	4.1 (Sdv = 1.893)	5.8 (Sdv = 2.834)

Each gene re	LAS IN results	s of average n heir correspoi	eme gene cop nding n-value	scores.	ı genes wım b		праниз пеща	ind snokendor	otera and sipir	onapiera anu i	юп-пешаюрп	agous mipicia	i ann rebinob	icia iaxa.
Phenotype	Species	Order	Spo (P=0.012)	Cyp4d1 (P=0.045)	Nos (P=0.043)	ND-B16.6 (P=0.032)	ND-13A (P=0.033)	Hr51 (P=0.021)	Cyp 12e1 (P=0.014)	Hmt-1 (P=0.002)	Cysu (P=0.008)	Spok (P=0.025)	Pxt (P=0.031)	Cd (P=0.004)
Blood	A. aegypti	Diptera	4	2.33	2.22	4	2	1	3	3.75	1.5	2	2.75	1.5
Blood	A. gambiae	Diptera	4	2.33	3.78	4	2	2	2	4	1.5	2	2.6	1.5
Blood	quinquefacia	Diptera	4	2.67	2.89	2	2	1	5	4.8	1	2	1.8	1.5
Blood	C. tainanus	Diptera	4	1.67	2.22	4	2	1	0	6	1.5	2	2	1
Blood	G. fuscipes	Diptera	6	2	2.18	4	2	1	1	4.2	1.4	4	2	2.33
Blood	L. longipalpis	Diptera	6	1.75	3.33	6	0	1	6	10.5	1.5	2	1.83	1.5
Blood	P. paptasi	Diptera	6	1.67	2.2	2	2	1	6	13	1.75	2	1.5	1.5
Blood	S. bullata	Diptera	6	2.25	2	4	2	1	5	4.8	1.2	3	1.67	1.33
Blood	S. calcitrans	Diptera	6	3.75	2.18	2	2	1	3	4.2	1.4	4	1.83	2
Blood	C. felis	Siphonaptera	4	0	5	8	2	1.33	5	10.5	3	2	2.67	1
No	B. tryoni	Diptera	2	2	4.5	4	4	1	7	5.4	1.4	2	1.83	1.5
No	B. antarctica	Diptera	4	4	3.6	2	2	1	0	3.75	1.6	0	1.67	1
No	3. coprophila	Diptera	4	4.75	5.11	4	2	1	1	8.25	1.75	2	2	1.5
No	C. capitata	Diptera	4	1.75	3.83	4	4	1.5	5	6	1.6	2	1.67	1.5
No	C. marinus	Diptera	2	4.67	2.29	2	2	1	0	4.5	1.75	2	1.8	1
No	. albipunctate	Diptera	2	4.33	2	2	0	1	0	8.25	1.75	2	1.8	1
No	C. nasturtil	Diptera	4	2.5	2.55	2	2	2	0	10.5	2.25	0	1.83	2
No	C. sericata	Diptera	8	4	2.18	4	0	1.2	1	5.4	1.2	4	3.17	2
No	H. illucens	Diptera	8	5	2.2	2	2	1	0	10.5	1.25	4	2	1.5
No	M. domestica	Diptera	6	3	2.73	4	2	1	1	4.2	1.2	3	1.83	2
No	P. pembai	Diptera	2	1.75	2.4	0	2	1	0	6	1	0	2.5	1
No	. vanderplant	Diptera	0	2.75	2.4	0	2	1	0	8.25	1.25	0	3	1
No	P. akamusi	Diptera	6	1	2	2	2	1	0	3	1.75	2	2	1
No	R. pomonella	Diptera	4	3	7.67	4	6	1	3	4.8	1.2	2	1.33	1.5
No	R. zephyria	Diptera	4	2.33	7	4	4	1	3	10.2	3.4	2	2.33	2.5
No	T. dalmanni	Diptera	4	5.75	2.18	4	2	1.75	9	12	2.6	2	2.5	5
No	Z. cucurbitae	Diptera	4	3.67	4	2	4	1	5	5.4	1.2	2	1.5	1.33
No	A. agestis	Lepidoptera	2	1	4	2	2	0	0	14	3.5	2	4	5
No	B. mori	Lepidoptera	2	1	4.25	2	2	0	0	11.25	3.5	0	3.33	4
No	D. plexippus	Lepidoptera	2	1	4.5	2	2	0	0	21.75	3.25	2	4	2.5
No	H. armigera	Lepidoptera	4	3	4	2	4	0	0	17	3	2	3.33	5
No	M. sexta	Lepidoptera	4	3	3.75	2	2	1	0	16	5	0	3.67	2.5
No	P. aegeria	Lepidoptera	0	3	3.75	2	2	0	0	15	3.25	2	3	3.5
No	P. xylostella	Lepidoptera	2	5	3.78	2	2	0	0	20	3.25	2	2.33	3.5
No	S. frugiperda	Lepidoptera	8	2	3.75	2	4	0	0	19	4.5	0	3.67	3.5
No	T. ni	Lepidoptera	4	1	5.25	2	2	0	2	15	3	2	3.67	3.5
No	Z. cesonia	Lepidoptera	0	2	3.75	2	4	0	0	14.25	З	2	ы	6

