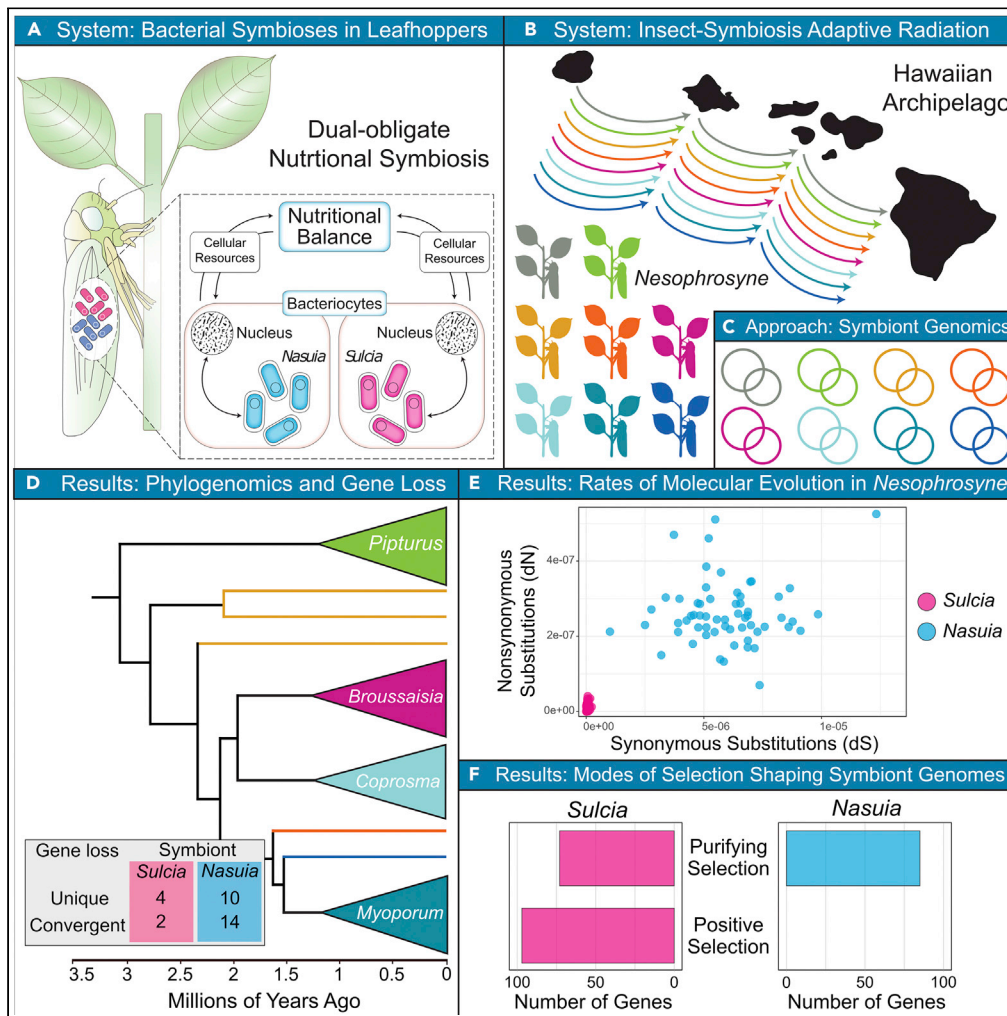


Article

A complex interplay of evolutionary forces continues to shape ancient co-occurring symbiont genomes



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Highlights

Ancient, reduced symbiont genomes are evolutionarily dynamic

Leafhopper symbionts have two of the slowest and fastest molecular evolution rates

Shared symbionts exhibit divergent and convergent gene losses among related hosts

Multiple symbionts in the same host experience different evolutionary pressures



Article

A complex interplay of evolutionary forces continues to shape ancient co-occurring symbiont genomes

Yumary M. Vasquez^{1,*} and Gordon M. Bennett^{1,2,*}

SUMMARY

Many insects depend on ancient associations with intracellular bacteria for essential nutrition. The genomes of these bacteria are often highly reduced. Although drift is a major driver of symbiont evolution, other evolutionary forces continue to influence them. To understand how ongoing molecular evolution and gene loss shape symbiont genomes, we sequenced two of the most ancient symbionts known, *Sulcia* and *Nasuia*, from 20 Hawaiian *Nesophrosyne* leafhoppers. We leveraged the parallel divergence of *Nesophrosyne* lineages throughout Hawaii as a natural experimental framework. *Sulcia* and *Nasuia* experience ongoing—but divergent—gene loss, often in a convergent fashion. Although some genes are under relaxed selection, purifying and positive selection are also important drivers of genome evolution, particularly in maintaining certain nutritional and cellular functions. Our results further demonstrate that symbionts experience dramatically different evolutionary environments, as evidenced by the finding that *Sulcia* and *Nasuia* have one of the slowest and fastest rates of molecular evolution known.

INTRODUCTION

Bacterial symbionts have enabled many animal groups to take advantage of unsuitable ecological niches leading to their biological diversification (Takiya et al., 2006; Moran, 2007; Sudakaran et al., 2015; Hendry et al., 2016; Sogin et al., 2020; Myers et al., 2021). In insects, bacterial endosymbionts are a key source of essential nutrition for many species, and even entire orders, that specialize in diets limited in essential nutritional resources (Moran, 1996; Baumann, 2005; Douglas, 2009). These symbionts are generally restricted to within specialized insect organs (bacteriomes) and cells (bacteriocytes) that enable host-bacterial interaction and strict vertical transmission from mother to offspring (Buchner, 1965; Koga et al., 2012). However, owing to their intracellular lifestyle, bacteria often lose over 90% of their genes (McCutcheon and Moran, 2012; McCutcheon et al., 2019). As symbioses age, bacteria continue to experience ongoing gene losses from their most basic—and essential—cellular processes and metabolisms (e.g., DNA replication and repair). Thus, endosymbionts require extensive resources from their hosts and other bacterial partners to function (Nakabachi et al., 2005; Hansen and Moran, 2011; McCutcheon and von Dohlen, 2011; Sloan et al., 2014; Luan et al., 2015; Mao et al., 2018; Weglarz et al., 2018; Kobiakka et al., 2018). Although we have a good picture of how symbiont genomes shrink on the scale of 10–100s millions years, we understand comparatively little about how this process continues to shape the tiny genomes of ancient symbionts among closely related host species (Wernegreen, 2002; McCutcheon et al., 2009a, 2009b; Moran et al., 2009; Mao et al., 2017; Chong et al., 2019).

In many cases, our understanding and inference of the evolutionary processes that influence the structure and function of symbiont genomes are derived from a single, or just a few, representative genomes (McCutcheon et al., 2009a, 2009b; Bennett and Moran, 2013; Koga and Moran, 2014; Bennett and Mao, 2018; Michalik et al., 2021). These processes include selection to retain essential functions, selection to adapt to changing host and environmental conditions, and strong genetic drift owing to small population sizes and strong intergenerational bottlenecks (Moran, 1996; Wernegreen, 2002; Woolfit and Bromham, 2003; Campbell et al., 2015; Perreau et al., 2021). These processes are ongoing among related symbiont lineages that are separated into distinct host species and their populations. The intensity of these forces may vary depending on the bacterial symbiotic roles and cellular environments, as well as the biology

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and ecology of their different host insect species (Wernegreen, 2002; Sabater-Muñoz et al., 2017; Chong et al., 2019). As a result, there is likely to be tremendous variation among the symbiont genomes of closely related host species. For example, the process of drift is known to cause independent symbiont lineages to differ widely in their genetic capabilities even between closely related host species (McCutcheon and Moran, 2010; Patiño-Navarrete et al., 2013; Husnik et al., 2013; Campbell et al., 2015, 2017; Bennett et al., 2016b; Husnik and McCutcheon, 2016; Boscaro et al., 2017; Łukasik et al., 2018; Chong et al., 2019; Monnin et al., 2020; Santos-Garcia et al., 2020). It is less clear how drift and selection work together to shape and maintain the genes and functions of symbionts as they diversify along with their hosts. Thus, an investigation into the patterns of gene loss and molecular evolution (e.g., the relative roles of selection vs. drift in gene evolution) among closely related host sister species has the potential to illuminate the fine-scale evolutionary processes that underlie ongoing symbiont genome evolution and diversification.

To better understand the evolutionary processes that shape ancient symbiont genome evolution, we analyzed the genomes of “*Candidatus Sulcia muelleri*” (*Bacteroidetes*; hereafter referred to as *Sulcia*) and “*Ca. Nasuia deltocephalinicola*” (*Betaproteobacteria*; hereafter *Nasuia*) lineages from insect sister species belonging to the endemic Hawaiian leafhopper genus, *Nesophrosyne* (Hemiptera: Auchenorrhyncha: Cicadellidae). *Nesophrosyne* is one of the largest insect adaptive radiations in Hawaii (Hembry et al., 2021). It originated ~3.2 million years ago, rapidly diversifying into over 200 species that specialize in endemic plant species in a one-to-one fashion (Zimmerman, 1948; Bennett and O’Grady, 2012, 2013). *Nesophrosyne* lineages established these host-plant relationships early in their diversification on the archipelago. These lineages then maintained their host-plant associations as they diversified across newly formed islands in a parallel and replicated fashion (Bennett and O’Grady, 2013). Thus, *Nesophrosyne* can provide a natural evolutionary experiment to test questions of how evolutionary processes shape symbiont genome evolution across diverging insect lineages and to further understand to what extent evolutionary processes are predictable.

