Mutational pressure drives differential genome conservation in two bacterial endosymbionts of sap feeding insects

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ABSTRACT

Compared to free-living bacteria, endosymbionts of sap-feeding insects have tiny and rapidly evolving genomes. Increased genetic drift, high mutation rates, and relaxed selection associated with host control of key cellular functions all likely contribute to genome decay. Phylogenetic comparisons have revealed massive variation in endosymbiont evolutionary rate, but such methods make it difficult to partition the effects of mutation vs. selection. For example, the ancestor of Auchenorrhynchan insects contained two obligate endosymbionts, Sulcia and a betaproteobacterium (BetaSymb; called Nasuia in leafhoppers) that exhibit divergent rates of sequence evolution and different propensities for loss and replacement in the ensuing ~300 Ma. Here, we use the auchenorrhynchan leafhopper Macrosteles sp. nr. severini, which retains both of the ancestral endosymbionts, to test the hypothesis that differences in evolutionary rate are driven by differential mutagenesis. We used a high-fidelity technique known as duplex sequencing to measure and compare low-frequency variants in each endosymbiont. Our direct detection of de novo mutations reveals that the rapidly evolving endosymbiont (Nasuia) has a much higher frequency of single-nucleotide variants than the more stable endosymbiont (Sulcia) and a mutation spectrum that is potentially even more AT-biased than implied by the 83.1% AT content of its genome. We show that indels are common in both endosymbionts but differ substantially in length and distribution around repetitive regions. Our results suggest that differences in long-term rates of sequence evolution in Sulcia vs. BetaSymb, and perhaps the contrasting degrees of stability of their relationships with the host, are driven by differences in mutagenesis.

KEYWORDS
mutation spectra, insertion, deletion, variant, genome decay, extinction

SIGNIFICANCE STATEMENT
Two ancient endosymbionts in the same host lineage display stark differences in genome conservation over phylogenetic scales. We show that the rapidly evolving endosymbiont has a higher frequency of mutations, as measured with duplex sequencing. Therefore, differential mutagenesis likely drives evolutionary rate variation in this system.
INTRODUCTION

Sap feeding insects in the order Hemiptera feed exclusively on nutrient deficient plant-saps (phloem and xylem) and rely on bacterial endosymbionts to provide lacking essential amino acids and vitamins (Buchner 1965; Sandström and Moran 1999). Such endosymbionts are housed intracellularly in specialized organs called bacteriomes and are subject to strict vertical transmission from mother to offspring. This repeated transmission bottlenecking and confinement within a host lineage results in genetic drift and endosymbiont genome decay (Moran 1996; Mira and Moran 2002; McCutcheon and Moran 2012; Bennett and Moran 2015; McCutcheon et al. 2019).

Ancient endosymbiont genomes are characterized by their extreme nucleotide composition bias and small size. In fact, many of these genomes are among the smallest known, lacking genes considered essential in free living bacteria (e.g., cell envelope biogenesis, gene expression regulation, and DNA replication/repair; McCutcheon and Moran 2012; Moran and Bennett 2014). Ancient endosymbiont genomes generally experience rapid sequence evolution and extensive lineage-specific gene deletions, leaving only a set of ‘core’ genes necessary to sustain a functioning nutritional symbiosis (Moran et al. 2009; Bennett, McCutcheon, et al. 2016). At the same time, these genomes exhibit remarkable structural stability, often displaying perfectly conserved synteny between lineages that diverged 10s to 100s of millions of years ago (McCutcheon et al. 2009a).

Missing DNA replication and repair genes, and conditions of relaxed selection, allow the AT mutation bias (which is common to most bacteria; Hershberg and Petrov 2010; Hildebrand et al. 2010; Long, Sung, et al. 2018) to drive AT content to levels above 75% in many endosymbionts (Moran et al. 2009; Moran and Bennett 2014; Wernegreen 2016). Notable exceptions are found in the GC rich genomes of Candidatus Tremblaya princeps (58.8% GC) and Candidatus Hodgkinia cicadicola (hereafter Hodgkinia; 58.4% GC; Baumann et al. 2002; McCutcheon and Von Dohlen 2011). Hodgkinia is an endosymbiont of cicadas with an extremely small genome (144 kb) (McCutcheon et al. 2009b). However, population level estimates of the Hodgkinia mutation spectrum reveal an AT mutation bias, indicating that GC prevalence in Hodgkinia likely results from selection on nucleotide identity (Van Leuven and McCutcheon 2012).
In extreme cases of genome decay, ancient endosymbionts can go extinct when they are outcompeted and replaced by newly colonizing bacteria or yeast (Matsuura et al. 2018; McCutcheon et al. 2019). Comparisons amongst ancient endosymbionts reveal massive variability in genome stability and extinction rates. Insects in the suborder Auchenorrhyncha maintain two bacterial endosymbionts: “Candidatus Sulcia muelleri” (hereafter Sulcia) and a betaproteobacterium (hereafter BetaSymb), which are both thought to be ancestral to the group (~300 Ma) (Moran et al. 2005; Dietrich 2009; Bennett and Moran 2013; Bennett and Mao 2018). The partner endosymbionts provision distinct and complementary sets of the 10 essential amino acids that animals are unable to synthesize and are unavailable in the host diet. Sulcia has been nearly universally retained over the ~300 million year diversification of the Auchenorrhyncha (~40,000 described species); in contrast, BetaSymb has been replaced in at least six major lineages, highlighting its relative instability (Dietrich 2009; Bennett and Moran 2015).

The apparent unreliability and relatively frequent loss of the BetaSymb lineage is mirrored by rapid rates of molecular evolution. Pairwise divergence estimates for the two ancestral symbionts in closely related leafhopper species (family Cicadellidae) reveal that Sulcia genomes are highly similar (99.68% nucleotide identity), while BetaSymb genomes are 30-fold more divergent (90.47% nucleotide identity) (Bennett, Abbà, et al. 2016). The dramatic differences in Sulcia and BetaSymb rates of molecular evolution have also been documented across more divergent Auchenorrhyncha lineages (Bennett and Moran 2013; Mao et al. 2017; Bennett and Mao 2018).

