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A phylogenetic examination of host use evolution in the quinaria and testacea groups of Drosophila

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Abstract

Adaptive radiations provide an opportunity to examine complex evolutionary processes such as ecological specialization and speciation. While a well-resolved phylogenetic hypothesis is critical to completing such studies, the rapid rates of evolution in these groups can impede phylogenetic studies. Here we study the *quinaria* and *testacea* species groups of the *immigrans-tripunctata* radiation of *Drosophila*, which represent a recent adaptive radiation and are a developing model system for ecological genetics. We were especially interested in understanding host use evolution in these species. In order to infer a phylogenetic hypothesis for this group we sampled loci from both the nuclear genome and the mitochondrial DNA to develop a dataset of 43 protein-coding loci for these two groups along with their close relatives in the *immigrans-tripunctata* radiation. We used this dataset to examine their evolutionary relationships along with the evolution of feeding behavior. Our analysis recovers strong support for the monophyly of the *testacea* but not the *quinaria* group. Results from our ancestral state reconstruction analysis suggests that the ancestor of the *testacea* and *quinaria* groups exhibited mushroom-feeding. Within the *quinaria* group, we infer that transition to vegetative feeding occurred twice, and that this transition did not coincide with a genome-wide change in the rate of protein evolution.

Graphical Abstract

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Keywords

ancestral state reconstruction; mycophagy; host specialization; phylogeny; Drosophila

1. Introduction

Recent adaptive radiations are a fertile ground for understanding evolutionary processes such as how speciation occurs and how adaptive traits evolve (Givnish and Sytsma, 1997; Grant and Grant, 2008; Schluter, 2000). Critical to these inferences is a well-resolved and accurate phylogenetic history of the group being studied (Hahn and Nakhleh, 2016; O'Meara, 2012). In this study, we investigate the phylogenetic history of the quinaria and testacea groups of the genus Drosophila, which are a developing model system for ecological genetic studies. Both of these groups occur within the *immigrans-tripunctata* radiation of the subgenus *Drosophila* and are found in temperate and boreal forests of the Northern hemisphere. The *quinaria* group is a young adaptive radiation $\left(\sim\right)$ 19.5MYA; Izumitani et al., 2016) composed of 26 species, and the testacea group is smaller, comprising four species.

Within the *quinaria* and *testacea* groups, there is a remarkable amount of morphological and life history variation. Most species in the quinaria group and all members of the testacea group exhibit mushroom-feeding, though within the *quinaria* group several species have switched to utilizing decaying vegetation. In addition to variation in food substrate preference, within and among these species there is well characterized variation in body and wing pigmentation (Dombeck and Jaenike, 2004; Werner et al., 2010), circadian rhythms (Simunovic and Jaenike, 2006), parasite prevalence and resistance (Haselkorn et al., 2009; Perlman and Jaenike, 2003; Perlman et al., 2003), mycotoxin tolerance (Jaenike et al., 1983; Spicer and Jaenike, 1996; Stump et al., 2011), endosymbiont infection and resistance (Dyer and Jaenike, 2004; Haselkorn et al., 2013; Jaenike, 2007; Stahlhut et al., 2010; Unckless and Jaenike, 2012; Werren and Jaenike, 1995), cold and desiccation tolerance (Gibbs and

Matzkin, 2001; Kimura, 2004), behavioral isolation (Arthur and Dyer, 2015; Humphreys et al., 2016; Jaenike et al., 2006), sexual signals used in mate discrimination (Dyer et al., 2014; Giglio and Dyer, 2013), and meiotic drivers and suppressors (Dyer, 2012; Dyer et al., 2007; Jaenike, 1999).

Several aspects promote the use of these species groups as a model for evolutionary ecology studies. The mushroom-feeders in both groups are composite generalists on fleshy basidiomycete mushrooms, and all life stages revolve around mushrooms: the adults are attracted to mushrooms to mate, and then females lay their eggs on mushrooms that serve as a substrate for larval development. Species that consume decaying vegetation are specialists on water plants such as skunk cabbage (Grimaldi and Jaenike, 1983). These species are easy to collect from baits or naturally occurring hosts, and large numbers of flies can be reared in the lab, where they have a 2–3 week generation time. In addition, many can be hybridized in the lab ($e.g.,$ Bray et al., 2014; Humphreys et al., 2016; Shoemaker et al., 1999), which facilitates forward genetic studies. Genetic transformation has been performed in D. guttifera, enabling reverse genetics (Werner et al., 2010), and genomic resources are being developed that will aide in downstream genetic studies.

Here we use a multi-locus genome-wide approach to resolve the phylogeny of the *quinaria* and testacea group species, which will allow ecological and genetic questions to be addressed in an evolutionary framework. When branch lengths are short as in recent radiations, processes such as incomplete lineage sorting and hybridization can alter the evolutionary history of molecular characters (Edwards, 2009; Mallet et al., 2016). Thus, when attempting to reconstruct a species tree for a recently diverging group, it is important to consider the type of data, such as the genomic region being sampled, as well as the analytical method(s), to avoid being misled by non-representative phylogenetic signal. To date, the phylogenetic inference of these groups (Dyer et al., 2011b; Perlman et al., 2003; Spicer and Jaenike, 1996), as well as of their placement in the immigrans-tripunctata radiation (Hatadani et al., 2009; Izumitani et al., 2016; Morales-Hojas and Vieira, 2012; O'Grady and DeSalle, 2018), has used datasets primarily composed of mitochondrial (mtDNA) markers, while some datasets have also included the Y-chromosome and a few nuclear loci. The results across loci and studies are largely incongruent.