Sulcia and *Nasuia*, like most other symbionts in the Auchenorrhyncha suborder, complement each other to provide their hosts with the 10 essential amino acids (EAAs) that are lacking in their xylem and phloem plant sap diets (Bennett and Moran, 2013). Genomic evidence suggests that both bacteria are ancient, having partnered with insects ~300 million years ago (Moran et al., 2005; Bennett and Mao, 2018). As a result, *Sulcia* and *Nasuia* have highly reduced genomes of 190 kilobases (kb) and 112 kb, respectively (Bennett and Moran, 2013). Both genomes maintain a core set of essential nutritional genes, but are lacking genes in most other essential functions that include translation and transcription, energy synthesis, and DNA replication and repair (Bennett and Moran, 2013; Bennett et al., 2014; Mao et al., 2017).

Here, we leverage the *Nesophrosyne* leafhopper radiation and their endosymbionts to understand (i.) whether and how ongoing gene losses continue to shape ancient symbiont genomes, (ii.) how selection and drift lead to symbiont genome diversification, and (iii.) whether these evolutionary forces are shared or distinct between multiple symbionts in a shared host. We further test our questions in absolute time provided by the *Nesophrosyne* radiation to gain general insights into the tempo and mode of bacterial symbiont genome evolution.

RESULTS AND DISCUSSION

Host-symbiont taxon sampling and nomenclature for genomic analysis

To compare the genomes of symbionts associated with *Nesophrosyne*, we strategically sub-sampled 20 species that span the ecological and phylogenetic diversity of the leafhopper genus. Our selected species comprise monophyletic groups that specialize in eight distinct endemic Hawaiian plants, encompassing the diversity of host-plant associations known in the *Nesophrosyne* (Bennett and O’Grady, 2012; see Figures 1 and 2). The sister species of each host lineage diversified across the Hawaiian Archipelago, with distinct species occurring on each island and even on a volcanic mountain (Bennett and O’Grady, 2013).

Sulcia and *Nasuia* strains are hereafter identified by the island and host plant genera associated with the leafhopper host as follows: the initial for the island location (e.g., Hawai’i Island = HI) and the first two letters of the plant genus (e.g., *Pipturus* = PI). For example, the *Sulcia* strain associated with the *Nesophrosyne* species restricted to Hawai’i Island and the plant genus *Pipturus* is referred to as *Sulcia-HIPI* (Table 1).

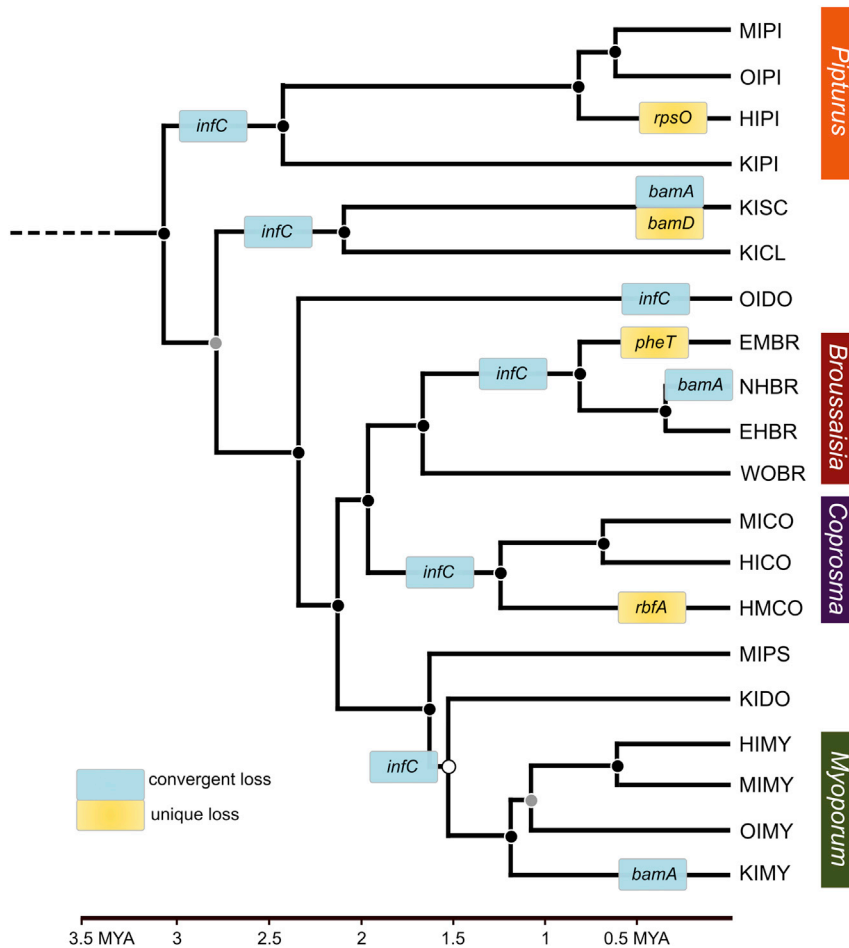


Figure 1. Convergent and unique gene loss among *Sulcia* genomes from endemic Hawaiian leafhoppers (*Nesophrosyne*)

Patterns of gene loss were estimated with maximum likelihood ancestral state reconstruction on an absolute time-calibrated phylogenetic tree in millions of years, using complete host mitochondrial genes (see [Supplemental information](#)). Convergent gene losses (*i.e.*, multiple repeated losses) are indicated by blue boxes. Unique gene losses (*i.e.*, losses that occurred once) are indicated by yellow boxes. Shorthand gene names are provided in each box. Posterior support for each node is shown as colored circles as follows: black >95, gray = 90-95, and white <90. See section methods for details on tree search parameters. See [Table 1](#) for species shorthand nomenclature. MYA = million years ago.

Phylogenetic relationships among *Nesophrosyne* and their symbionts are congruent

To test our questions in absolute time, we generated a time-calibrated phylogeny from complete *Nesophrosyne* mitochondrial genomes (15 genes, 14,304 sites) using absolute time calibration points determined previously by [Bennett and O’Grady \(2013\)](#). To compare phylogenetic topologies of the host and symbionts, we also reconstructed phylogenies for *Nasuia* (99 genes, 86,095 sites) and *Sulcia* (184 genes, 181,781 sites) independently using their complete genomes. The relationships among our sub-sampled host species agree with previous phylogenetic work and there is strong support for the monophyly of leafhoppers associated with their host-plant groups ([Bennett and O’Grady, 2012, 2013](#); see [Figure S1](#)). However, two *Nesophrosyne* species are weakly supported in their placements and vary between trees derived from the symbiont and mitochondrial datasets (KIDO and MIPS; see [Figures S1–S3](#)). Previous work also observed a similar mid-depth polytomy, suggesting a rapid early diversification in the *Nesophrosyne* that is unable to be split by available genetic and genomic data (*i.e.*, a hard polytomy; see [Bennett and O’Grady 2012](#)), or the possible introgression of symbiont lineages between hybridizing hosts. Hybridization events are known to occur among rapidly diversifying, host-plant restricted auchenorrhynchan lineages in Hawai’i ([Roesch Goodman et al., 2012](#)). However, we do not currently have a suitable sampling of host species and their populations to thoroughly test this hypothesis in the *Nesophrosyne*.

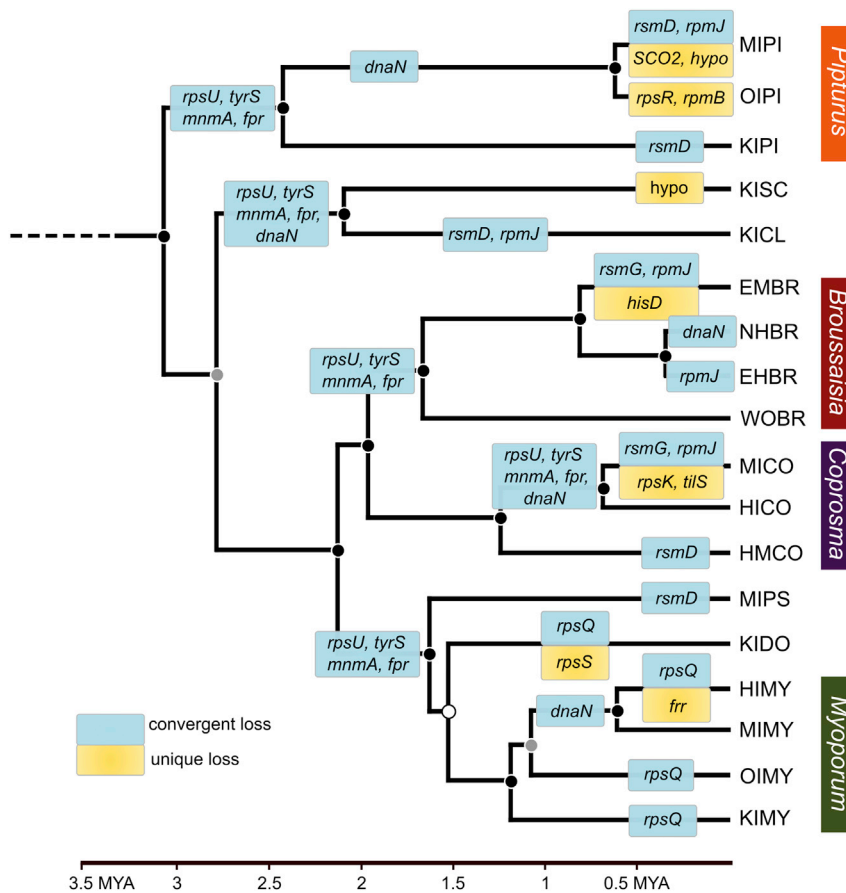


Figure 2. Convergent and unique gene loss among *Nasuia* genomes from endemic Hawaiian leafhoppers (*Nesophrosyne*)

Patterns of gene loss were estimated with maximum likelihood ancestral state reconstruction on an absolute time-calibrated phylogenetic tree in millions of years, using complete host mitochondrial genes (see [Supplemental information](#)). Convergent gene losses (i.e., multiple repeated losses) are indicated by blue boxes. Unique gene losses (i.e., losses that occurred once) are indicated by yellow boxes. Shorthand gene names are provided in each box. Posterior probability support for each node is shown as colored circles as follows: black >95, gray = 90-95, and white <90. See section methods for details on tree search parameters. See [Table 1](#) for species shorthand nomenclature. MYA = million years ago. hypo = hypothetical protein.