Table 5: Average Heme gene copy numbers for contracted and expanded genes comparing the hematophagous and non-hematophagous taxa. Spook (Spo), Cytochrome P450 12e1 (Cyp12e1), Hormone receptor 51(Hr51), Spookier (Spok) and NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6) expanded. Cardinal (Cd), Curly Su (Cysu), Heavy metal tolerance factor 1 (Hmt-1), Peroxinectin-like (Pxt), NADH dehydrogenase (ubiquinone) 13 kDa A subunit (ND-13A), Nitric oxide synthase (Nos), and Cytochrome P450 4d1 (Cyp4d1) contracted. "Sdv" = standard deviation scores.

Gene Family Name	Hematophagous average gene copy number	Non-hematophagous average gene copy number
Spo	5 (Sdv = 1.0540)	3.5 (Sdv =2.284)
Cyp12e1	$\overline{3.6 \text{ (Sdv} = 2.119)}$	1.3 (Sdv = 2.462)
Hr51	1.1 (Sdv = 0.322)	0.7 (Sdv = 0.607)
Spok	2.5 (Sdv = 0.850)	1.7 (Sdv = 1.164)
ND-B16.6	4 (Sdv =1.886)	2.4 (Sdv = 1.134)
Cyp4d1	2 (Sdv =0.946)	2.9 (Sdv = 1.445)
Nos	2.8 (Sdv =0.974)	3.5 (Sdv =1.212)
ND-13A	1.8 (Sdv = 0.632)	2.4 (Sdv =1.134)
Hmt-1	6.6 (Sdv = 3.409)	10.6 (Sdv = 5.449)
Cysu	1.6 (Sdv =0.539)	2.4 (Sdv = 1.110)
Pxt	2.1 (Sdv = 0.445)	2.6 (Sdv = 0.812)
Cd	1.5 (Sdv =0.403)	2.5 (Sdv = 1.508)

Table 6: TB	LASTN results of avera	ige iron gene co	py numbers for anted has their	or genes with p	0 < 0.05 in con	nparing hemate	phagous Dipte	era and Siphon	naptera and
Phenotype	Species	Order	Cyp4da1 (P = 0.009)	Cyp6a16 (P = 0.014)	Cyp6a17 (P = 0.037)	Cyp6a23 (P = 0.037)	ND-B14.5A $(P = 0.047)$	Cyp6t3 (P = 0.047)	ND-PDSW $(P = 0.048)$
Blood	A. aegypti	Diptera	4.67	2	2	2	0	0	4
Blood	A. gambiae	Diptera	4.67	0	2	2	0	2	4
Blood	C. quinquefaciatus	Diptera	5.33	2	2	2	0	0	4
Blood	C. tainanus	Diptera	3.33	0	2	3	0	0	2
Blood	G. fuscipes	Diptera	4	0	0	2	2	4	2
Blood	L. longipalpis	Diptera	3.5	2	10	8	0	0	2
Blood	P. paptasi	Diptera	3.33	3	12	15	0	0	2
Blood	S. bullata	Diptera	4.5	4	9	9	2	4	2
Blood	S. calcitrans	Diptera	7.5	2	17	17	2	8	2
Blood	C. felis	Siphonaptera	0	0	16	18	0	0	4
No	B. tryoni	Diptera	4	7	14	15	2	16	2
No	B. antarctica	Diptera	8	4	3	7	2	0	2
No	B. coprophila	Diptera	9.5	0	9	9	0	0	2
No	C. capitata	Diptera	3.5	4	14	13	2	18	2
No	C. marinus	Diptera	9.33	6	10	9	2	4	2
No	C. albipunctata	Diptera	8.67	8	34	45	0	0	2
No	C. nasturtil	Diptera	5	0	3	3	0	0	2
No	C. sericata	Diptera	8	6	17	14	2	2	2
No	H. illucens	Diptera	10	5	9	18	2	4	0
No	M. domestica	Diptera	6	3	14	14	2	6	2
No	P. pembai	Diptera	3.5	0	16	16	0	4	2
No	P. vanderplanki	Diptera	5.5	0	13	12	0	0	2
No	P. akamusi	Diptera	2	0	3	3	2	0	2
No	R. pomonella	Diptera	6	10	12	13	2	5	4
No	R. zephyria	Diptera	4.67	8	14	18	2	12	2
No	T. dalmanni	Diptera	11.5	0	23	20	2	24	2
No	Z. cucurbitae	Diptera	7.33	12	18	20	2	6	2