Several aspects about the biology of these flies may complicate phylogenetic inferences. First, based on previous molecular evolutionary results, divergence times are low and thus the speciation rates appear to be high (Perlman et al., 2003; Spicer and Jaenike, 1996), and based on population genetic work the effective population sizes of these species are large (e.g. Dyer et al., 2013; Dyer et al., 2007; Dyer and Jaenike, 2005; Dyer et al., 2011b); both of these factors can increase the amount of incomplete lineage sorting. In lineages where incomplete lineage sorting has occurred, phylogenetic analyses based on concatenated multigene datasets can recover strong support for an incorrect topology (Kubatko and Degnan, 2007; Mendes and Hahn, 2018; Roch and Steel, 2014). An alternative approach is to recover species trees using a multispecies coalescent model, treating each locus as a distinct unit and then inferring the species tree from individual gene trees (Edwards et al., 2016). Second, many species within each group can hybridize and some show evidence of recent hybridization (e.g. Bray et al., 2014; Dyer et al., 2011b; Humphreys et al., 2016; Jaenike et

al., 2006; Patterson and Stone, 1952). Third, several species are infected with Wolbachia and other maternally-inherited endosymbionts, which can cause the linked mtDNA to show nonneutral or incongruent evolutionary patterns (Cariou et al., 2017; Hurst and Jiggins, 2005; Shoemaker et al., 2004).

In this study, we sampled all of the *testacea* group species, most of the *quinaria* group species, and several closely related outgroups in the *immigrans-tripunctata* radiation. We constructed a dataset composed of sequence data for 43 protein-coding loci that are located throughout the genome $(i.e.,$ on the autosomes, sex chromosomes, and mtDNA). To account for the natural history of these species, we reconstructed phylogenetic hypotheses for the quinaria and testacea groups using both analyses of a concatenated molecular dataset as well as a coalescent-based model approach that recovers a species tree from individual gene trees. We then compared the resulting topologies and examined phylogenetic incongruence in the context of specific genomic regions. Finally, we used these results to infer the evolutionary history of feeding behavior within the group using comparative approaches. Specifically, we examined the transitions between mushroom- and vegetation-feeding life histories and the associated rates of protein evolution.

2. Materials and Methods

2.1 Taxon sampling

Fly stocks used in this study were either derived from wild-caught strains or obtained from the Drosophila Species Stock Center. In Table 1, we list the 27 species used, along with the origin of the sequenced strain and reported feeding behavior of the species. We used one strain per species, and our species sampling includes all four of the known testacea group species and 18 of the 26 known *quinaria* group species. As there is strong behavioral isolation among populations of D. subquinaria (Jaenike et al., 2006), we included two strains of this species, originating from inland (Hinton, Alberta) and coastal (Portland, Oregon) regions of its geographic range. Several *quinaria* group species were not represented in this study because fresh genomic DNA was not available, and they include the North American species *D. macroptera, D. rellima,* and *D. palustris*, the European species *D. limbata*, and the Asian species D. curvispina, D. kuntzei, and D. unispina. We also included representative members of other species groups known to occur in the *immigrans-tripunctata* radiation, including D. immigrans, D. tripunctata, and D. bizonata from the immigrans, tripunctata and bizonata species groups respectively, and D. cardini and D. acutilabella from the cardini group.

2.2 DNA sequencing

We sequenced portions of 43 protein-coding loci from each species (Table S1). The names of the loci, their genomic location using the corresponding Müller Element in Drosophila melanogaster, and the primers for PCR and sequencing are in Tables S2 and S3. These loci are spread throughout the genome and based on gene location in D. melanogaster this includes ten loci on Müller element A (which is the X-chromosome in these species), five on element B, eight on element C, three on element D, ten on element E, two on the 'dot' element F, three on the mtDNA, and two on the Y chromosome. There is general

conservation across Drosophila of genes to Müller elements, but substantial scrambling within chromosomes due to rearrangements (Bhutkar et al., 2008). Some of the sequences for the Y-chromosome and mtDNA loci were obtained from previous studies (Dyer et al., 2011b; Perlman et al., 2003).

Genomic DNA was extracted from single flies using the Qiagen Puregene Kit, and PCR used standard methods. PCR amplicons were purified using Exosap-IT (Thermo Fisher Scientific), sequenced in both directions using Big Dye 3.1 chemistry (Applied Biosystems), and run on an ABI 3730xl DNA Analyzer at the Georgia Genomics Facility at the University of Georgia. Chromatograms were analyzed using Sequencher (Gene Codes). Sites that were heterozygous (as evident by double peaks) were left as heterozygous in the analyses. All sequences have been deposited in Genbank (Accession numbers MK016667-MK017684).

The sequences were aligned in Geneious v10 (Kearse et al., 2012). Open reading frames were assigned using annotated *D. melanogaster* orthologs as a guide (Flybase.org). Initial alignments of coding sequences were completed using the protein translation implementing a BLOSUM weight matrix, and then the sequences were converted back to DNA sequences and manually adjusted. Noncoding sequences (introns and untranslated regions) were removed; these contained many gaps among species and thus were of limited use at this phylogenetic scale. For one locus (period) a highly repetitive region of the coding sequence that we could not align with confidence was also excluded from the analyses. The final dataset included 21,486 base pairs per species. We used MEGA v7 (Kumar et al., 2016) to assess the number of informative sites in each alignment (Table S3).