While phylogenetic comparisons have greatly improved our understanding of the changes that endosymbiotic genomes experience, they do not adequately parse the relative contributions of mutagenesis, genetic drift, and natural selection. Although it is possible that selection differentially constrains sequence change in Sulcia and BetaSymb (Wernegreen 2016), it has been suggested that the low rate of evolution in Sulcia may correlate with decreased DNA replication (Silva and Santos-Garcia 2015) or low mutational input (Bennett et al. 2014). However, these hypotheses are difficult to test directly as traditional approaches for measuring bacterial mutation, such as mutation accumulation lines (Kucukyildirim et al. 2016; Long, Miller, et al. 2018), are not feasible for obligate endosymbionts. In one case, researchers were able to use the recent demographic history of the insect host to infer endosymbiont mutation
rates (Moran et al. 2009), but the challenge of directly measuring mutation in endosymbionts continues to inhibit efforts to disentangle mutagenesis, drift, and selection in endosymbionts.

A direct measurement of mutation would ideally capture all variants, before some can be filtered by natural selection. While modern DNA sequencing technologies have facilitated an enormous increase in sequenced endosymbiont genomes, they have not necessarily translated to improvements in measuring de novo mutation in endosymbiont genomes. This is because typical sequencing error rates are too high (often above $10^{-3}$ errors per base pair [bp]) to detect rare variation in DNA samples (Schirmer et al. 2016). As a result, these methods can only detect single-nucleotide variants (SNVs) that have existed long enough to rise to a substantial frequency within the sample, either via drift or positive selection.

Fortunately, the recent advent of several high-fidelity DNA sequencing techniques provides the opportunity to obtain a snapshot of extremely low frequency SNVs in endosymbiont DNA (Sloan et al. 2018). One technique, called duplex sequencing, is particularly suited for such a measurement because it has an exceptionally low error rate of less than $2 \times 10^{-8}$ errors per bp (Kennedy et al. 2014; Wu et al. 2020). The high accuracy of this method is achieved by tagging each original DNA fragment with random barcodes and producing multiple sequencing reads from both strands to obtain a consensus sequence. Importantly, this method is robust to single-stranded DNA damage and individual errors introduced during PCR amplification or sequencing because it separately tracks reads originating from the two complementary strands of the original biological molecule and requires support from each.

Here, we measured low frequency variants in the endosymbionts of a leafhopper (Macrosteles. sp. nr. severini) to test the hypothesis that differences in mutagenesis may be responsible for differential evolutionary stability between Sulcia and BetaSymb (called "Candidatus Nasuia deltocephalinicola" in leafhoppers; hereafter Nasuia). We assembled Sulcia and Nasuia reference genomes from a population of M. sp. nr. severini isolated from Hawaii, using standard Illumina and Oxford Nanopore (MinION) libraries. We then created duplex libraries, which were mapped to the reference genomes to detect de novo variants. We found a dramatically higher frequency of independent SNVs in Nasuia than in Sulcia, which suggests the elevated rate of evolution observed in Nasuia over phylogenetic scales is driven, at least in part, by large differences in mutagenesis.
MATERIALS AND METHODS

Growth and DNA extractions of *Macroteles* sp. nr. *severini* lines

A laboratory stock population of field collected *Macroteles* sp. nr. *severini* from Sumida Farms, Aiea, Oahu Island, Hawaii was established on December 8, 2016. Identification of this species, which has not been formally described, followed Le Roux and Rubinoff (2009). For this experiment, eight laboratory populations were established on barley plants from a single randomly selected foundress on June 17, 2017. Populations were grown with occasional plant replacements to maintain colony health for approximately six months and harvested December 18-22, 2017. Two populations, referred to as A and B, survived to produce sufficient number of individuals for downstream genomic sequencing. Populations were further subdivided into two replicates each (A1, A2, B1, B2) for dissection and sequencing. For each replicate, bacteriomes from approximately 100 individuals were dissected out, pooled, and stored in 95% ethanol. Total DNA was extracted using a DNeasy Blood and Tissue kit (Qiagen) following the manufacturer’s protocol, with a 12-hour Proteinase K digestion.

To ensure variants detected with duplex sequencing were not an artifact of endosymbiont DNA transferred to the *M*. sp. nr. *severini* genome nuclear genome, we sequenced DNA isolated from *M*. sp. nr. *severini* heads, which should contain no true endosymbiont DNA. DNA was extracted from a pool of 20 dissected heads using the Qiagen DNeasy Blood and Tissue Kit.

Construction of shotgun Illumina libraries

Standard shotgun Illumina libraries were created from the most concentrated bacteriome DNA sample (B1) as well as the head DNA sample using the NEBNext Ultra II FS DNA Library Prep Kit. We used 50 ng of input DNA, with 15-minute and 10-minute fragmentation steps for the bacteriome and head DNA, respectively. Samples were amplified with 4 cycles of PCR. The libraries had average lengths of 318 bp and 319 bp for the bacteriome and head DNA, respectively.

Construction of duplex libraries from bacteriome DNA for variant detection

Separate duplex sequencing libraries were generated for each of the four replicate bacteriome DNA samples (A1, A2, B1, and B2). Duplex library preparation followed protocols described elsewhere (Wu et al. 2020). Briefly, samples were fragmented with the Covaris M220
Focused-Ultrasonicator and subsequently end-repaired (NEBNext End Repair Module), A-tailed (Klenow Fragment enzyme, 1 mM dATP), adapter ligated (NEBNext Quick Ligation Module) and treated with a cocktail of three repair enzymes (NEB CutSmart Buffer, Fpg, Uracil-DNA Glycosylase, Endonuclease III). 50 pg of repaired and adapter-ligated products were amplified for 19 cycles with the NEBNext Ultra II Q5 Master Mix (New England Biolabs M0544) and dual-indexed with custom IDT Ultramer primers. Adapter dimers were detected when DNA was assessed on an Agilent TapeStation 2200 (High Sensitivity D1000 reagents) and subsequently removed with size selection on a 2% BluePippin gel (Sage Science), using a target range of 300-700 bp. The pooled duplex libraries had an average length of 386 bp.