Gene content of *Nesophrosyne*'s symbiont genomes varies despite their highly reduced size

From the Illumina sequenced *Nesophrosyne* host species, we recovered 20 *Sulcia* and 18 *Nasuia* genomes (see [Table S1](#)). All symbiont genomes are complete and circular. Even coverage across all circularized genomes was verified with read mapping that verified complete, high-quality assemblies (see Methods). Two *Nasuia* genomes (*Nasuia*-HIPI and *Nasuia*-OIDO) were omitted because sequencing coverage was too low to assemble reliable contigs and complete genomes required for our downstream molecular assays.

The average genome size of *Sulcia* is 190 kilobases (kb; range = 190.3-190.9 kb) with an average of 190 protein-coding genes (range = 188-192 genes). These genomes further retain a single conserved 16S/23S/5S rRNA operon and 30 tRNAs. In contrast, *Nasuia* exhibits more variation between host species. Its average genome size is 112 kb (range = 107.7-116.1 kb) with an average of 132 protein-coding genes (range = 125-139 genes), a single 16S/23S/5S rRNA operon, and 18-21 tRNAs (see [Table S1](#)). *Nesophrosyne*'s *Sulcia* genomes are highly conserved, varying by up to six genes in the most extreme cases (~3% of its genome). In contrast, *Nasuia*'s genome is highly variable among the *Nesophrosyne*, differing by up to 24 genes (>20% of its genome).

Table 1. Shorthand nomenclature used for bacterial symbiont strains

Host species	Island location	Habitat type	Host-plant group	Shorthand naming
<i>N. sp. 295</i>	East Hawai'i Island	Rain Forest	<i>Broussaisia</i>	EHBR
<i>N. ogradyi</i>	East Maui Island			EMBR
<i>N. kanawao</i>	North Hawai'i Island			NHBR
<i>N. makaihe</i>	West O'ahu Island			WOBR
<i>N. sp. 23</i>	Kaua'i Island	Rain Forest	<i>Clermontia</i>	KICL
<i>N. haleakala</i>	Haleakalā Mtn, Maui	Spanning all habitats	<i>Coprosma</i>	HMCO
<i>N. sp. 302</i>	Hawai'i Island			HICO
<i>N. sp. 58</i>	Maui Island			MICO
<i>N. sp. 29</i>	Kaua'i Island	Dry-Mesic	<i>Dodonea</i>	KIDO
<i>N. maratima</i>	O'ahu Island			OIDO
<i>N. sp. 281</i>	Hawai'i Island	Spanning all habitats	<i>Myoporum</i>	HIMY
<i>N. sp. 126</i>	Kaua'i Island			KIMY
<i>N. sp. 242</i>	Moloka'i Island			MIMY
<i>N. sp. 246</i>	O'ahu Island			OIMY
<i>N. montium</i>	Hawai'i Island	Rain Forest	<i>Pipturus</i>	HIPI
<i>N. sp. 17</i>	Kaua'i Island			KIPI
<i>N. sp. 48</i>	Maui Island			MIPI
<i>N. ponapona</i>	O'ahu Island			OIPI
<i>N. sp. 15</i>	Moloka'i Island	Dry-Mesic	<i>Psychotria</i>	MIPS
<i>N. sp. 21</i>	Kaua'i Island	Spanning all habitats	<i>Scaevola</i>	KISC

Shorthand naming are the first letters of the island location and the first two letters of the host-plant group. Habitat type of each host-plant group is added as well. (*N.* = *Nesophrosyne*, *sp.* = species).

Globally, patterns of genome evolution and gene retention among *Sulcia* and *Nasuia* are similar to patterns observed in other Auchenorrhyncha lineages. Both retain complementary essential amino acid (EAA) pathways in an 8 + 2 arrangement for *Sulcia* and *Nasuia*, respectively, as observed in other leafhoppers and related insects (Chang et al., 2015; Bennett et al., 2016a; Mao et al., 2017; see also Bennett and Moran, 2013 for a list and pathways in *Nasuia* and *Sulcia*). The highly conserved nature of *Sulcia*'s genome has been widely observed across the other major auchenorrhynchan lineages that retain it (e.g., sharpshooter leafhoppers, cicadas, spittlebugs; McCutcheon and Moran, 2007, 2010; Koga and Moran, 2014; Campbell et al., 2015; Łukasik et al., 2018; Matsuura et al., 2018). The most dramatic differences observed among *Sulcia* genomes occur between the major infraorders Fulgomorpha (planthoppers) and Cicadomorpha (leafhoppers and kin). Among the planthoppers, *Sulcia* genomes are much smaller than in the cicadomorph lineages (<149 kb in Fulgomorpha vs. an avg. of 251 kb in Cicadomorpha [range = 179-288 kb]; Bennett and Mao, 2018; Michalik et al., 2021). Among other gene losses, planthopper *Sulcia* lineages provide only three of the seven to eight EAAs typically retained in strains found in cicadomorph hosts (McCutcheon and Moran, 2010; Michalik et al., 2021).

Nasuia's genomes, in contrast to *Sulcia*, exhibit more variation in the genes they retain. The number of gene losses involves >20% of *Nasuia*'s genome ($n = 24$ genes) among the *Nesophrosyne*. This variation is significant considering that *Nasuia*'s genome is among the smallest known of any bacterium. The loss of any single gene that *Nasuia* lineages still retain likely requires direct adaptation by the host, or its partner symbionts, to support lost genetic and functional capabilities (Moran and Bennett, 2014; McCutcheon et al., 2019). The diversity of co-primary symbionts associated with *Sulcia* also shows similarly higher genomic variation. For example, the cicada co-primary symbiont, "Ca. *Hodgkinia*" (hereafter *Hodgkinia*), has the most dramatic genomic variation among a symbiont yet observed. Although some cicada species harbor *Hodgkinia* with typical circular chromosomes (avg. size = 142 kb), in other hosts its genome is broken into mini circles of varying size and complexity (e.g., fragments range from 71 to 150 kb in 13-year *Magicalcidas*; Van Leuven et al., 2014; Campbell et al., 2015; Łukasik et al., 2018). Similarly, the lineages of the co-primary symbiont "Ca. *Baumannia*," which replaced *Nasuia* in sharpshooter leafhoppers >60 million years ago, vary

by the loss of large chunks of its genome spanning >100 kb and >100 protein-coding genes (Wu et al., 2006; Bennett et al., 2014, 2016b).

Ancestral gene losses are ongoing and to some extent evolutionarily convergent

To determine patterns of gene loss (*i.e.*, unique vs. convergent losses) among *Sulcia* and *Nasuia* genomes, we reconstructed ancestral patterns of gene loss and retention with maximum likelihood approaches (see Figures S4 and S5 and STAR Methods). We also used *Sulcia* and *Nasuia* lineages from two previously sequenced species from the Membracoidea superfamily, the aster leafhopper (*Macrostoteles quadrilineatus*) and the keeled treehopper (*Entylia carinata*), to determine ancestral patterns of gene loss leading to the *Nesophrosyne* lineage (Bennett and Moran, 2013; Mao et al., 2017). In general, the membracoidean *Sulcia* and *Nasuia* lineages are structurally conserved and perfectly syntenic, aside from differences in their patterns of individual gene losses (reviewed by Mao et al., 2017). Later in the discussion, we summarize gene losses in both symbionts. We caution that gene loss counts presented here are minimum counts, as we did not sequence the symbiont genomes from all of *Nesophrosyne*'s 200+ species.

Sulcia in the common ancestor to Membracoidea likely retained at least 210 protein-coding genes and 30 tRNAs (reviewed by Mao et al., 2017). Prior to the diversification of the Deltocephalinae leafhoppers (*Nesophrosyne*'s leafhopper subfamily), *Sulcia*'s genome was reduced to 192 genes and 30 tRNAs (Shcherbakov, 2002; Mao et al., 2017). Among the *Nesophrosyne*, it has further undergone at least six instances of gene loss (*infC*, *bamAD*, *rbfA*, *rpsO*, and *pheT*; see Figure 1). The loss of these genes impacts a range of key cellular functions in *Sulcia*, which have significant implications for how the symbiosis functions and is maintained (Wilson and Duncan, 2015; Bublitz et al., 2019; Mao and Bennett, 2020). Some of these gene losses appear to be unique independent events among individual species, while others have been convergently lost multiple times during *Nesophrosyne* diversification.