2.3 Phylogenetic methods

To reconstruct species trees, we completed phylogenetic analyses from concatenated datasets using maximum likelihood and Bayesian inference methods as well as from individual gene trees using a coalescent-based approach. For the analyses of the concatenated datasets and individual gene sequences, data partitions and models of evolution were determined using a 'greedy' search and the Bayesian Information Criterion (BIC) implemented in PartitionFinder v2.1.1 (Guindon et al., 2010; Lanfear et al., 2012; Lanfear et al., 2017). See Appendix S1 for the optimal partitioning scheme and models of evolution used in the phylogenetic analyses.

For the phylogenetic analysis of the concatenated datasets, we used maximum likelihood (ML) and Bayesian inference (BI) analyses on three datasets: one that contained all 43 loci (21,486 bp), another that excluded the three mtDNA loci (40 loci, 19,973 bp), and one composed only of the three mtDNA loci. The ML analysis was completed with RAxML-HPC2 v8.2.8 (Stamatakis, 2006, 2014). To obtain branch support for the most likely topology we completed 1,000 bootstrap replications, and only support values greater than 50% are provided in the results. We conducted the BI analysis using MrBayes v3.2.6 (Ronquist et al., 2012). When the best-fit model of evolution identified by PartitionFinder could not be implemented due to software limitations, we used the next most complex model that could be implemented. The BI analyses were performed with four chains, one cold and three hot, using the default temperature settings. The analysis consisted of two simultaneous, independent runs of 20,000,000 generations with samples drawn every 1,000 generations.

The first 5,000,000 generations were discarded as 'burn-in'. To confirm that the two runs had converged, the potential scale reduction factor (PSRF; Gelman and Rubin 1992) was calculated. The PSRF should approach 1.000 as independent runs converge. Posterior probability percent values provided the branch support values. Both the ML and BI analyses were completed on the CIPRES scientific gateway (Miller et al., 2010), and the resulting phylogenies were rooted using D. immigrans.

We inferred species trees using the coalescent-based model implemented in ASTRAL-III v5.6.1 (Zhang et al., 2017). First, using RAxML v8.2.4 we obtained the most likely unrooted topology (i.e., gene tree) along with 1,000 bootstrap replicates for each individual locus from the nuclear DNA. We concatenated the three mtDNA loci and analyzed them as a single locus. Then, we reconstructed a species tree using the unrooted gene trees of the 40 nuclear loci plus the mtDNA topology. From these gene trees, we also estimated species trees using several subsets of the data, including the loci from each Müller element separately, the Y-chromosome, and all 40 nuclear loci excluding the mtDNA. We used the 'multi-locus bootstrapping' option in ASTRAL-III to infer the species tree, and we used the '-q' option in ASTRAL-III to score the species trees produced by the analyses of the concatenated datasets. The ML analyses of the individual loci and the ASTRAL-III analyses were completed using the Georgia Advanced Computing Resource Cluster. The species trees resulting from the summary coalescent model analysis were rooted using *D. immigrans*.

2.4 Ancestral state reconstruction

To reconstruct the evolutionary history of the feeding behaviors in this group, the feeding behaviors of the 27 species included in the phylogenetic analyses were divided into the following categories: (A) mushroom, (B) vegetative, and (C) fruit (Table 1; Appendix S2). Species that exhibit multiple feeding behaviors were coded as possessing multiple states. For instance, D. tripunctata feeds on both mushrooms and fruit and thus was coded as AC rather than a unique state. We reconstructed the evolution of feeding behavior using a Bayesian approach, implementing the Bayesian Binary MCMC (BBM) ancestral state reconstruction method in RASP (Yu et al., 2015). The analysis was run for 5,000,000 generations using four chains that were sampled every 100 generations. A fixed Jukes-Cantor model with null root distribution was used for the analysis, and the first 1,000 samples were discarded as burn-in.

2.5 Rates of molecular evolution

The vegetative species in the *quinaria* group are thought to feed only on a limited range of decaying water plants, whereas the mushroom-feeding species are generalists on fleshy basidiomycete mushrooms and thus may have a broader host range. Host specialization may decrease the effective population size (e.g., Li et al., 2014), which in turn may render purifying selection less efficient and result in a higher rate of protein evolution due to the fixation of slightly deleterious mutations. We tested for relaxation of purifying selection in the vegetative species by analyzing the rates of protein evolution (ω , or d_N/d_S) of each locus. We implemented branch models in codeml of PAML v4.8a (Yang, 2007), and for each locus we compared a model with a single ω value across the entire phylogeny (ω-all) versus a model with two ω values, one for the clade(s) with the vegetative species (ω -veg) and

another for the rest of the phylogeny (ω -nonveg). For each locus, we accounted for gene and species tree differences by completing the analyses using the individual gene tree from the RAxML analyses, the pruned summary species tree from the coalescent model analysis run in ASTRAL-III, and the pruned ML tree from the concatenated dataset. We used likelihood ratio tests to ask whether the model with two ω values was a better fit than the model with one ω value. We determined each P-value using a χ^2 -distribution with one degree of freedom; for each phylogeny we implemented a Bonferroni correction and thus the adjusted significance threshold is $P = 0.05/40$ loci = 0.0013. We used nonparametric statistics to compare the ω values across loci from each type of analysis.