**Illumina sequencing of shotgun and duplex libraries**

Shotgun and duplex libraries were sequenced on a NovaSeq 6000 platform (2×150bp) at the University of Colorado Cancer Center in separate sequencing runs. Sequencing resulted in 9.4 M read pairs for the bacteriome shotgun library. The shotgun library from *M. sp. nr. severini* head DNA was sequenced in two runs (the first run did not produce enough reads), which resulted in a total of 359 M read pairs. Duplex libraries were also sequenced in two runs (again, the first round of sequencing did not produce enough reads), which resulted in 37.7 M to 44.9 M read pairs per library. The raw sequencing reads are available via the NCBI Sequence Read Archive (SRA) under accessions SRR12112868 (A1), SRR12112867 (A2), SRR12112866 (B1), SRR12112865 (B2), SRR12112862 (*M. sp. nr. severini* head tissue) and SRR12112864 (B1: shotgun) (all in BioProject PRJNA642181).

**MinION library construction and sequencing**

To aid in genome assembly, we also sequenced the (B1) bacteriome sample with the Oxford Nanopore MinION (Jain et al. 2016), with 150 ng of input DNA on a FLO-MINSP6 flow cell, using the Rapid Sequencing Kit (SQK-RAD004). Data were processed using the MinION software release v19.10.1 with default MinKNOW parameters except that base calling was set to high accuracy mode. Only the first 1.1 M reads were used for genome assembly (which accounted for ~ 2/3 of the data generated by the run and averaged 2972 bp in length). The resulting reads are deposited in the NCBI SRA under accession SRR12112863.
Genome assembly and characterization

Endosymbiont genomes were assembled from the bacteriome Illumina shotgun and MinION libraries with the SPAdes genome assembler (v3.11.1) using the -nanopore flag (Bankevich et al. 2012). Scaffolds returned by SPAdes were searched (blastn v2.9.0+) against available *Nasuia* and *Sulcia* genomes from a *M. quadrilineatus* (GenBank: CP006059.1 and CP006060.1, respectively) (Bennett and Moran 2013), and the two longest scaffolds were identified as near complete matches to *Sulcia* and *Nasuia*.

Evidence of chromosome circularity was assessed by searching for the scaffold beginning and ending sequences in the shotgun and MinION reads. Reads that contain the sequence of both scaffold ends (and therefore span the circular gap) were identified for both genomes. In both cases, the scaffolds ended in tandem repeat regions where the number of repeat units varied in different spanning reads (repeat length heterogeneity).

In the *Sulcia* assembly, the scaffold was broken at a 6-bp microsatellite (corresponding to positions 67046-67105 in the final assembly), with anywhere from eight to 11 repeat units in different spanning reads. The most common repeat number in spanning reads was 10, and the genome was manually adjusted accordingly. The same approach was used to confirm the most common tandem repeat number for three other regions in the *Sulcia* assembly that exhibited repeat length heterogeneity (corresponding to regions 16069-16134, 174128-174181 and 178159-178299 in the final assembly), which were identified as scaffold breakpoints or ambiguous calls (Ns) in a separate SPAdes assembly run without the MinION data.

In the *Nasuia* genome, the repeat structure that broke the assembly is far more complicated, and its resolution was only possible through the use of the long-read MinION data (supplementary figure S1). In spanning sequences, we found seven to 53 repeats of a ~72 bp sequence (which contains only one G and one C), with 22 repeats being the most common repeat number. A complicating factor is the intermittent presence of an additional 7-bp sequence, which itself is a tandem repeat of the first seven bp of the 72-bp repeat, resulting in a mix of 72-bp and 79-bp repeats. The shotgun Illumina data were used to determine that 53.3% of repeats were 72 bp and 46.7% of repeats were 79 bp. The spanning MinION read of median length, which contained 22 repeats total, was selected as the basis for resolving the *Nasuia* genome gap (supplementary figure S1).
After the most common repeat numbers were determined and circularity was confirmed, the breakpoints of the chromosomes were shifted, and the *Nasua* scaffold was reverse complemented to reflect the orientation and genome positions of *Nasua* and *Sulcia* accessions from the closely related *M. quadrilineatus* (Bennett and Moran 2013). We then aligned our new assemblies to *M. quadrilineatus* accessions using MAFFT (v7.453) under default parameters (Katoh and Standley 2013). Gaps in the alignments were tabulated with a custom script. Pairwise distances were calculated with MEGA (v10.1.8) (Kumar et al. 2018), using the Tamura 3-parameter model and a gamma distribution. Repetitive sequences were identified with custom scripts that report the position and length of homopolymers and microsatellites of 4+ bp in length (supplementary table 1) (Temnykh et al. 2001).

Both *Sulcia* and *Nasua* were annotated with initial protein-coding gene predictions using Glimmer3 in Geneious v11.1.5 and were then checked against existing symbiont genomes from two other *Macrosteles* host species (Delcher et al. 2007; Bennett, Abbà, et al. 2016). All RNA annotations were done using the “annotate from reference” function in Geneious with a match threshold of >80% similarity. *Nasua* and *Sulcia* genomes from *M. quadrilineatus* were used as the reference genome (Bennett and Moran 2013). GC content at 4-fold degenerate sites was calculated using custom scripts.

**Variant detection with duplex data**

Duplex libraries were processed with our custom data analysis pipeline (Wu et al. 2020; https://github.com/dbsloan/duplexseq), which uses the random barcodes on read ends to first group raw reads into single stranded consensus sequences, which require at least three raw Illumina reads. Complementary single stranded consensus sequences with matching barcodes are then paired to form a duplex consensus sequence (DCS). Therefore, each DCS must contain at least six Illumina reads, though the mean raw read redundancy in these libraries was higher (supplementary figure S2). Duplex consensus sequences were subsequently mapped against the newly assembled reference genomes and the resulting alignment files were parsed to identify indels, single nucleotide variants (SNVs), and coverage per bp. Supplementary table 2 reports total DCS coverage and percent mapping to endosymbiont genomes for the four replicates. A filtering step then checked for the presence of identified variants in a *k*-mer database (*k* = 39) created from the *M. sp. nr. severini* head DNA shotgun library (KMC v. 3.0.0; http://mc.manuscriptcentral.com/gbe)
https://github.com/refresh-bio/KMC) to ensure variants are not derived from bacterial DNA that has been transferred to the host genome (Hotopp et al. 2007; Nikoh et al. 2010). All variants had counts well below those detected for *M. sp. nr. severini* nuclear and mitochondrial sequences.