Two genes removed from *Sulcia*'s genomes were lost multiple times in convergent evolutionary events: *infC* and *bamA* (Figure 1). The outer membrane protein assembly factor gene (*bamA*) was lost at least three independent times among our sampled *Nesophrosyne* species (Figure 1). It is part of a multi-gene complex essential for bacterial outer membrane assembly and metabolite exchange (Malinverni et al., 2006; Charles et al., 2011). Interestingly, the loss of *bamA* in *Sulcia*-KISC co-occurs with the loss of its interacting partner protein (*bamD*) and may be linked (Wu et al., 2005). Thus, some gene losses may instigate the loss of others in a domino-like fashion, as has been proposed to occur in the *Blattabacterium*-cockroach system (Kinjo et al., 2021). Perhaps more remarkable is the loss of the translation initiation factor IF-3 gene (*infC*) at least six times during the diversification of *Nesophrosyne* (see Figure S4). The *infC* gene is part of a three protein complex involved in translation initiation (Sabol et al., 1970). All *Nesophrosyne Sulcia* lineages still retain the other two *infAB* genes, suggesting support from partner symbionts or that they have moonlighting functions. The *infABC* gene set is generally retained in most other bacterial symbiont genomes across the Hemiptera, indicating that the gene has an essential functional role (Nakabachi et al., 2006; McCutcheon et al., 2009a, 2009b; McCutcheon and Moran, 2010; McCutcheon and von Dohlen, 2011).

Nasuia in the common ancestor to Membracoidea retained at least 163 genes and 29 tRNAs (Mao et al., 2017). Early on in the divergence of the Deltocephalinae leafhoppers, *Nasuia*'s genome was further reduced to a mere ~142 genes and just over 112 kb in size (Bennett and Moran, 2013). Across the *Nesophrosyne*, *Nasuia* has undergone at least 24 instances of gene loss (>20% of its genome). Ten of these genes have been lost once among our sampled *Nesophrosyne* host species, including genes involved in ribosome function (*rpsKRS* and *rpmB*), tRNA synthesis (*tilS* in *Nasuia*-MICO), and histidine synthesis (*hisD* from *Nasuia*-EMBR). The latter two genes are essential for EAA and general protein synthesis (Soma et al., 2003; Van Leuven et al., 2019). The concentrated losses of ribosome-associated genes (8 out of 24) suggest that the host either easily replaces them, or the ribosomal holoenzyme can adapt to their absence (Akanuma et al., 2012; Galperin et al., 2021; Nikolaeva et al., 2021).

More than half of gene losses in *Nasuia* ($n = 14$) have been convergently lost in at least two or more host insect species (Figure 2). The most extreme case of convergent loss is that of a gene cassette that includes four complete genes: 30S ribosomal subunit protein S21 (*rpsU*), tyrosine-tRNA ligase (*tyrS*), tRNA-specific 2-thiouridylase (*mnmA*), and flavodoxin/ferredoxin-NADP+ reductase (*fpr*) (see ancestral state reconstructions in Figure 3B). Additionally, an essential component of the DNA replication holoenzyme, *dnaN*, has been lost at least five times and is missing from more than half of our *Nasuia* genomes (Johanson and

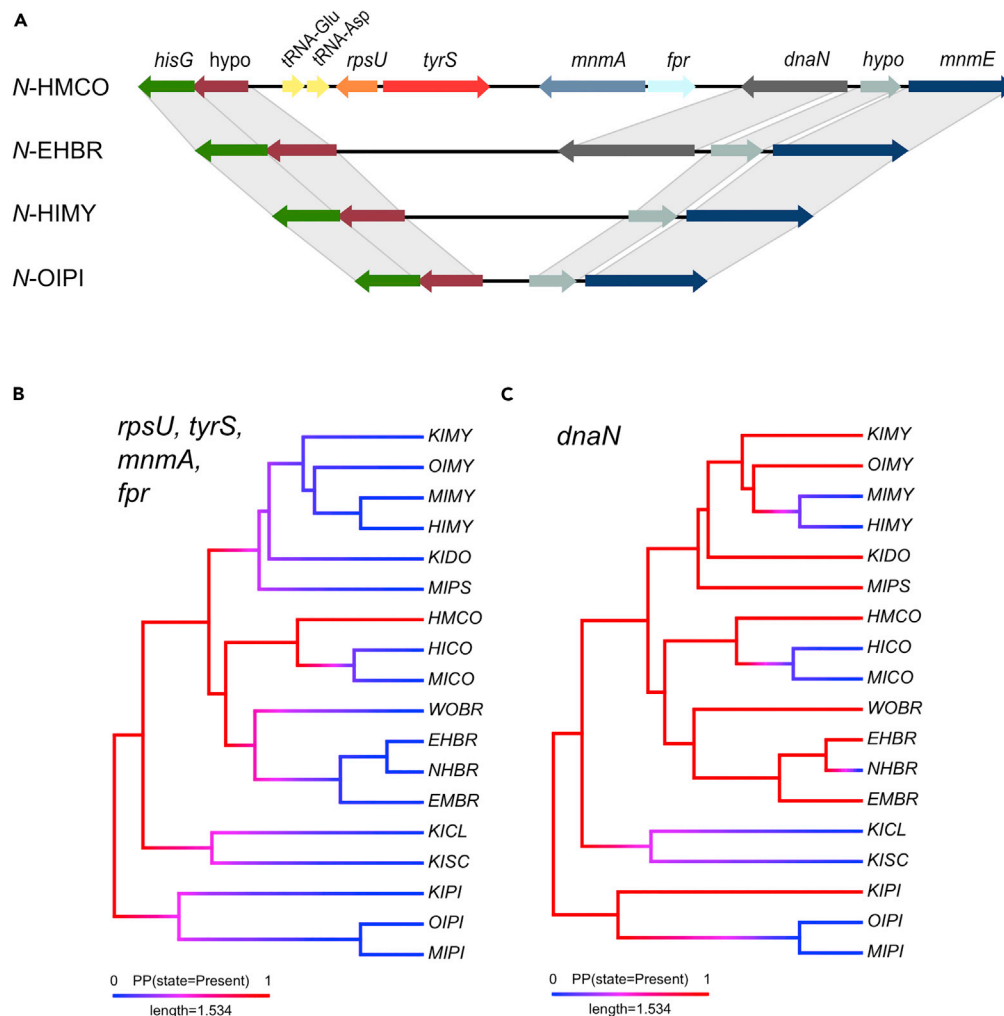


Figure 3. *Nasuia* gene losses of *rpsU*, *tyrS*, *mnmA*, *fpr* and *dnaN* cassette

(A) *Nasuia* gene losses of *rpsU*, *tyrS*, *mnmA*, *fpr*, and *dnaN* cassette for three representative genomes, as well as the genes maintained in *Nasuia*-HMCO (N-HMCO) sequence. The genes *rpsU*, *tyrS*, *mnmA*, and *fpr* have been lost in all genomes except N-HMCO (top). The *dnaN* gene has been convergently lost in nine sequences. Ancestral state reconstruction of *rpsU*, *tyrS*, *mnmA* and *fpr* (B) suggests that these genes have been lost at least five times. Ancestral state reconstruction of *dnaN* (C) suggests that this gene has been lost at least five times. Colors correspond to the same gene in each genome segment (hypo = hypothetical protein). (B and C) Ancestral state reconstruction of *rpsU*, *tyrS*, *mnmA*, and *fpr*, as well as *dnaN* from phyttools v.1.0-1 package on genes convergently lost in *Nasuia* (Revell, 2012). The loss of *rpsU*, *tyrS*, *mnmA*, and *fpr* may have instigated the loss of *dnaN*. See section methods for details on ancestral state reconstruction.

McHenry, 1980; Figure 3C). Finally, several ribosomal proteins (*rsmD*, *rpmJG*, and *rpsQ*) have been convergently lost across our sampled *Nasuia* species.

The ongoing loss of more than 20% of *Nasuia*'s genes among these lineages similarly presents major challenges to its hosts and partner symbionts, particularly because several of these genes are essential to its nutritional role in the symbiosis and its cellular functions (e.g., *hisD* and *tilS*, respectively). Although the number of genes *Nasuia* is capable of losing stands in stark contrast to *Sulcia*, both symbionts require the host or companion symbionts to adapt stabilizing support mechanisms (Mao et al., 2018). However, *Nasuia* is apparently a more demanding partner requiring independent host lineages to innovate novel support strategies. The reason for *Nasuia*'s more exaggerated rates of gene losses over *Sulcia* is likely owing to its rapid rates of molecular evolution discussed below (Bourguignon et al., 2020).