3. Results

To examine the evolutionary relationships within the *quinaria* and *testacea* groups, we constructed a dataset of 43 protein-coding loci that represents every autosome, both sex chromosomes, and the mtDNA from 27 species (Table S1). The final dataset contained 21,486 bp per species, of which 7,051 were variable sites (VS) and 5,022 were parsimony informative sites (PI) (Table S3). Not every locus could be sequenced in every species (Table S1). The average number of loci sequenced per species was 39 of 43 loci (std dev = 2.7) and each locus was sequenced in an average of 26 of 27 species (std dev $= 2.8$). The proportion of VS and PI sites did not differ across the chromosomes ($p = 0.96$ and $p = 0.94$) respectively; Fig. S1).

3.1 Phylogenetic Analyses

The ML and BI analyses of the concatenated dataset that included all of the nuclear and mtDNA loci recovered well-resolved and strongly supported species trees (Fig. 1 A–B). The topologies were largely congruent, and both contained three clades composed of the same species, which we refer to as clades A, B, and C. Clade A (Bootstrap $(BS) = 100$; Posterior Probability (PP) = 100), which contained the species representing the *tripunctata* and *cardini* groups, is sister to Clades $B + C$. Clade B is composed of the *bizonata* and *testacea* groups $(BS = 100; PP = 100)$ and is sister to a monophyletic *quinaria* group $(BS = 100; PP = 100)$. We refer to the entire *quinaria* group as Clade C, and within the *quinaria* group, the analyses recovered two divergent subgroups, which we refer to as Clades C1 and C2 ($BS = 100$; $PP =$ 100 for both). Clade C2 is comprised of D. angularis, D. falleni, D. innubila, D. brachynephros, and D. phalerata, while Clade C1 is comprised of all other quinaria group species (Fig. 1). In both the ML and BI species trees, the two subgroups were comprised of the same species, and the relationships among the species were also identical.

The coalescent-based model analysis in ASTRAL-III using all the loci also recovered a well-resolved, strongly supported species tree (Fig. 1C). While this phylogeny contained two of the same main clades from the ML and BI analyses (Clades A and B; BS = 100 for both), it did not support the monophyly of the *quinaria* group (Clade C). Instead, the *quinaria* group was paraphyletic, though the two *quinaria* subgroups were recovered (Clade C1, BS = 100; Clade C2, $BS = 100$). The C2 *quinaria* subgroup ($BS < 50$) was sister to the lineage ($BS >$ 78.6) containing the *tripunctata + cardini* (Clade A; $BS = 100$) and *bizonata + testacea*

clades (Clade B; $BS = 100$). The C1 subgroup was sister to the other three clades (A, B, and C2).

In addition to differences among the species relationships recovered in the analyses of the concatenated dataset and the coalescent-based model analysis, we scored the species trees from each analysis of the 43 locus dataset in ASTRAL-III to determine the fraction of the induced quartet trees resulting from the input set of gene trees that are in the given species tree $(i.e.,$ normalized quartet score (NQS)). The NQS can serve as a measure of the presence of incomplete lineage sorting (Mai et al., 2017; Sayyari and Mirarab, 2016). For each of the three analytical methods, there was only moderate congruence of the gene trees and the species trees. Specifically, the discordance between the gene trees and species trees was approximately the same in the coalescent-based (NQS = 0.7895), the BI analysis (NQS = 0.7858), and the ML analysis ($NQS = 0.7858$). These findings indicate that 79% of the quartet trees generated from the gene trees could be found in the species trees of each of the three analytical methods.

As previous examinations of the evolutionary relationships of these species utilized datasets composed primarily of mtDNA markers (Dyer et al., 2011b; Hatadani et al., 2009; Izumitani et al., 2016; Morales-Hojas and Vieira, 2012; Perlman et al., 2003; Spicer and Jaenike, 1996), we examined the phylogenetic signal in the three mtDNA loci separately from the nuclear loci using ML and BI analyses (Fig. S2 A, B). Both analyses recovered topologies that contained the three main clades (Clades A-C) and the two quinaria subgroups (Clades C1 and C2) found in the dataset containing all loci (Fig. 1 A, B). However, the relationships among the three clades differed from what was observed in the analysis of all the loci and neither analysis was able to resolve the relationships of these groups (Fig S2 A, B). Both analyses recovered moderate to weak support for the monophyly of the *quinaria* group (BS) 88; PP > 88), but the evolutionary relationships among the species within each subgroup differed from those found in the all gene analyses.