Table 7: Average Iron gene copies for hematophagous Diptera & Siphonaptera and nonhematophagous Diptera. NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW) expanded. Cytochrome P450 4d1 (Cyp4d1), Cytochrome P450 6a16 (Cyp6a16), Cytochrome P4 50 6a17 (Cyp6a17), Cytochrome P450 6a23 (Cyp6a23), NADH dehydrogenase (ubiquinone) B14.5 A subunit (ND-B15.5A) and Cytochrome P450 6t3 (Cyp6t3) contracted. "Sdv" = standard deviation scores.

Gene Family Name	Hematophagous Average Gene Copy Number	Non-hematophagous Average Gene Copy Number
ND-PDSW	2.8 (Sdv = 1.033)	2 (Sdv = 0.707)
Cyp4d1	4.1 (Sdv = 1.893)	6.6 (Sdv = 2.687)
Сурба16	1.5 (Sdv = 1.434)	4.3 (Sdv = 3.917)
Cyp6a17	7.2 (Sdv = 6.391)	13.3 (Sdv = 7.647)
Сурба23	7.8 (Sdv = 6.663)	14.6 (Sdv = 9.387)
ND-B15.5A	0.6 (Sdv = 0.966)	1.4 (Sdv = 0.939)
Cyp6t3	1.8 (Sdv = 2.741)	5.9 (Sdv = 7.284)

and non-hema	atophagous Diptera ta	xa. Each gene	represented ha	is their correspondences	onding p-val	u comparing in ue scores.	onawpnagous	արագ այս թ	трионариста
Phenotype	Species	Order	Cyp4d1 (P = 0.009)	Cyp6a16 (P = 0.014)	Cyp6a17 (P = 0.037)	Cyp6a23 (P = 0.037)	ND-B14.5A (P = 0.047)	Cyp6t3 (P = 0.047)	ND-PDSW (P = 0.048)
Blood	A. aegypti	Diptera	2.33	1	1	1	0	0	4
Blood	A. gambiae	Diptera	2.33	0	1	1	0	1	4
Blood	C. quinquefaciatus	Diptera	2.67	1	1	1	0	0	4
Blood	C. tainanus	Diptera	1.67	0	1	1.5	0	0	2
Blood	G. fuscipes	Diptera	2	0	0	1	2	2	2
Blood	L. longipalpis	Diptera	1.75	1	5	4	0	0	2
Blood	P. paptasi	Diptera	1.67	1.5	9	7.5	0	0	2
Blood	S. bullata	Diptera	2.25	2	4.5	4.5	2	2	2
Blood	S. calcitrans	Diptera	3.75	1	8.5	8.5	2	4	2
Blood	C. felis	Siphonaptera	0	0	8	9	0	0	4
No	B. tryoni	Diptera	2	3.5	7	7.5	2	8	2
No	B. antarctica	Diptera	4	2	1.5	3.5	2	0	2
No	B. coprophila	Diptera	4.75	0	4.5	4.5	0	0	2
No	C. capitata	Diptera	1.75	2	7	6.5	2	9	2
No	C. marinus	Diptera	4.67	3	S	4.5	2	2	2
No	C. albipunctata	Diptera	4.33	4	17	22.5	0	0	2
No	C. nasturtil	Diptera	2.5	0	1.5	1.5	0	0	2
No	C. sericata	Diptera	4	3	8.5	7	2	1	2
No	H. illucens	Diptera	5	2.5	4.5	9	2	2	0
No	M. domestica	Diptera	3	1.5	7	7	2	ы	2
No	P. pembai	Diptera	1.75	0	8	8	0	2	2
No	P. vanderplanki	Diptera	2.75	0	6.5	6	0	0	2
No	P. akamusi	Diptera	1	0	1.5	1.5	2	0	2
No	R. pomonella	Diptera	3	5	6	6.5	2	2.5	4
No	R. zephyria	Diptera	2.33	4	7	9	2	6	2
No	T. dalmanni	Diptera	5.75	0	11.5	10	2	12	2
No	Z. cucurbitae	Diptera	3.67	6	9	10	2	3	2