***Sulcia* and *Nasuia* have among the slowest and fastest rates of symbiont molecular evolution**

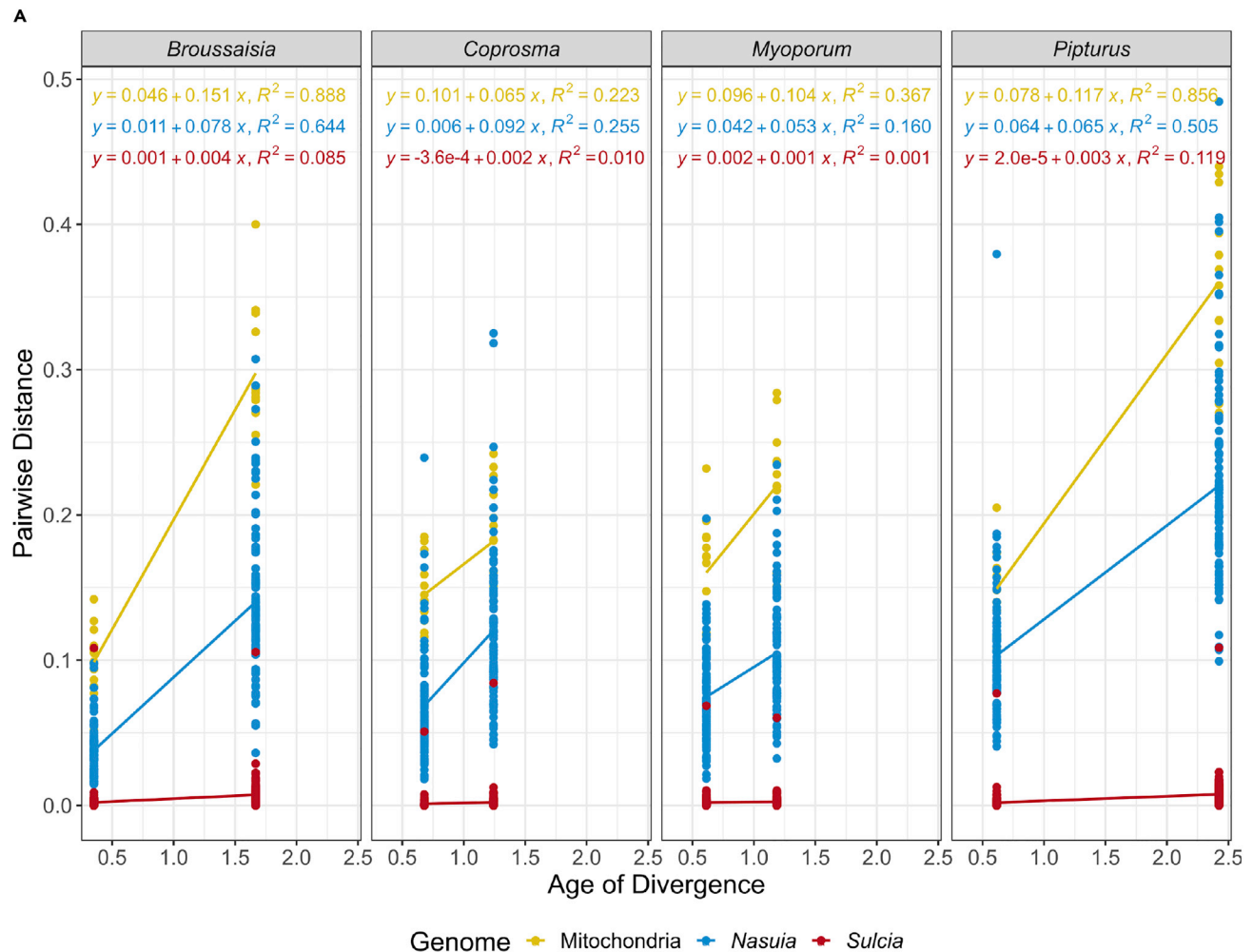
To determine the underlying drivers of symbiont genome evolution among *Nesophrosyne*, we estimated genome-wide substitution rates in absolute time for *Sulcia* and *Nasuia*. We further compared these against host mitochondrial rates (Figure 4). The average substitution rates for *Sulcia* genomes are 3.24×10^{-9} substitutions/site/year. In contrast, *Nasuia* genomes exhibit a 34.8-fold higher rate of molecular evolution (avg. = 1.13×10^{-7} substitutions/site/year). The evolutionary rates of both symbiont genomes do not exceed that of the mitochondria (avg. = 2.15×10^{-7} substitutions/site/year: 1.9-fold from *Nasuia* and 66-fold from *Sulcia*). *Nesophrosyne* mitochondrial rates of evolution are in-line with observations from other hemipteran insects, which are generally elevated relative to other insects (Dowton et al., 2009; Song et al., 2012; Cui et al., 2013; Cameron, 2014; Li et al., 2017).

To compare the rate of molecular evolution in *Sulcia* and *Nasuia* to available symbionts from other insect hosts, we estimated nonsynonymous (dN) and synonymous (dS) substitutions over their divergence times. We used the general M0 model, which averages substitution rates across whole genes and phylogeny. We further converted rates to dN/time (dN/t) and dS/time (dS/t) (Yang 2007; Silva and Santos-Garcia, 2015). The rates for *Sulcia* among the *Nesophrosyne* are 9.46×10^{-9} dN/t and 4.21×10^{-8} dS/t (time = 3.2 MYA). In contrast, *Nasuia*'s rates are highly elevated, averaging 1.65×10^{-7} for dN/t and 3.76×10^{-6} for dS/t. The average dN/t in *Nasuia* is 17.5-fold higher than in *Sulcia*, while the differences in average dS/t are even higher (89.3-fold).

Compared to other insect symbionts, for which data are available, *Nasuia* has among the highest evolutionary rates yet identified. The well-known symbiont of aphids, *Buchnera*, has an average rate of 2.58×10^{-9} dN/t and 1.43×10^{-8} dS/t, which is 64.2 and 262-fold less than *Nasuia*'s rates, respectively (using 20 *Buchnera* protein coding genes; Clark et al., 1999). Several of the highest rates of molecular evolution previously documented for insect symbionts belong to *Baumannia*, which replaced *Nasuia* in sharpshooter leafhoppers >60 million years ago, and *Blochmannia* found in carpenter ants (Silva and Santos-Garcia, 2015). *Nasuia* exceeds these, with 42 and 341-fold higher for dS/t and 20 and 165-fold higher for dN/t than *Blochmannia* and *Baumannia*, respectively.

One factor that may influence the rates of evolution in *Nasuia* and *Sulcia*, as well as other symbionts more broadly, is differences in host and symbiont generation times (e.g., bacteria with shorter, more frequent generations can incur more mutations and substitutions per some unit of time; Degnan et al., 2005; Silva and Santos-Garcia, 2015). However, we do not have a clear understanding of generation times in our insects for several reasons. First, they are difficult to rear owing to their highly restricted habitat ranges and species further experience differences in rain-fall-associated seasonality (Bennett and O'Grady, 2012; see also Degnan et al., 2005). In addition, it is not known whether symbiont replication rates are even coupled with host generations in our system, nor among most other insect symbioses. It is, however, worth noting that *Sulcia* and *Nasuia* exhibit dramatically different rates of evolution despite sharing the same host lineages. Thus, host generation time alone cannot explain observed differences.

In contrast to *Nasuia*'s highly elevated rates of molecular evolution, *Sulcia* has one of the most depressed rates of any biological system (McCutcheon et al., 2009a, 2009b; Bennett et al., 2014; see Figure 4). It has been widely observed among the Auchenorrhyncha (e.g., in cicadas and spittlebugs), that *Sulcia* has a nearly inert rate of molecular evolution even across divergences spanning 100s of millions of years (Takiya et al., 2006; Koga et al., 2013; Bennett et al., 2014; Bennett and Mao, 2018; Waneka et al., 2021; Arab and Lo, 2021; Michalik et al., 2021). *Sulcia*'s depressed evolutionary rates are an enigmatic biological phenomenon. One possible explanation for *Sulcia*'s reduced rates of molecular evolution may be its retention of mutation repair systems. In a *Macrosteles* leafhopper, *Sulcia* was found to have an overall low rate of mutagenesis compared to *Nasuia*, possibly owing to its retention of the DNA mismatch repair protein, *mutS* (Waneka et al., 2021). The *mutS* gene recognizes and initiates the repair of mismatched bases and small indels, which can lower substitution rates (Dettman et al., 2016; Long et al., 2018; Waneka et al., 2021). Most *Sulcia* lineages in the Auchenorrhyncha still retain *mutS*. This gene retention pattern may explain *Sulcia*'s universally conserved rates of molecular evolution (McCutcheon and Moran, 2007, 2010; Woyke et al., 2010; Bennett and Moran, 2013; Koga and Moran, 2014; Bennett and Mao, 2018). In contrast, the *mutS* gene is widely lost from most of *Sulcia*'s co-primary symbiont partners, which have characteristically elevated rates of molecular evolution—with the exception of "*Candidatus Zinderia insecticola*" in spittlebugs, which still maintains the gene (Takiya et al., 2006; McCutcheon and Moran, 2012; Bennett et al., 2014, 2016b; Campbell et al.,



B

	Summary of p-values				
	Across All Groups	Myoporum	Coprosma	Pipturus	Broussaisia
Mitochondria - Nasuia	<2e-16	4.20E-13	6.80E-11	2.80E-07	2.30E-14
Mitochondria - Sulcia	<2e-16	<2e-16	<2e-16	<2e-16	<2e-16
Sulcia - Nasuia	<2e-16	<2e-16	<2e-16	<2e-16	<2e-16

Figure 4. Summary of uncorrelated rates of evolution between symbiont and host genes

(A) Linear regression between pairwise distances and age of divergence among all *Nasuia*, *Sulcia*, and mitochondrial genes for Hawaiian leafhoppers (*Nesophrosyne*). Colors indicate pairwise distance of protein-coding genes in each genome. Genomes are separated by the host-plant group that the host species has specialized in. A linear regression line is mapped between pairwise distance values from the closest related species to the most divergent species in each host-plant group. The regression equation and the coefficient of determination (R^2) are also reported and colored by the genome. Outliers with a pairwise distance >0.5 were removed (4 from *Nasuia*, 3 from Mitochondria).

(B) Associated p values of statistical tests between evolutionary rates of genomes across all plant groups and within plant groups. Significant p value between groups indicates no correlation between means of evolutionary rate.

2015; Arab and Lo, 2021). These higher rates may drive genomic volatility and variation among *Sulcia*'s partner symbionts, which are more frequently lost or replaced among the Auchenorrhyncha (Bennett and Moran, 2015; Sudakaran et al., 2017).