RESULTS

Higher divergence in *Nasuia* than in *Sulcia* derived from closely related leafhoppers

We assembled *Nasuia* and *Sulcia* genomes (GenBank accession CP060019.1 and CP060020.1, respectively), using a combination of short-read (Illumina) and long-read (MinION) sequences from *M. sp. nr. severini* bacteriome DNA. The resulting genomes (referred to hereafter as *Nasuia-MSEV* and *Sulcia-MSEV*) were generally comparable to those previously sequenced from other hosts in terms of size (*Nasuia-MSEV*: 113 kb, *Sulcia-MSEV*: 190 kb), gene count (*Nasuia-MSEV*: 173, *Sulcia-MSEV*: 224), and GC content (*Nasuia-MSEV*: 16.9%, *Sulcia-MSEV*: 24.0%). We aligned our new assemblies to the previously published *M. quadrilineatus* accessions and found pairwise distances of 7.68% and 0.01% for the *Nasuia* and *Sulcia* pairs, respectively (fig. 1). The ~700-fold higher pairwise divergence in *Nasuia* than in *Sulcia* is much larger than 30-fold difference previously shown in pairwise comparisons between endosymbionts of the closely related *M. quadrilineatus* and *M. quadripunctulatus* (Bennett, Abbà, et al. 2016). This disparity is driven by the near perfect sequence conservation in *M. sp. nr. severini* and *M. quadrilineatus Sulcia* genomes, which only differ at approximately one of every 10,000 positions. Transitions are responsible for 79.6% of the observed pairwise distance in *Nasuia*. The extremely small number of changes in the *Sulcia* comparison prohibits a meaningful calculation of the contribution of transitions to pairwise distance.

We found 1.00 and 0.28 gaps per kb in the *Nasuia* and *Sulcia* pairs, respectively (supplementary table 3). A large (1,352 bp) gap in the *Nasuia* pairs (supplementary table 3) corresponds to the mix of 72 and 79 bp repeats (22 total) in the *Nasuia-MSEV* assembly that we characterized with the MinION data (supplementary figure S1). The lack of this region in the *Nasuia* accession from *M. quadrilineatus* likely reflects differences in methods of genome assembly, rather than an actual difference in genome structure between the two *Nasuia* strains. We found complete conservation of synteny in both endosymbiont pairs, reflecting the high level of gene-order stability seen in many other ancient endosymbionts (fig. 1) (McCutcheon et al. 2019).
Higher frequency of SNVs in *Nasua-MSEV* than in *Sulcia-MSEV*

We then used our new endosymbiont genome assemblies as references for mapping duplex sequence data. We detected a 106-fold higher SNV frequency in *Nasua-MSEV* ($2.18 \times 10^{-5}$) than in *Sulcia-MSEV* ($2.04 \times 10^{-7}$) (fig. 2A, supplementary data S1). For both endosymbionts, observed SNV frequencies are above the noise threshold of $2 \times 10^{-8}$ errors per bp that we have previously established for our duplex protocol (Wu et al. 2020). All 17 of the SNVs identified in *Sulcia-MSEV* were ‘singletons’ (captured by only a single DCS). In contrast, 35 of the 126 positions with SNVs in *Nasua-MSEV* were detected at ‘high frequency’ (captured by more than one DCS). Accordingly, the 106-fold higher SNV frequency in *Nasua-MSEV* is partially driven by variants present at high frequency in the experimental populations. We therefore performed a comparison based on independent SNV events (sites in the genome with a SNV) and found a 18-fold higher independent event SNV frequency in *Nasua-MSEV* than in *Sulcia-MSEV* after normalizing for sequence coverage (fig. 2B, supplementary table 4).

For both *Sulcia-MSEV* and *Nasua-MSEV*, the SNV type with the highest independent event frequency was CG→TA transitions, though SNV type comparisons are low powered in *Sulcia-MSEV* due to the small number of variants (supplementary figure S3). In *Nasua*, AT→GC transitions and CG→AT transversions were also detected at relatively high frequencies. The proportion of transitions out of all independent SNVs was 0.82 and 0.52 for *Nasua-MSEV* and *Sulcia-MSEV*, respectively, though the *Sulcia-MSEV* proportion is likely unreliable given that it is based on so few events. The proportion of transitions in *Nasua-MSEV* appears to be within the range reported for other bacteria (Hershberg and Petrov 2010), and does not deviate from the proportion observed in the comparison with *Nasua* from *M. quadrilineatus* (0.80; binomial test, $p = 0.51$).

Given the extreme AT bias of the genomes (*Nasua-MSEV*: 83.1% AT, *Sulcia-MSEV*: 76% AT), we tested if genomic AT content is at equilibrium, in which case the number of AT-increasing mutations would equal the number of AT-decreasing mutations. Of the 126 (independent event) SNVs detected in *Nasua-MSEV*, significantly less than half (only 43) decreased AT content, while 75 mutations increased AT content (eight changes were AT-neutral) (binomial test, $p = 0.0004$). However, the probability of detecting AT-increasing and AT-decreasing SNVs may not be equal, as relatively GC-rich regions have higher average sequence
coverage due to bias against extreme AT-rich sequences during library amplification. The *Nasua-MSEV* genome has a GC content of only 16.9%, but GC positions accrued 24.0% of the total sequencing coverage. This difference in depth of coverage may lead to the preferential detection of rare variants at GC sites. We therefore adjusted AT-decreasing SNVs to account for decreased detection probability, which we calculated as a ratio of AT coverage/bp and GC coverage/bp (AT-decreasing SNVs were 35.5% less likely to be detected). Adjusted AT-decreasing counts of 56 and AT-increasing counts of 63 were not significantly different from 50:50 equilibrium expectations (binomial test, $p = 0.64$). The five AT-decreasing and six AT-increasing SNVs we detected in *Sulcia-MSEV* provide little power for testing for AT content equilibrium, but neither raw counts nor counts adjusted for the probability of detection deviate significantly from equilibrium expectations (binomial test, $p = 0.77$).