Given these findings, we completed a ML, BI, and coalescent-based analysis using only the 40 nuclear loci (Fig. 2 A–C). In each of the three analyses, the main clades and the relationships among them did not change with the exclusion of the mtDNA relative to those same analyses with the full dataset. With the ML analysis (Figs. 1A, 2A), the only difference between the topologies occurred among the species relationships recovered for D. magnaquinaria, D. deflecta, and D. subpalustris in subgroup C1. Overall the support values for some relationships showed minor decreases, and the resulting ML species tree exhibited slightly more discordance (NQS = 0.7847) with the gene trees. For the BI analysis (Figs. 1B, 2B), the only topological change occurred among the same three species as in the ML analysis, but the support values for all relationships remained at $PP = 100$. As with the species tree from the ML analysis, the BI species tree obtained exhibits greater discordance with the gene trees ($NQS = 0.7847$). The species tree obtained from the coalescent-based analysis (Fig. 2C) was also identical to that recovered by the analysis of the 43 loci (Fig. 1C). Exclusion of the mtDNA increased support in the sister relationship between the C2 clade and Clades $A + B$ (BS > 60.5). In addition, the amount of discordance between the species tree and gene trees did not change for the coalescence-based analysis (NQS = 0.7894). Because of the incongruence of the mtDNA vs. nuclear loci and the well-known

issues with using mtDNA in phylogenetic analyses of insects (Cariou et al., 2017; Hurst and Jiggins, 2005), we used the ML and coalescent-based species tree topologies obtained from the nuclear loci for all downstream analyses.

3.2 Genomic Incongruence

To identify incongruence among the loci in the different genomic regions, we used a coalescent-based model analysis to assemble species trees from subsets of the molecular dataset corresponding to different genomic regions. To assess the incongruence among the nuclear loci, we recovered species trees for each Müller element and the Y chromosome (Fig. S3A-G) and compared them to species tree recovered by the ASTRAL analysis of the 40 nuclear loci dataset (Fig. 2C). Every genome region except Müller D recovered a phylogeny that contained the A, B, C1, and C2 clades each as a monophyletic group. In the Müller D species tree (Fig. S3D), both the C1 and C2 clades were found to be paraphyletic. The trees recovered from the Müller B, C, and F supported a monophyletic *quinaria* group $(i.e., Claude C)$, whereas Müller elements A, D, E, and the Y-chromosome did not support a monophyletic *quinaria* group. When the *quinaria* group was not monophyletic, it was always the C2 clade that was more closely related to either the *testacea + bizonata* groups (Clade B) or both this and the *tripunctata* + *cardini* groups (Clades A and B). In general, the different genomic regions did not recover consistent sister relationships among the A, B, and C clades. Non-recombining and/or regions involved in reproductive isolation (*i.e.*, Müller F and the sex chromosomes) may show reduced incomplete lineage sorting and/or reduced introgression, but even comparing these three regions they all differed in the sister relationships of the main clades and the monophyly of the *quinaria* group.

3.3 Feeding behavior evolution

To infer the instances of host switching in the *quinaria* group, we reconstructed the most likely ancestral feeding behavior states on the ML and ASTRAL species trees recovered from the analysis of all loci except the mtDNA (Fig. 3A,B). Despite the differences in the evolutionary relationships identified by the two phylogenetic analyses, they each suggest similar patterns of feeding behavior evolution. First, within the *quinaria* group there were two independent instances of switching from mushrooms to vegetation, both within the C1 clade. Three of the vegetative species, *D. deflecta*, *D. magnaquinaria*, and *D. subpalustris*, form a monophyletic group whose common ancestor was likely vegetative feeding (ML: 89.18%, Probability (P)=0.885; ASTRAL: 90.76%, P=0.901). The fourth vegetative species in our analysis, *D. quinaria*, is a distinct lineage in the group and a separate evolutionary instance of host switching, and we recovered moderate to strong support that its ancestor most likely exhibited mushroom feeding (ML: 87.26%, P=0.873; ASTRAL: 82.81%, P=0.694). Second, the common ancestor of the *testacea* and *quinaria* groups was likely mushroom feeding. This is more strongly supported in the ML species tree, where these groups are monophyletic (60.74%, P=0.536). This node on the ASTRAL tree also contains the *tripunctata* and *cardini* groups that consume fruit and mushrooms, thus the support for this finding is substantially weaker (75.57%, P=0.26). Third, we recovered moderate to strong support (ML: P=0.511; ASTRAL: P=0.552) that the most likely feeding behavior of the ancestor of the immigrans-tripunctata radiation (Node 1) was fruit feeding (ML: 84.11%; ASTRAL: 73.08%), and weak to moderate support that the ancestor of the ingroup taxa

(Node 2) exhibited mushroom feeding behavior (ML: 60.74%, P=0.536; ASTRAL: 75.57%, $P=0.26$).

3.4 Rates of protein evolution

Within the *quinaria* group some species are generalists on fleshy basidiomycete mushrooms while others are specialists on decaying water plants. A narrower host range of the vegetative species may result in a decrease in the effective population size and thus a relaxation of purifying selection. However, we do not find compelling evidence for relaxation of purifying selection in these lineages (Fig. S4). For each locus, we compared a model with two ω (or d_N/d_S) values, one for the clade(s) of the vegetative species (ω -veg; see Table 1) and another for the rest of the phylogeny (ω -nonveg), with a model with one ω for the entire tree (ω -all). We found a two ω model to be a better fit for one locus using the ML species tree (per), for four loci using the ASTRAL species tree (ebony, mof, per, skp), and for two loci using the gene trees (*per, skp*) (Table S4). The *per* locus was the only locus with consistent evidence of an elevated d_N/d_S value in the vegetative lineages (Table S4). Most of the loci appear to be under strong purifying selection (Fig. S4), and across all 40 nuclear loci overall we found no significant differences between the ω values calculated from any of the three topologies (e.g., ML species tree: median ω -all = 0.031, median ω -veg $= 0.031$, median ω-nonveg $= 0.033$; Steel-Dwass pairwise tests, all $P > 0.85$).