Rates of molecular evolution are uncorrelated between each symbiont

To further test whether *Sulcia* and *Nasuia* show correlated rates of evolution with each other and the host mitochondria, suggesting a shared evolutionary environment (Arab and Lo, 2021), we estimated substitution rates for both symbionts and mitochondria globally across and within the *Nesophrosyne*'s monophyletic host-plant associated clades (*Broussaisia*, *Coprosma*, *Pipturus*, and *Myoporum*; see Figure 4A). Correlations between substitutions rates among host and symbiont genomes could be explained if similar forces of selection are acting on the genomes, such as shared population bottlenecks and dependence on shared genes (e.g., host mutation repair genes taking over for those lost in symbiont genes; McCutcheon and Moran, 2012; Mao et al., 2018; McCutcheon et al., 2019). Overall, substitution rates between *Nesophrosyne*'s *Sulcia*, *Nasuia*, and the mitochondria are not correlated across host-plant affiliated clades (Figure 4B). Rates between symbionts and mitochondrial genes within host-plant clades are also uncorrelated (Figure 4B). Our results support a recent analysis of *Sulcia* from more widely divergent auchenorrhynchan clades that found similar decoupling of rates (see Arab and Lo, 2021). However, both of these findings are in contrast to mono-symbiont systems (*Blochmannia*-Carpenter Ants, *Blattabacterium*-Cockroaches, and *Buchnera*-Aphids) that tend to show a significant correlation between the rate of evolution in mitochondrial and symbiont genes (Degnan et al., 2004; Arab et al., 2020; Arab and Lo, 2021).

The disparity in rates of molecular evolution between *Sulcia* and its partners strongly suggests that different molecular and cellular processes, and evolutionary pressures, are likely shaping symbiont genomes (see Figure 4; Takiya et al., 2006; Bennett et al., 2014; Campbell et al., 2015; Arab and Lo, 2021). Although there is evidence that more ancient mitochondrial and plastid symbiont genomes do tend to show correlated rates of molecular evolution, these genomes are highly integrated into the general biology of most eukaryotic cellular and metabolic processes (Smith and Lee, 2010; Sloan et al., 2012; Hua et al., 2012). In contrast, the biological roles of insect symbionts are arguably less integrated into system-wide biological functions of their host insects. Nutritional symbionts are sequestered to distinct organs in the host, retaining relatively enriched genetic autonomy, distinct population sizes, and distinct cellular replication and repair capabilities (Buchner, 1965; Mira and Moran, 2002; Koga et al., 2012; Bennett et al., 2014; Chong and Moran, 2016; Mao and Bennett, 2020; Stever et al., 2021). As a result, they likely do not experience the same patterns and processes of molecular evolution as their partner symbionts, mitochondria, or host nuclear genes. Nevertheless, more fine-scale analyses that include a broad-scale sampling of host nuclear and symbiont genes, as well as focus on protein domains, may further find rate correlations on interacting genes.

Sulcia and *Nasuia* experience differential selection patterns across their genomes

To understand how different modes of selection are shaping the evolution and function of *Sulcia* and *Nasuia* genomes, we tested for selection both across genes and across sites using the ratio of nonsynonymous to synonymous substitutions (denoted by ω ; see STAR Methods). To determine whether symbiont genes are generally under strong purifying selection ($\omega < 0.1$), relaxed purifying selection ($0.95 < \omega > 0.1$), or positive selection ($\omega > 1$), we initially used the M0 model in codeml (ω estimated across the whole gene; Yang et al., 2000; Z. Yang 2007; Sloan and Moran, 2012; Sabater-Muñoz et al., 2017; Perreau et al., 2021). In *Sulcia*, a small subset of genes are undergoing strong purifying selection (avg. $\omega = 0.0615$ [range = 0.0001-0.0970, $n = 34$]), while most genes are undergoing relaxed purifying selection (avg. $\omega = 0.331$ [range = 0.112-0.931, $n = 133$]). Additionally, three genes in *Sulcia* show signatures of positive selection (*atpH*, *putA*, and *trpC*; $\omega > 1$). In *Nasuia*, most genes are undergoing strong purifying selection (avg. $\omega = 0.0449$ [range = 0.0162-0.0988, $n = 73$]), while comparatively few genes are experiencing relaxed purifying selection (avg. $\omega = 0.243$ [range = 0.105-0.555, $n = 12$]). This approach did not detect any *Nasuia* genes under positive selection.

Although it is useful to obtain an average ω value for a gene, it is not a sufficiently realistic model to detect signatures of positive selection that operate at finer scales (Anisimova et al., 2001). Specific codons related to intrinsic protein function may be under positive selection, while the majority of the gene can experience relaxed purifying selection (Yang et al., 2000; Anisimova et al., 2001). Thus, to test for positive selection on

different codon sites within symbiont genes, we used two nested models: M1a-M2a and M7-M8 in codeml (Yang 2007). We interpret consistent results between the two models (and also the M0 model from above) as strong global support for positive selection operating on sites within a gene (Anisimova et al., 2001; Padhi et al., 2009; Price et al., 2011; Alves et al., 2013). We applied this approach to *Sulcia* and *Nasuia* across our sampled *Nesophrosyne*.

Sulcia site-based selection analyses recovered the same genes undergoing positive selection from the M0 model, as well as an additional 94 genes (97 in total, $p < 0.05$ with BH correction). Genes under positive selection are functionally enriched for the Clusters of Orthologous Groups (COG) Translation (J; 31 genes; Fisher exact test, $p = 0.0008$; Figure 5) and Energy Production and Conversion (C; 14 genes; $p = 0.0486$; Figure 5). It is notable that, although not significant, 27 genes under positive selection are involved in essential amino acid synthesis (E; $p = 0.1030$; Figure 5).

Sulcia's genes under positive selection are primarily involved in essential amino acid synthesis, buffering degraded protein function, and other cellular processes. Some examples of proteins with positive sites include protein chaperonins (*groEL* and *dnaK*), EAA metabolite synthesis (e.g., arginine [*argB-DEG*], lysine [*asd*, *dapBD*, and *lysAC*], valine [*ilvBCEN*], and phenylalanine and tryptophan [*aroABCEGK* and *trpABCE*]), transcription and translation (*rplA* and *rpsDFMNPT*), transcription release factors (*prfAB*), aminoacyl tRNA synthetases (*glnS*, *leuS*, *serS*, *tyrS*, and *valS*), and energy synthesis (*gapA* and *atpBCFGH*). In addition to genes involved in the incomplete TCA cycle of *Sulcia* (*aceF*, *acoA*, *lpdA*, *korB*, and *sucA*) and the pathway for the conversion of glutamine to carbamoyl phosphate (*carAB*). Although some of these genes are clearly important to the host (e.g., amino acid-related genes), others are key components of *Sulcia*'s independent cellular stability. For example, *Sulcia*'s *groEL* gene, which assists in the folding of damaged proteins (reviewed in Kupper et al., 2014), exhibits an overall pattern of strong purifying selection, but some sites within it are under positive selection. As suggested previously in pea-aphids, the sites within *groEL* under positive selection are involved in gene interaction domains that likely improve its ability to bind with other rapidly evolving proteins (Fares et al., 2002, 2004).

In *Nasuia*, only four genes are predicted to be under positive selection with site-based models. This result is congruent with results from the M0 model that show most *Nasuia* genes are under strong purifying selection (Figure 5). None of these genes are associated with amino acid synthesis and are too few for COG enrichment analyses. Given *Nasuia*'s elevated rates of molecular evolution, selection seems to be operating to maintain function, even though this symbiont has a higher rate of gene loss compared with its partner. The ongoing widespread gene losses from *Nasuia*'s genome may generally result from its high rates of molecular evolution. In this scenario, the probability of a random mutation disabling a gene that can become fixed through genetic drift is much higher (Wernegreen, 2015). Similar results have also been demonstrated in the genomes of *Blattabacterium* from cockroaches and *Buchnera* from pea-aphids, where the rate of genome reduction is also associated with an increased mutation rate rather than selection acting on genes (Bourguignon et al., 2020; Kinjo et al., 2021).

Finally, to further understand whether patterns of selection can predict convergent loss of particular genes, we performed a test of selection with the M0 model (described above) on genes that are not universally retained in *Nesophrosyne*'s *Sulcia* and *Nasuia* genomes. Some symbiotic genes that are only retained in one host species (e.g., the gene cassette containing *rpsU*, *tyrS*, *mnmA*, and *fpr*) could not be tested owing to the lack of information to make pairwise comparisons possible. In *Nasuia*, genes that have been lost in some taxa, but not in others, are generally undergoing strong purifying selection ($n = 4$; *rpmG*, *rpmJ*, *rpsQ*, and *rsmD*; avg. $dN/dS = 0.033$), while one gene is undergoing relaxed purifying selection (*dnaN*; $dN/dS = 0.183$). In *Sulcia*, we see a similar pattern ($n = 2$). One gene is undergoing strong purifying selection (*infC*; $dN/dS = 0.0001$), while the other is undergoing relaxed purifying selection (*bamA*; $dN/dS = 0.222$). These results suggest that types of selection alone are not strong predictors of whether genes will eventually be lost.