Interestingly, 12 of the 35 high-frequency SNVs in *Nasua-MSEV* were detected in both experimental populations A and B (fig. 3). Shared SNVs could have originated in the ancestor of the A and B foundresses and remained polymorphic during the six months that the experimental populations were maintained (approximately six host generations). Alternatively, the shared SNVs could have arisen independently in the two experimental lines (i.e. homoplasy). There were no SNVs shared across any replicates in *Sulcia-MSEV*.

In *Nasua-MSEV* protein-coding regions, we found that the proportion of nonsynonymous changes was significantly reduced in high-frequency SNVs compared to singletons (Fisher’s Exact Test, $p = 0.003$). Reduction in the proportion of nonsynonymous SNVs in high-frequency SNVs (compared to singletons) occurs in all six substitution types, though none of the six SNV type reductions are significant on their own. (Fisher’s Exact Test, $p > 0.079$; supplementary figure S4, supplementary table 5). The nucleotide substitution spectra also differ between singletons and high-frequency SNVs. Ignoring AT neutral substitutions (AT$\rightarrow$TA and CG$\rightarrow$GC), only 26 of 84 singletons decrease AT content, compared to 17 of 34 high-frequency SNVs (marginal significance with Fisher’s Exact Test, $p = 0.06$). We could not perform the same analysis for *Sulcia-MSEV* SNVs, as they were all singletons. For both endosymbionts, SNVs are not differentially distributed across intergenic, protein-coding, rRNA, or tRNA regions of the genome (supplementary figure S5, supplementary table 6).

Because high-frequency SNVs in *Nasua-MSEV* show evidence of filtering by selection (supplementary figure S4), and differ in spectrum compared to singletons, we repeated the
aforementioned AT equilibrium calculations with only singleton SNVs. AT-decreasing counts of 34 and AT-increasing counts of 50 (adjusted for probability of detection) are marginally different from equilibrium expectations (binomial test, \( p = 0.10 \)). We then used our normalized singleton counts to derive an expected GC equilibrium content of 12.4%, which is lower than the actual *Nasuia-MSEV* GC content of 16.9%, but almost identical to the GC content of 12.2% at 4-fold degenerate sites, which are expected to be under minimal selection.

**No strand-specific mutation asymmetry detected in *Nasuia-MSEV* protein-coding genes.**

Coding strands in CDS regions of the *Nasuia-MSEV* genome are enriched for G compared to C (positive GC skew of 0.16 or 1.4 Gs for every C). In endosymbiotic genomes, positive GC skew on the leading strand of DNA replication is thought to result from asymmetrical C→T changes on the leading strand, which is more susceptible to cytosine deamination due to its prolonged existence in a single-stranded state (Klasson and Andersson 2006). During transcription, the increased single-stranded exposure of the coding DNA strand has also been proposed to lead to asymmetrical C→T changes in bacterial genomes (Francino and Ochman 1997).

We tested if mutation drives GC skew on coding strands of *Nasuia-MSEV* by comparing reciprocal mutations (C→T vs. G→A) to expected counts given C and G coverage. We did not find evidence for asymmetry in reciprocal mutations in CG→TA transitions, or for any of other five substitution types in *Nasuia-MSEV* coding strands in CDS regions (binomial test, all p-values \( \geq 0.16 \)) (supplementary table 7). We did not attempt to perform the reciprocal strand analysis for *Sulcia-MSEV* given the small amount of SNVs.

**Comparison of indels in *Nasuia-MSEV* and *Sulcia-MSEV***

Overall, average indel frequencies for *Nasuia-MSEV* and *Sulcia-MSEV* are similar (4.92×10^{-6} and 4.51×10^{-6} respectively). *Nasuia-MSEV* has a higher frequency of deletions than insertions while *Sulcia-MSEV* has a higher frequency of insertions than deletions (fig. 4a, supplementary data S2). However, it is important to recognize that during genome assembly, determination of the dominant repeat number in regions with repeat length heterogeneity can influence whether an indel is considered a deletion or an insertion, so comparisons of the frequencies of deletions vs. insertions should be cautiously interpreted. Indeed, when the three
hypervariable regions that were identified during genome assembly (see Materials and Methods) are excluded from *Sulcia* indel analysis, insertions become less frequent than deletions (supplementary table 8). Duplex sequencing did not detect any indels at the 72/79 bp tandem repeat in the *Nasua* genome (supplementary figure S1), despite MinION data revealing this region to be hypervariable in terms of repeat number. Most repeats in this region are likely too long to be spanned by duplex sequences (the shortest repeat we found in the MinION data was seven repeats long) though the extreme AT content and frequent homopolymers in the region could also explain why only one DCS read from all four libraries mapped to the region (visual inspection in IGV; Thorvaldsdóttir et al. 2013). For all downstream indel analysis, no regions were excluded (i.e., the full data set was considered).

Both endosymbionts have indels that are present at high frequencies, which are sometimes shared across experimental populations A and B. Like with high-frequency SNVs in *Nasua-MSEV*, these indels could arise from homoplasy or from shared ancestry and persistence in a polymorphic state during the growth of experimental cultures. Local sequence context plays a large role in indel distribution across both genomes as 94.1% and 98.4% of indels overlap with repetitive genomic sequence (microsatellites and homopolymers) for *Nasua-MSEV* and *Sulcia-MSEV*, respectively. Given the apparent influence of local sequence context on indel occurrence, and the lack of local sequence effects on SNV occurrence, we suspect that high frequency indels often reflect multiple homoplasious events (e.g., long homopolymers can be subject to “multiple hits”). Our indel analysis therefore emphasizes total indel frequency rather than independent event indel frequency (fig. 4b).