4. Discussion

Previous phylogenetic studies of the *quinaria* and *testacea* groups and their placement within the *immigrans-tripunctata* radiation have produced results that are largely incongruent due, in part, to limited sampling of genomic compartments (*i.e.*, just mtDNA, Y-chromosome, and/or a few nuclear loci) (Dyer et al., 2011a; Dyer et al., 2011b; Hatadani et al., 2009; Izumitani et al., 2016; Morales-Hojas and Vieira, 2012; Perlman et al., 2003; Spicer and Jaenike, 1996). In addition, hybridization and incomplete lineage sorting can obscure phylogenetic signal in recent adaptive radiations, and both may be occurring in species groups. Our study represents the first attempt to analyze evolutionary relationships within and among the *quinaria* and *testacea* groups and several other closely species groups in the immigrans-tripunctata radiation using many loci that are distributed across all of the chromosomes of the genome.

4.1 Congruence among loci and approaches

Given our knowledge of the natural history and genetics of the *quinaria* group, there is a strong possibility for the phylogenetic history of the individual loci in our dataset to differ from that of the "true" species tree. Thus, we reconstructed species trees using ML and BI analyses of a concatenated dataset and also using a coalescent-based model analysis of the individual gene trees. We found that the species tree from each of three analyses contained the same four clades $(A, B, C1, C2)$, though the evolutionary relationships among these clades varied depending on the type of analysis. Most significantly for our species of interest, the C1 and C2 clades that comprise the *quinaria* group species were not always monophyletic, though the same species were always found within each clade. While the analyses of the concatenated dataset recovered strong support for the monophyly of the

quinaria group, the coalescent-based species analysis found the *quinaria* group to be paraphyletic with respect to the other ingroup taxa. Given this incongruence across analysis methods, we examined subsets of the data to understand the factors that may be contributing to topology differences both within the *quinaria* group as well as among the other species groups we sampled.

mtDNA is especially prone to incongruent phylogenetic patterns due to transfer between species through hybridization and linkage with maternally-transmitted endosymbionts such as Wolbachia and Spiroplasma. We know that both of these phenomena occur in the quinaria group – for instance, some *D. quinaria* harbor a highly divergent mtDNA haplotype at low frequency that originates from an unknown and/or extinct species (Dyer et al., 2011a) and D. subquinaria harbors a low frequency of D. recens mtDNA (Bewick and Dyer, 2014; Jaenike et al., 2006). Both of these events are likely due to past hybridization events. Indeed, in our study, the phylogenies constructed from the mtDNA are incongruent with the nuclear loci. For instance, the mtDNA trees did not support the monophyly of the testacea group, nor did they support the sister relationship between *D. occidentalis* and *D. suboccidentalis* (Fig. S3). This is in contrast to all of our species trees, no matter the analytical method, as well as the previous parsimony-based analyses of these species that used only these mtDNA loci ($e.g.$ Perlman et al. 2003). Thus, our findings support many previous suggestions that the mtDNA can bias species-level phylogenies, and we suggest that the species tree excludes this genomic region. Given the extensive sampling of nuclear loci in this study, exclusion of the mtDNA loci from the analysis did not significantly alter the evolutionary relationships recovered for any of the three species trees (Figs. 1, 2). The relationships of the four clades (A, B, C1, and C2) were all the same, and the only difference in the species relationships within the groups was in the relationships of some of the species in the C1 clade of the ML and BI tree.

We also assessed concordance of the different chromosomes of the nuclear genome. While the major groupings (A, B, C1, and C2 clades) were present in the species tree from each chromosome, the relationships among and within them varied. This incongruence among the different genomic regions is likely to be in part responsible for the low branch support values along the backbone of the phylogeny in the coalescent analysis. Our findings of the major species groups (Clades A, B, C1, and C2) and the species found within each group are consistent with other recent studies that used a smaller number of loci (Dyer et al., 2011b; Hatadani et al., 2009; Izumitani et al., 2016; Morales-Hojas and Vieira, 2012; Perlman et al., 2003). Thus, these clades likely represent natural groups. To recover the relationships among these groups it will be necessary to sample additional taxa from other species groups within the *immigrans-tripunctata* radiation (e.g., funebris, guarani, and others) and employ a larger number of markers.

Other studies have also found that the quinaria group is comprised of two divergent subgroups, and we cannot resolve whether these subgroups form a monophyletic group. The species trees from Müller B, C, and F supported the monophyly of the *quinaria* group (C1) and C2 clades), whereas for the remaining four genomic regions (Müller $A(X)$, D, and E and the Y-Chromosome), the quinaria group is paraphyletic. While regions of the genome that have reduced recombination (*i.e.* Müller F and the Y-chromosome) or are generally involved

in reproductive isolation (*i.e.* the sex chromosomes) may show reduced incomplete lineage sorting due to a smaller effective population size and/or reduced introgression in other species, these regions do not show any consistent pattern in our dataset. Given the large effective population sizes and potential for incomplete lineage sorting and hybridization, the resolution of the monophyly of the *quinaria* group will require larger-scale approaches than used here.