Conclusion: Divergent evolutionary forces shape co-primary symbiont genomes

Our sampling of *Nesophrosyne* leafhoppers and their symbionts provides a fine-scale look into the processes that underlie symbiont genome evolution. Gene losses among the *Nesophrosyne*'s *Sulcia* and *Nasuia* lineages demonstrate that there is still ongoing volatility, even among symbionts with two of the

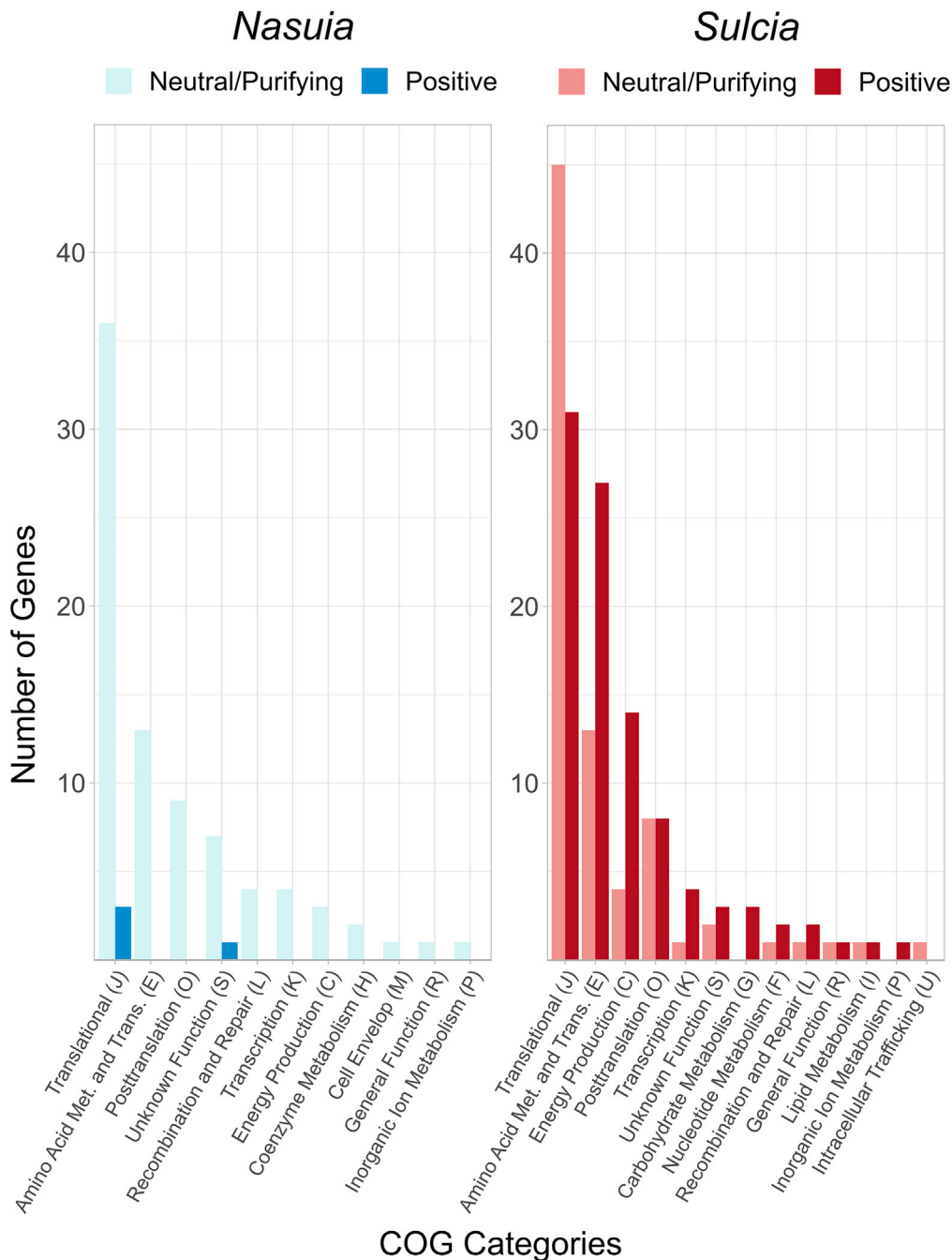


Figure 5. Bar chart showing genes undergoing positive selection or purifying/neutral selection across *Sulcia* and *Nasuia* genomes from Hawaiian leafhoppers (*Nesophrosyne*)

Genes are binned into their Clusters of Orthologous Groups (COGs) functional categories (Tatusov et al., 2000). Bars are color coded according to their genome (i.e., *Sulcia* or *Nasuia*) and selection (i.e., Positive or Neutral/Purifying). We used two nested models, M1a-M2a and M7-M8, to determine overall support for selection among genes (See section methods for further explanation; Anisimova et al., 2001). The likelihood scores were compared within paired models (chi-squared test; $p \leq 0.05$) to indicate significant positive selection for genes.

smallest known genomes. However, these patterns vary widely between these symbiont species. *Sulcia* has one of the slowest evolving genomes known, while *Nasuia* has an exceptionally fast evolving one. *Sulcia* also has far more genes under positive or relaxed purifying selection (80% of genes tested) than *Nasuia*'s

genome, which is largely under strong purifying selection (86% of genes tested). Taken together, our results indicate that the two symbionts experience independent cellular, metabolic, and evolutionary pressures. These differences may further lead to a high level of retention of *Sulcia* among the Auchenorrhyncha and the relatively high turnover of its partner symbionts (reviewed in Koga et al., 2013; Bennett and Moran, 2015; Sudakaran et al., 2017; Bourguignon et al., 2020).

Regardless of the differences between *Sulcia* and *Nasuia*, our results show that there is repeatability in gene losses that may be more easily accommodated if mutations render them non-functional. Both symbionts exhibit convergent gene loss events, particularly regarding genes involved in transcription and translation. These findings suggest a pre-adapted genetic or cellular host environment that permits these genes to be repeatedly lost (see Figures 1 and Figure 2). The host, or partner symbionts, may be capable of filling gaps in these bacterial cellular processes, as is predicted to occur in a wide range of hemipteran insect systems (Hansen and Moran, 2011; Sloan et al., 2014; Luan et al., 2015; Mao et al., 2018; Van Leuven et al., 2019; Mao and Bennett, 2020).

Finally, even though ancient symbiont genomes of insects show some level of conservation among host lineages, a closer look among host sister species reveals complex patterns of gene loss and modes of selection. The evolutionary processes acting on even the tiniest symbiont genomes are clearly dynamic and highly variable. This understanding can be overlooked when comparing lineages among disparately related taxonomic groups.

Limitations of the study

While we sequenced leafhopper species that range the diversity of the genus *Nesophrosyne*, we do not have symbiont lineages sequenced from all 200 + species. Therefore, the range of gene loss exhibited with our species may be greater in other lineages. Additionally, we assembled the mitochondrial sequences to compare rates of evolution; however, we do not have insight into the role of the host nuclear genome in supporting symbiont genome loss or correlated rates of evolution. Finally, a more thorough investigation within and between populations of leafhopper species would provide more information on how host-symbiont co-evolve on an ecological scale.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.isci.2022.104786>.

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AUTHOR CONTRIBUTIONS

This study was conceived by G.M.B. and Y.M.V., field species collection was conducted by G.M.B., dissections were conducted by G.M.B. and Y.M.V., genome assembly and bioinformatic analyses were performed by Y.M.V., and the article was written by Y.M.V. and G.M.B.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological samples		
<i>N. sp.</i> 295	USA: East Hawaii Island	EHBR
<i>N. ogradyi</i>	USA: East Maui Island	EMBR
<i>N. kanawao</i>	USA: North Hawaii Island	NHBR
<i>N. makaihe</i>	USA: West Oahu Island	WOBR
<i>N. sp.</i> 23	USA: Kauai Island	KICL
<i>N. haleakala</i>	USA: Haleakala Mtn, Maui	HMCO
<i>N. sp.</i> 302	USA: Hawaii Island	HICO
<i>N. sp.</i> 58	USA: Maui Island	MICO
<i>N. sp.</i> 29	USA: Kauai Island	KIDO
<i>N. maratima</i>	USA: Oahu Island	OIDO
<i>N. sp.</i> 281	USA: Hawaii Island	HIMY
<i>N. sp.</i> 126	USA: Kauai Island	KIMY
<i>N. sp.</i> 242	USA: Molokai Island	MIMY
<i>N. sp.</i> 246	USA: Oahu Island	OIMY
<i>N. montium</i>	USA: Hawaii Island	HIPI
<i>N. sp.</i> 17	USA: Kauai Island	KIPI
<i>N. sp.</i> 48	USA: Maui Island	MIPI
<i>N. ponapona</i>	USA: Oahu Island	OIPI
<i>N. sp.</i> 15	USA: Molokai Island	MIPS
<i>N. sp.</i> 21	USA: Kauai Island	KISC
Orosius sp.		
Deposited Data		
BioProject	NCBI – GenBank	PRJNA816609
<i>Sulcia</i> sequence data	NCBI – GenBank	CP093890 to CP093909
<i>Nasuia</i> sequence data	NCBI – GenBank	CP094180 to CP094197
Insect mitochondrial sequence data	NCBI – GenBank	ON135504 to ON135524
Software and algorithms		
Trimmomatic	Bolger et al., 2014	v0.39
FastQC	Andrews, 2010	v0.11.9
SPAdes	Bankevich et al., 2012	v3.14
GLIMMER	Delcher et al., 2007	v3.02
Geneious Pro	Drummond et al., 2011	
BLAST	Altschul et al., 1990	
Bowtie2	Langmead and Salzberg, 2012	v2.3.5.1
RAST	Overbeek et al., 2014	v2
RNAmmer	Lagesen et al., 2007	v1.2
tRNAscan-SE	Lowe and Chan, 2016	v2.0
MITOS	Bernt et al., 2013	
MAFFT	Katoh and Standley, 2013	v7.455

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Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
PartitionFinder 2	Lanfear et al., 2017	
BEAST	Drummond and Rambaut, 2007	v.1.10.4
Tracer	Rambaut et al., 2018	v1.7.1
RWTY	Warren et al., 2017	v1.0.2
Phytools	Revell, 2012	v.1.0–1
RStudio	RStudio Team, 2018	
MEGAX	Kumar et al., 2018	v.10.2.4
JModelTest2	Darriba et al., 2012	v2.1.10
PAML (codeml)	Yang 2007	v4.8
Other		
<i>Sulcia</i> genome assembly from <i>Macrostes quadrilineatus</i>	Bennett and Moran (2013)	NCBI: CP006060
<i>Nasua</i> genome assembly from <i>Macrostes quadrilineatus</i>	Bennett and Moran (2013)	NCBI: CP006059
<i>Sulcia</i> genome assembly from <i>Entylia carinata</i>	Mao et al., 2017	NCBI: CP021172
<i>Nasua</i> genome assembly from <i>Entylia carinata</i>	Mao et al., 2017	NCBI: CP021173

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources, data, and codes should be directed to and will be fulfilled by the lead contact, Gordon Bennett (gbennett2@ucmerced.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Sequence data have been deposited with NCBI under the BioProject number: PRJNA816609 and are publicly available as of the date of publication. *Sulcia* genomes can be found under the accession numbers GenBank: CP093890 to GenBank: CP093909. *Nasua* genomes can be found under the accession numbers GenBank: CP094180 to GenBank: CP094197. Insect mitochondrial genomes can be found under the accession numbers GenBank: ON135504 to GenBank: ON135524. These accession numbers are also listed in the [key resources table](#).
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Sample collection

We sampled a targeted set of 20 species that span the ecological and phylogenetic diversity of the genus. Adult female and male leafhoppers were field-collected and stored in ethanol to be used for downstream analysis. The selected species are also specific to eight different host-plant species that encompass the diversity of host-plant families and genera that the genus is associated with (see [Table 1](#)).