Table 9: Average heme gene copies for hematophagous (Diptera & Siphonaptera) and nonhematophagous taxa (Diptera). NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW) expanded. Cytochrome P450 4d1 (Cyp4d1), Cytochrome P450 6a16 (Cyp6a16), Cytochrome P450 6a17 (Cyp6a17), Cytochrome P450 6a23 (Cyp6a23), NADH dehydrogenase (ubiquinone) B14.5 A subunit (ND-B15.5A) and Cytochrome P450 6t3 (Cyp6t3) contracted. "Sdv" = standard deviation scores.

Gene Family Name	Hematophagous Average Gene Copy Number	Non-hematophagous Average Gene Copy Number
ND-PDSW	2.8 (Sdv =1.033)	2 (Sdv = 0.707)
Cyp4d1	2 (Sdv = 0.946)	3.3 (Sdv = 1.344)
Cyp6a16	0.8 (Sdv = 0.717)	2.1 (Sdv = 1.959)
Cyp6a17	3.6 (Sdv = 3.195)	6.6 (Sdv = 3.823)
Cyp6a23	3.9 (Sdv = 3.332)	7.3 (Sdv = 4.694)
ND-B15.5A	0.6 (Sdv = 0.966)	1.4 (Sdv = 0.939)
Cyp6t3	0.9 (Sdv = 1.370)	3 (Sdv = 3.642)

Table10a: *D. melanogaster* associated genes with corresponding FlyBase ID of each gene. Blue highlights potential gene expansions in the analysis of hematophagous Diptera, Siphonaptera, and non-hematophagous Diptera and Lepidoptera.

D.melanogaster associated genes	FlyBase ID
Spook (Spo)	FBgn0003486
Cytochrome P450 12e1 (Cyp12e1)	FBgn0037817
Hormone receptor 51 (Hr51)	FBgn0034012
Spookier (Spok)	FBgn0086917
NADH dehydrogenase (ubiquinone) B16.6 subunit (ND-B16.6)	FBgn0029868
Cardinal (Cd)	FBgn0263986
Curly su (Cysu)	FBgn0038511
Heavy metal tolerance factor 1(Hmt-1)	FBgn0038376
Peroxinectin-like (Pxt)	FBgn0261987
NADH dehydrogenase (ubiquinone) 13 kDa A subunit (ND-13A)	FBgn0031684
Nitric oxide synthase (Nos)	FBgn0011676
Cytochrome P450 4d1 (Cyp4d1)	FBgn0005670

Table10b: *D. melanogaster* associated genes with corresponding FlyBase ID of each gene. Blue highlights potential gene expansions in the analysis of hematophagous Diptera, Siphonaptera, and non-hematophagous Diptera.

D.melanogaster associated genes	FlyBase ID
NADH dehydrogenase (ubiquinone) PDSW subunit (ND-PDSW)	FBgn0021967
Cytochrome P450 4d1 (Cyp4d1)	FBgn0005670
Cytochrome P450 6a16 (Cyp6a16)	FBgn0031726
Cytochrome P450 6a17 (Cyp6a17)	FBgn0015714
Cytochrome P450 6a23 (Cyp6a23)	FBgn0033978
Cytochrome P450 6t3 (Cyp6t3)	FBgn0033697
NADH dehydrogenase (ubiquinone) B14.5 A subunit (ND-B14.5A)	FBgn0033978



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