4.2 Patterns of reproductive isolation and divergence

Within the quinaria group, both of the C1 and C2 clades contain New and Old-World species. While Patterson and Stone (1952) lists instances of potential hybridization between C1 and C2 species, all of the cases we have been able to confirm in the laboratory are between species pairs in the C1 clade. Furthermore, within the C1 clade, species that hybridize generally do not co-occur, even if they are closely related (Coyne and Orr, 1989). For instance, *D. tenebrosa* is found only at high elevation in the mountains of the southeast United States and Mexico, and can hybridize with *D. suboccidentalis, D. recens*, and *D.* palustris, all of which occur only in northern North America (Dyer, unpublished). Likewise, D. suboccidentalis and D. occidentalis can produce fertile F1 hybrids but do not co-occur (Arthur and Dyer, 2015). The exception is D. recens and D. subquinaria, which co-occur in central Canada (Jaenike et al. 2006). These species along with D. transversa show incomplete reproductive isolation: hybrid sons of *D. recens* and either *D. subquinaria* or *D.* transversa are sterile, while hybrid sons and daughters between D. subquinaria and D. transversa are fully fertile (Humphreys et al., 2016; Shoemaker et al., 1999). The three species tree analyses all result in a different relationship of these species, which is likely due to recent hybridization and incomplete lineage sorting. Even the two geographic populations of D. subquinaria (coastal and inland) are not consistently monophyletic among analyses. Given the complex demography and incomplete isolation among these species, studies are underway that use more samples from each species, which may sort out some of these incongruences.

Within the *testacea* group, our results are consistent with previous studies in identifying that the North American species D. putrida is sister to the other three species in the group (*i.e.*, Perlman et al. 2003, Dyer et al. 2011b, Izumitani et al. 2016). This is also consistent with patterns of reproductive isolation, as *D. putrida* will not form hybrids with any of the other testacea group species and it is both ecologically and morphologically distinct from the others. Considering the other three species in this group, most of our analyses suggest that D. neotestacea and D. testacea are most closely related, with D. orientacea as the outgroup. However, this is contrary to patterns of reproductive isolation: D. orientacea and D. testacea can form fertile F1 female hybrids, whereas neither of these species form hybrids with D. neotestacea (Dyer et al., 2011b; Grimaldi et al., 1992). A further puzzle is that in the analyses that do not replicate the pattern of *D. orientacea* being the outgroup of this species trio, *D. orientacea* is found to be sister to *D. neotestacea* and not *D. testacea* (Müller C and ML of the full dataset, also Perlman et al. 2003), or the three species form a polytomy (Müller D and Y-chromosome). All three of these species are morphologically identical; D. neotestacea is found in the New World and the other two species occur in the Old World, where it is unknown whether their ranges overlap. Finally, little is known about the Asian

species *D. bizonata*, which is also now found in Hawaii (Leblanc et al., 2009), but it is morphologically very similar to the *testacea* group species. It is found to be sister to the testacea group in most of our analysis as well as in another recent analysis (Izumitani et al., 2016).

4.3 Evolution of feeding strategies

Species within the immigrans-tripunctata radiation feed on mushrooms, decaying vegetation, sap fluxes, fermenting fruits, flowers, or combinations of multiple host sources (Markow and O'Grady, 2008). We reconstructed the feeding history of the species we sampled, focusing on the *quinaria* and *testacea* groups. First, within the *quinaria* group, species are primarily mushroom feeders, but several have switched to utilizing decaying vegetation (Markow and O'Grady, 2008; Morales-Hojas and Vieira, 2012; Spicer and Jaenike, 1996). We found that there were likely two independent host-switching events within the C1 clade of the *quinaria* group from mushroom-feeding to vegetation-feeding (Fig. 3). It will be interesting to determine whether the preference for vegetation has the same or different genetic basis in these two lineages. We do not find a difference in the rate of protein evolution in the vegetative versus non-vegetative lineages (Fig. S4). However, it is interesting to note that period is both the fastest evolving locus in our dataset and the only locus to have a consistently higher rate of protein evolution in vegetative versus non-vegetative species (Table S4), though it does not display evidence of positive selection (*i.e.* d_N/d_S is not greater than 1). This gene is involved in circadian rhythms and courtship behaviors (reviewed in Nitabach and Taghert (2008) and Tataroglu and Emery (2015)). Previous work in the quinaria group found that the vegetative species are active throughout the day, whereas the mushroom species tended to restrict their activity to the morning and evening (Simunovic and Jaenike, 2006), thus an intriguing possibility is that a shift in the selective pressures of circadian feeding behaviors is reflected in the molecular evolution of the per locus.

At a deeper phylogenetic scale, we found that the common ancestor of the *quinaria* and testacea groups was probably mushroom-feeding, though additional species group sampling within this radiation will be needed to confirm this finding. It will be interesting to determine if traits present in these groups associated with mushroom-feeding such as the ability to tolerate mushroom toxins also had a single origin. Finally, at the base of our phylogeny we found that the ancestral state of the immigrans-tripunctata radiation was probably fruit feeding. This seems likely given that species within the immigrans group and closely related groups outside of the radiation are primarily fruit feeders (Kimura, 1980; Mitsui et al., 2010; Morales-Hojas and Vieira, 2012). Within the *cardini, immigrans*, and tripunctata groups a variety of feeding strategies occur, many of which were not included in our study. For instance, some species only exhibit only the fruit feeding strategy, others are fruit or flower feeders $(e.g., guarani$ and *pallidipennis*), and others $(e.g., function)$ functions group) exhibit mushroom feeding but only consume polypore rather than basidiomycete mushrooms (Lacy, 1984; Markow and O'Grady, 2008; Morales-Hojas and Vieira, 2012). In general, the inclusion of additional taxa from these groups in future studies will help resolve the ancestral feeding strategy along the backbone of the phylogeny.