We found that *Nasua-MSEV* has a higher frequency of homopolymer-associated indels, while *Sulcia-MSEV* has a higher frequency of microsatellite-associated indels (fig. 5a). These differences do not merely reflect a difference in the repetitive landscape of the symbiont genomes, as the frequencies are normalized by the specific coverage of a given repeat type (microsatellite or homopolymer), and *Nasua-MSEV* contains more of both repeat type per kb than *Sulcia-MSEV* (supplementary table 1). For both endosymbionts, we find that as homopolymers increase in length, indel frequency also increases (fig. 5b).

**DISCUSSION**

**Rapid evolution of BetaSymb likely driven by higher mutation rate.**
BetaSymb (represented by Nasuia in leafhoppers and other names depending on the insect host; Vidania in planthoppers and Zinderia in spittlebugs) evolves much more rapidly than Sulcia, despite the partner endosymbionts having occupied a nearly identical ecological niche (i.e., bacteriocytes in a shared host) for ~300 million years (Bennett, Abbà, et al. 2016; Bennett and Mao 2018). In addition, BetaSymb experiences increased evolutionary turnover and has been replaced in at least six Auchenorrhyncha lineages (Bennett and Moran 2015). This accelerated evolution with respect to Sulcia does not appear to be limited to BetaSymb, as some of its lineage-specific replacements such as Hodgkinia (Alphaproteobacteria) and Baumannia (Gammaproteobacteria) show similar asymmetries (Bennett et al. 2014; Campbell et al. 2017). Our finding of a higher frequency of de novo SNVs in Nasuia-MSEV (fig. 3) suggests the increased evolutionary rate in BetaSymb is likely driven by mutation.

Our results therefore support the hypothesis that genome decay and extinction in endosymbionts is driven by increases in mutation rate (Itoh et al. 2002). Such increases intensify Muller’s ratchet, which is already prevalent in endosymbionts experiencing strict vertical transmission and host confinement (Rispe and Moran 2000). That Sulcia also has a tiny genome (ranging from 190 to 244 kb; Woyke et al. 2010), but exhibited a depressed mutation frequency in our study, and greatly reduced synonymous site divergence compared to multiple partner lineages (Bennett et al. 2014; Bennett and Mao 2018), illustrates that mutation is not the only force responsible for genome reduction (Bennett and Moran 2015).

Because variant frequencies are the product of both the mutation rate (μ) and the effective population size (Nₑ), it is important to recognize that a higher Nₑ in Nasuia-MSEV could also lead to the 18 to 106-fold higher SNV frequency we observed in Nasuia-MSEV. However, current evidence does not support a larger Nₑ for Nasuia. Sequence coverage comparisons from our bacteriome shotgun library reveal a 1.5-fold higher per bp depth of coverage in Sulcia-MSEV than in Nasuia-MSEV (Pedersen and Quinlan 2018). This suggests the standing population size may actually be smaller in Nasuia-MSEV than in Sulcia-MSEV, though bias against AT-rich sequences during library amplification could also drive lower coverage of the Nasuia-MSEV genome, as coverage for GC base pairs is only 1.2-fold higher in Sulcia-MSEV.

The standing population size does not preclude the possibility that there may be differences in the intensity of the transmission bottlenecks for the two endosymbionts. Microscopy has revealed that, while Sulcia and Nasuia are housed in distinct bacteriocyte types,
they both successfully migrate to and infect terminal oocytes (Szklarzewicz et al. 2016). It is not known, however, if the number of transmitted bacteria differs for Nasuia vs. Sulcia in leafhoppers. In cicadas, Sulcia and partner symbiont Hodgkinia are transmitted at similar numbers, though Hodgkinia transmission dramatically increases in lineages where the Hodgkinia genome has been fragmented into multiple chromosomes (Campbell et al. 2018). Finally, a larger Ne in Nasuia-MSEV should lead to more efficient purifying selection, which would be inconsistent with the fact that Nasuia exhibits more extensive genome decay (high AT content, reduced size, etc.) over phylogenetic scales. This final argument relies on indirect evidence, and a study of actual Nasuia vs Sulcia transmission is needed to definitively rule out the possibility that differences in Ne are driving different observed variant frequencies. Furthermore, the idea that reduced Ne drives genome size reduction has recently been challenged, as mutation rates are apparently better at predicting gene loss than dS/dN (a proxy for Ne) in several independent endosymbiont lineages (Bourguignon et al. 2020).

Interestingly, 35 variants in Nasuia-MSEV were present in ‘high-frequency’ (detected in more than one DCS). Compared to singletons, high-frequency SNVs in Nasuia-MSEV are less likely to result in nonsynonymous changes (supplementary figure 2), suggesting they have been filtered by selection. This idea is further supported by the observation that high frequency SNVs differ in AT spectra compared to singletons (marginal significance in Fisher’s Exact Test, p = 0.06). Singletons in our data thus most accurately represent the true mutation spectra, before it can be shaped by selection. When we used singletons to predict equilibrium GC content, we got a value nearly identical to the observed GC content at 4-fold degenerate sites (12.4% and 12.2%, respectively), which are predicted to experience minimal selection. The actual GC content in Nasuia is higher (16.9%), suggesting that selection on nonsynonymous sites and tRNAs/rRNAs keeps genomic GC content above equilibrium values. As such, it appears that despite the already extreme AT-richness of the Nasuia genome, mutation bias would further erode GC content in the absence of selection (Hershberg and Petrov 2010; Hildebrand et al. 2010; Long, Sung, et al. 2018). Our observation of rapid filtering of high frequency events in Nasuia-MSEV raises the possibility that a previous study finding no AT mutation bias in Buchnera aphidicola (an ancient endosymbiont of many aphids) (Moran et al. 2009) likely did so based on a set of mutations that had been subject to substantial effects of selection. While our data demonstrate that selection is
able to maintain some level of *Nasuia* sequence conservation, it remains possible that *Nasuia* and *Sulcia* are differentially constrained.