METHOD DETAILS

Genome sequencing

For each target species, ten field-collected individuals were pooled to obtain enough DNA for genomic sequencing. DNA was extracted with a DNAeasy kit (Qiagen) and concentration quantified with a Qubit 3.0 fluorometer (ThermoFisher). Libraries and Illumina MiSeq sequencing were conducted at UC Berkeley qB3 Functional Genomics Lab for 4 million 2 × 300 base pair (bp), paired-end reads.

Genome assembly and annotation

Raw reads were quality filtered and cleaned of adapters using Trimmomatic v0.39 and checked with FastQC v0.11.9 (Andrews, 2010; Bolger et al., 2014). Assembly of symbiont genomes was done using SPAdes v3.14 (program settings: -k 127, -only-assembler, -meta; Bankevich et al., 2012). Since extracts contain both host and bacterial DNA, symbiont and mitochondrial genomes were manually extracted by using features unique to each symbiont (*i.e.*, high relative coverage and high AT content). Extracted contigs were verified and confirmed with BLAST searches of open reading frames predicted with GLIMMER v3.02 in Geneious Pro (Altschul et al., 1990; Drummond et al., 2011). To confirm consistent assembly coverage and circularization of bacterial genomes, quality filtered reads were aligned to the completed symbiont genome using Bowtie2 v2.3.5.1 (program settings: -local; Langmead and Salzberg, 2012). Linear chromosomes were circularized by breaking contigs and attaching ends. High, consistent coverage across these ends were verified to confirmed closure of the circular bacterial chromosomes. No plasmids were identified, as is to be expected for tiny symbiont genomes (Bennett and Moran 2013).

Initial genome annotations were done with RAST v2 (Overbeek et al., 2014). Annotations were then verified with GLIMMER v3.02 gene predictions that were checked with BLASTP searches against the nr database (Altschul et al., 1990; Delcher et al., 2007). Bacterial RNA genes were further identified with RNAmmer v1.2 and tRNAscan-SE v2.0 (Lagesen et al., 2007; Lowe and Chan, 2016). Mitochondrial genes were identified with MITOS (Bernt et al., 2013).

QUANTIFICATION AND STATISTICAL ANALYSIS

Phylogenetic tree construction

To construct a phylogenetic tree in absolute time for downstream analysis, and to verify relationships with the *Nesophrosyne* genus, we extracted and aligned complete mitochondrial genomes of our sampled insect species. We included the mitochondrial DNA from *Orosius* sp. as the known outgroup for *Nesophrosyne* (Bennett and O'Grady, 2012; Fletcher et al., 2017). We also tested co-cladogenesis between host and symbiont by extracting and aligning all *Nasuia* and *Sulcia* genes. For the mitochondria and symbiont genomes, each gene was individually aligned with MAFFT v7.455 using the L-INS-i model (Katoh and Standley, 2013). Genes that did not occur in all genomes, or that were difficult to align with confident site homology, were omitted.

The resulting phylogenetic datasets included concatenated protein coding and ribosomal genes for a total of 184 genes (181,781 sites) from *Sulcia*, 99 genes (86,095 sites) from *Nasuia*, and 15 genes (14,304 sites) from leafhopper mitochondria. Best-fit models of nucleotide substitutions and partitioning schemes were determined using PartitionFinder 2 (program settings: branchlengths = unlinked, models = all, model_selection = bic; Lanfear et al., 2017). Bayesian time calibrated phylogenies were then inferred using BEAST v.1.10.4, using the generated partition scheme and corresponding molecular substitution models (Drummond and Rambaut, 2007). The tree prior included the yule process speciation with a random starting tree. Five internal node calibrations were selected following our previous phylogenetic study of the *Nesophrosyne* (Bennett and O'Grady, 2013). Briefly, internal node calibrations were determined from *Nesophrosyne* species divergences that match the sequential geological formation of the Hawaiian Islands (*i.e.*, progression rule). Calibrations were applied with a normal prior distribution since absolute species divergence could have occurred earlier or after island formation (see Bennett and O'Grady, 2012 for additional information; Bennett and O'Grady, 2013). Multiple BEAST chains were run per genome alignment and sampled every 1000 generations following Bayesian recommendations (two chains with four million generations; Huelsenbeck et al., 2002). Runs were performed with an uncorrelated relaxed clock with a lognormal distribution. Convergence and stationarity of chains were verified with ESS values were >200 using Tracer v1.7.1 and RWTY v1.0.2 (Rambaut et al., 2018; Warren et al., 2017).

Ancestral genome reconstruction and ancestral state reconstruction

To determine patterns of gene loss (*i.e.*, unique vs convergent gene losses) among symbionts between host lineages, we reconstructed the ancestral gene retention across *Nesophrosyne's* symbionts. We further included symbiont genomes from previously sequenced lineages including the aster leafhopper (*Macrostelus quadrilineatus*) and the keeled treehopper (*Entylia carinata*) (Bennett and Moran, 2013; Mao et al., 2017). Maximum likelihood ancestral state reconstructions were estimated with phytools v.1.0-1 package (Revell, 2012; RStudio Team, 2018). We used a custom model that allows for the loss of genes to occur (no

gene gain) to account for the inability of symbionts to recombine with other environmental or symbiotic bacteria. Posterior density of stochastic character maps was generated by simulating 100 trees.

Patterns of molecular evolution

To test for genome wide substitution rates across *Sulcia* and *Nasuia* genes, as well as host mitochondria, we used MEGAX v.10.2.4 (Kumar et al., 2018). Model selection for pairwise evolutionary distances were selected with JModelTest2 v2.1.10 with constricted model selection to those available for MEGAX analyses (e.g., Jukes-Cantor, Tamura-Nei, etc.; Darriba et al., 2012). To test the rate of substitutions between islands within the same plant family, two pairwise analyses were done: (i.) the oldest diverging species (e.g., Kaua'i and Hawai'i species) and (ii.) the closest diverging species (e.g., Maui and Hawai'i species). These ages ranged from the most recent divergence (0.351 MYA in *Broussaisia*) to the most ancient divergence (2.4239 MYA in *Pipturus*; see Figure 4A). Rates of substitutions were graphed across absolute time of divergence. To test whether *Sulcia* and *Nasuia* show correlation in their evolutionary rates with *Nesophrosyne* mitochondrial genes, we performed a Kruskal-Wallis test and a pairwise comparison with a Wilcoxon rank sum test and a Benjamini-Hochberg correction for multiple tests (Wilcoxon, 1945; Benjamini and Hochberg, 1995; McKight and Najab, 2010; RStudio Team, 2018). Additionally, in our analysis, we consider the rate of evolution in absolute time by testing across host-plant groups (specifically *Broussaisia*, *Coprosma*, *Pipturus* and *Myoporum*) and within host-plant clades (Figure 4).

Selection among symbiont genomes

In order to test for selection among symbiont genes, we measured rates of synonymous (d_S) and nonsynonymous (d_N) substitutions with codeml v4.8 using the mitochondrial time-calibrated phylogeny (Yang 2007). The ratio of nonsynonymous to synonymous substitutions ($\omega = d_N/d_S$) were calculated for each aligned *Sulcia* and *Nasuia* gene. Selection is calculated by measuring the ratio of nonsynonymous to synonymous substitutions (denoted by ω), where $\omega > 1$, $\omega = 0$ and $\omega < 1$ indicate positive, neutral and purifying selection, respectively (Yang and Bielawski, 2000).

We estimated rates of synonymous and nonsynonymous substitutions using three models. The M0 model was used to test for selection (ω) across all codon sites. This generated a single ω value that was evaluated further. Additionally, we used two nested models, M1a-M2a and M7-M8, to determine strong support for selection in codon sites (Anisimova et al., 2001). Models M1a and M7 are constrained and disallow positive selection while the M2a and M8 models are unconstrained, allowing for positive selection. The M7-M8 models offer a more stringent test of positive selection (Anisimova et al., 2001). However, by using multiple nested models, we verify positive selection in genes that are highly supported in both models (Anisimova et al., 2001). The likelihood scores were compared within paired models (chi-squared test; $p \leq 0.05$) to indicate significant positive selection for genes. To confirm the specificity of our results, we only consider genes that were identified as being under significant positive selection by both nested models for downstream analysis. Genes were further separated into Clusters of Orthologous Groups (COG) to test for functional group enrichment within functional categories, using a Fisher exact test with Benjamini-Hochberg Procedure multiple-testing correction (Benjamini and Hochberg, 1995; Fisher, 1992).