Our results represent the first well-sampled phylogeny of the *quinaria* and *testacea* groups of Drosophila. The findings from our different phylogenetic analyses highlight the importance of careful selection of molecular markers and types of analyses employed, particularly in recent radiations. This study also provides a framework phylogeny for the *immigrans*tripunctata radiation that can be used to guide the taxon selection for future studies. The inclusion of representatives of additional species groups and additional species from the immigrans, cardini, and tripunctata groups could help to resolve the deeper relationships within the radiation as well as whether the *quinaria* group should be split into two species groups. Given that we sampled almost all of the species currently placed within the testacea and *quinaria* groups, it is unlikely that additional taxon sampling will help to resolve relationships, particularly in subgroup C1. However, employing phylogenomic approaches may resolve some of the more recent nodes where reproductive isolation among species is not complete and they have a complex demographic history $(e.g.,)$ exhibit incomplete lineage sorting, hybridization, and/or a combination of both factors). Even with these uncertainties, we are able to infer that the evolution of mushroom-feeding in the *quinaria* and *testacea* groups likely occurred once, and that within the *quinaria* group there were probably two independent instances of host-switching from mushroom-feeding to vegetation feeding. Future studies will disentangle whether these host switches occurred via a similar genetic mechanism.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- **•** Phylogenetic analyses support monophyly of testacea group but not quinaria group
- **•** Phylogeny offers framework to guide future study of immigrans-tripunctata radiation
- **•** Ancestor of the tripunctata and quinaria groups exhibited mushroom-feeding
- Two transitions to vegetative feeding occurred in the *quinaria* group
- **•** No change in rate of protein evolution associated with switch to vegetative feeding

Figure 1.

Comparison of phylogenetic relationships within the quinaria and testacea groups using the entire dataset of 43 loci. (A) Analyses of the concatenated dataset using maximum likelihood. Shown is the most likely cladogram, with a $LnL = -111850.75$. BS \pm 50 is given above branches; (B) Analyses of the concatenated dataset using Bayesian inference. Shown is the most probable topology, with a Harmonic Mean = −108525.50. PP is given above the branches; (C) The species tree resulting from the ASTRAL analysis of all 43 gene trees, with BS $\,$ 50 given above branches. All analyses were rooted with *D. immigrans*. Clade A

represents the tripunctata and cardini group species; Clade B represents the testacea and bizonata group species, and Clade C represents the quinaria group species, with C1 and C2 representing the two divergent clades within this group. Within D. subquinaria, the Coastal and Inland samples are designated by C and I, respectively.

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Figure 2.

Phylogenetic relationships within the *quinaria* and *testacea* groups resulting from the analyses of 40 nuclear loci. BS 50 are given above branches. (A) Analyses of the concatenated dataset using maximum likelihood. Shown is the most likely cladogram, with a LnL = -102720.99 . BS $\,$ 50 is given above branches; (B) Analyses of the concatenated dataset using Bayesian inference. Shown is the most probable topology, with a Harmonic Mean = −99861.12. PP is given above the branches; (C) The species tree resulting from the ASTRAL analysis of all 40 gene trees, with BS $\,$ 50 given above branches. As in Fig. 1,

Clade A represents the tripunctata and cardini group species; Clade B represents the testacea and bizonata group species, and Clade C represents the quinaria group, with C1 and C2 representing the two divergent clades within this group. Within D. subquinaria, the Coastal and Inland samples are designated by C and I, respectively.

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Figure 3.

Evolutionary hypothesis of feeding behavior using species trees reconstructed from the 40 nuclear loci. (A) Feeding behavior reconstruction based on ML topology; (B) Feeding behavior reconstruction based on the ASTRAL species tree. The most likely ancestral feeding behavior reconstruction and alternative reconstructions with a likelihood > 5% is indicated on each branch. The probability (P) of the most likely ancestral state is provided for each node. Clade A represents the tripunctata and cardini group species; Clade B represents the testacea and bizonata group species, and Clade C represents a monophyletic quinaria group, with C1 and C2 representing the two divergent clades within this group; * represents Node 1, the ancestor of the immigrans-tripunctata radiation; ** represents Node 2, the ancestor of the ingroup taxa. Within D. subquinaria, the Coastal and Inland samples are designated by C and I, respectively.

Table 1.

Species included in this study.

NOTE – The species representing each of the species groups included in the analysis are listed. In addition, the collection location, Wolbachia infection status, and potential to hybridize are provided for each species. M: Mushrooms, V: Decaying vegetation, F: Fermenting fruit; Geographic Range: Asian (A), North America (NA), Europe (E).

 a Tuno *et al.* (2007);

b Colon-Parrilla & Perez-Chiesa (1999);

 c Stump *et al.* (2011);

 d Werner & Jaenike (2017);

e
Kimura *et al.* (1977);

 $f_{\mbox{\scriptsize Werren}}$ & Jaenike (1995)

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