Twelve of the 35 *Nasuia-MSEV* high frequency SNVs were shared between experimental populations A and B, which were reproductively isolated for six months (approximately six host generations; Capinera 2008). While it is possible that shared SNVs resulted from homoplasy, some shared SNVs have risen to very high frequency (greater than 45% in one case), which demonstrates they are at least old enough to have spread to a large portion of the experimental population. We find it most likely that shared SNVs were ancestral to endosymbiont populations present in the initial insect foundresses and persisted as polymorphisms through the duration of insect growth but cannot rule out the possibility that some or all shared SNVs have arisen through homoplasy.

**Can differences in DNA replication and repair machinery explain greater genome conservation in *Sulcia?***

While *Nasuia-MSEV* and *Sulcia-MSEV* lack most subunits of the primary bacterial polymerase (DNA Pol III), they both retain α and ε subunits (*dnaE* and *dnaQ*), which in *Escherichia coli* perform polymerization and 3’ → 5’ exonuclease activity, respectively (Fijalkowska et al. 2012). *Nasuia-MSEV* also uniquely retains the β subunit (*dnaN*), which facilitates binding between subunit α and the template DNA strand (Gui et al. 2011). *Nasuia-MSEV* retains two additional DNA replication and repair (RR) genes lost in *Sulcia-MSEV*: *dnaB* (helicase) and *ssb* (single stranded binding protein) (Bennett and Moran 2013).

Only one RR gene, *mutS*, is absent in *Nasuia-MSEV*, but retained in *Sulcia-MSEV*. In other bacteria, the MutS protein recognizes and initiates repair of mismatched bases and small indels (Dettman et al. 2016; Long, Miller, et al. 2018). While the role of *mutS* in *Sulcia* has never been experimentally validated, it contains all five of the domains present in the *E. coli mutS* (Ogata et al. 2011; Groothuizen and Sixma 2016). The extremely low frequency of SNVs we observed in *Sulcia-MSEV* (fig 1, Supp Table 4) could result from *mutS*-directed mismatch repair (MMR).

Mutation accumulation experiments with *Pseudomonas aeruginosa* reveal *mutS* mutants experience a 230-fold increase in indels (average length of 1.1 bp) compared to WT lines, demonstrating the importance of *mutS* in short indel surveillance (Dettman et al. 2016). While
we observed a high frequency of indels in *Sulcia-MSEV*, most were expansions or contractions of repeats at existing microsatellites. In fact, only 19 of the 385 indel containing DCS reads in *Sulcia-MSEV* were of 1 bp in length (compared to 145 of 154 in *Nasua-MSEV*) (supplementary table 9). Even more striking is the complete lack of 1-bp indels in the alignment of *Sulcia-MSEV* with *Sulcia* from *M. quadrilineatus* (compared to 48 1-bp indels in the equivalent *Nasua* alignment) (supplementary table 3). The rarity of 1-bp indels in *Sulcia-MSEV* thus further supports the idea that *mutS* is critical for maintaining *Sulcia* genome conservation through surveillance of homopolymer expansions/contractions.

Interestingly, the genome of *Sulcia* in leafhoppers actually has a reduced set of RR genes compared to those found in *Sulcia* genomes in other Auchenorrhynchan lineages. *Sulcia-CARI*, an endosymbiont of spittlebugs, possesses the two DNA Pol III subunits found in *Sulcia-MSEV*, but has also retained in subunits *dnaB, dnaN, dnaX*, and polymerase accessory subunits *holA* and *holB*. Additional RR genes retained by *Sulcia-CARI*, but absent in leafhopper *Sulcia* genomes, include gyrase genes *gryA, gryB* and MMR genes *mutL*, and *mutH* (McCutcheon and Moran 2010; Bennett and Moran 2013). Despite lacking the above listed RR genes, leafhopper *Sulcia* genomes remain remarkably stable, as demonstrated by phylogenetic comparisons at multiple levels of leafhopper divergence in this and previous studies (Bennett, Abbà, et al. 2016; Mao et al. 2017).

Given that MutS works in concert with *DnaN, mutL* and *mutH* products in typical bacterial MMR (Simmons et al. 2008; Jiricny 2013), it is curious that *DnaN* and *mutH* have been completely lost and *mutL* has been pseudogenized in leafhopper *Sulcia* genomes (Bennett and Moran 2013; Bennett, Abbà, et al. 2016). A recent assay of *M. quadrilineatus* gene expression revealed that in the bacteriome, leafhopper hosts often overexpresses homologs of RR genes that are absent from endosymbiont genomes, raising the possibility that leafhopper hosts may support endosymbiont genome replication and repair (Mao et al. 2018). Host support has been hypothesized to enable endosymbiont gene loss (Hansen and Moran 2011; Husnik et al. 2013; Russell et al. 2013; Sloan et al. 2014), and a recent study has demonstrated that the partner endosymbionts of cicadas likely rely of host encoded machinery to acquire correctly processed tRNAs (Van Leuven et al. 2019). However, none of the four putative *mutL/mutH* homologs in the *M. quadrilineatus* transcriptome showed substantial overexpression in the *Sulcia* specific bacteriocyte (compared to expression in the insect body) (supplementary table 9). Thus, it
remains unclear if Sulcia mutS is participating in conventional MMR or if it has acquired novel functions in support of Sulcia genome conservation. Functional characterization of MMR in leafhopper Sulcia would provide valuable insight into what appears to be an atypical role for an ancient and highly conserved mismatch recognition protein (Ogata et al. 2011; Jiricny 2013).

The missing mutS in Nasuia provides an additional opportunity for host support. Mao et al. (2018) reported strong overexpression of one host mutS homolog (referred to as TRINITY_DN66078_c1_g1 in that study; see supplementary table 10) in the Nasuia bacteriome, but it is highly unlikely that this is involved in MMR for two reasons. First, the identified homolog is related to eukaryotic MSH4, which is a gene family that lacks a mismatch recognition domain and is involved in recombination rather than MMR (Ogata et al. 2011). Furthermore, it appears to encode only a partial MSH4 protein fused with a C-terminal sequence of unidentified origin. Therefore, it is likely that Nasuia lacks conventional MMR function.

While mutS may play a role in Sulcia genome conservation, retention of mutS is not a cure-all. For example, BetaSymb in spittlebugs (named “Candidatus Zinderia insecticola”) retains the entire mismatch repair gene set (mutS, mutL, and mutH) but displays a rate of evolution typical of Nasuia and other BetaSymb lineages (Bennett and Moran 2013; Koga et al. 2013). We posit that the mutS in Sulcia may be particularly efficient at recognizing and removing mismatched bases and short indels. All Sulcia genomes analyzed to date retain mutS, but identification of a Sulcia lineage that has lost mutS would facilitate a test of the hypothesis that mutS may be the key to Sulcia genome conservation.

CONCLUSION

The two ancestral endosymbionts in auchenorrhynchan insects vary dramatically in their propensity for extinction vs. retention, thus providing a natural comparative framework for investigating what facilitates long-term, stable endosymbiotic relationships. Compared to Sulcia, which exhibits remarkable genome conservation for an ancient endosymbiont and has been nearly universally retained, BetaSymb displays substantially elevated rates of molecular evolution. We have shown that Nasuia-MSEV, a representative of BetaSymb from leafhoppers, has a substantially higher frequency of de novo mutations than Sulcia-MSEV. Our analysis supports the hypothesis that the evolutionary rate variation in Sulcia vs. BetaSymb is driven by differences in mutational input. We posit that the low mutation rate in Sulcia may also explain
why *Sulcia* is so consistently retained in diverse auchenorrhynchan lineages, while partner endosymbionts are apparently much more transient.

**DATA AVAILABILITY**

The raw sequencing reads are available via the NCBI Sequence Read Archive under accessions SRR12112868, SRR12112867, SRR12112866, SRR12112865, SRR12112862 and SRR12112864. Genome annotations for *Nasuia-MSEV* and *Sulcia-MSEV* are available via GenBank under accessions CP060019.1 and CP060020.1, respectively.

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FIGURES

Fig. 1. Synteny and sequence similarity of *Nasuia* and *Sulcia* from two closely related leafhopper species. Synteny plots reveal gene order to be perfectly conserved in both *Nasuia* and *Sulcia* of *M. quadrilineatus* and *M. sp. nr. severini*. Circles represent significant pairwise alignments between protein-coding genes in the two genomes, with circle diameter indicating alignment length. *Nasuia* exhibits a 700-fold higher sequence divergence than *Sulcia*, which is reflected in the lower amino acid identity (color scale on figure right) of *Nasuia* protein coding sequences. The compact nature of both genomes can be observed in the gene tracks on the top (*M. quadrilineatus* associated) and right (*M. sp. nr. severini* associated) of the synteny plots, where protein coding-genes are shown in purple, rRNA genes are in yellow, tRNA genes are in light blue, and intergenic regions are white. This figure was generated with GenomeMatcher (Ohtsubo et al. 2008)

Fig. 2. Duplex sequencing variant frequency in *Nasuia-MSEV* and *Sulcia-MSEV*. Variant frequencies are shown (A) in terms of total SNV frequency (DCS variant coverage/total DCS coverage) and (B) in terms of independent event frequency (sites with a variant/DCS coverage). In both, the six mutation types and are normalized by DCS coverage of the corresponding reference bases (for example the AT\(\rightarrow\)CG changes are normalized by total AT coverage). *Nasuia-MSEV* is red and *Sulcia-MSEV* is blue. Triangles and squares are experimental populations A and B, respectively. For *Nasuia-MSEV*, the average (of the two experimental populations) independent event frequency (4.97\(\times\)10\(^{-6}\)) is reduced relative to the average SNV Frequency (2.18\(\times\)10\(^{-5}\)) due to the presence of 32 sites where multiple DCS reads mapped with a variant. There is no difference in *Sulcia-MSEV* variant frequencies (2.04\(\times\)10\(^{-7}\)) because all *Sulcia-MSEV* SNVs are singletons (see also supp Fig 1).

Fig. 3. High-frequency variants in *Nasuia-MSEV*. SNV frequencies of the 32 positions in the *Nasuia-MSEV* genome captured by more than one DCS. Frequencies for experimental populations A and B are shown in red and orange, respectively. Experimental populations were subdivided into replicates 1 and 2, which are shown in triangles and squares, respectively. At 12 positions in the genome variants are shared between experimental populations A and B (teal stars) and at 14 positions in the genome variants are shared between replicates 1 and 2 within the two populations (purple stars).
**Fig. 4. Indel frequencies in *Nasua-MSEV* and *Sulcia-MSEV*.** Indel frequencies shown in terms of (A) total frequency (DCS indels/DCS coverage) and (B) Independent event frequency (sites with an indel/DCS coverage). Overall average indel frequencies for *Nasua-MSEV* and *Sulcia-MSEV* are similar – but deletions are more common in *Nasua-MSEV* and insertions are more common in *Sulcia-MSEV*. Independent event frequencies are higher for *Nasua-MSEV* than for *Sulcia-MSEV*. *Nasua-MSEV* is red and *Sulcia-MSEV* is blue. Triangles and squares are experimental populations A and B, respectively.

**Fig. 5. Frequency of repeat-associated indels in *Nasua-MSEV* and *Sulcia-MSEV*. (A)** Frequencies of indels associated with homopolymers and microsatellites and (B) frequencies of homopolymer-associated indels as function of homopolymer length. Frequencies are calculated as the total number of DCS indels in a repeat category divided by DCS coverage of that repeat category. 94.1% of all *Nasua-MSEV* indels and 98.4% of all *Sulcia-MSEV* indels overlap with homopolymers or microsatellites. 90.9 % and 96.1 % of all indels are expansions or contractions of existing repeats for *Nasua* and *Sulcia*, respectively. For *Nasua-MSEV*, the majority of repeat-associated indels are located in homopolymers, and deletions are more common than insertions. For *Sulcia-MSEV*, the majority of indels are located in microsatellites. For both endosymbionts, the frequency of insertions and deletions increase as homopolymer length increases. *Nasua-MSEV* is red, and *Sulcia-MSEV* is blue. Triangles and squares are experimental populations A and B, respectively.
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