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3	Conflicting evolutionary histories of the mitochondrial and nuclear genomes in New World Myotis
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16	Abstract
17	The diversification of Myotis into more than 100 species in just a few million years is one of the
18	most extensive mammalian radiations available for study. Efforts to understand relationships within
19	Myotis have primarily utilized mitochondrial markers, and trees inferred from nuclear markers lacked
20	resolution. Our current understanding of relationships within Myotis is therefore biased towards a set of
21	phylogenetic markers that may not reflect the phylogenetic history of the nuclear genome. To resolve
22	this, we sequenced the full mitochondrial genomes of 37 representative Myotis, primarily from the New
23	World, in conjunction with targeted sequencing of 3,648 ultraconserved elements (UCEs). We inferred the
24	phylogeny of Myotis and explored the effects of concatenation and summary phylogenetic methods, as
25	well as combinations of markers based on informativeness or levels of missing data, on our phylogenetic
26	results. Of the 295 phylogenies generated from the nuclear UCE data, all are significantly different from
27	phylogenies inferred using mitochondrial genomes. Even within the nuclear genome quartet frequencies
28	indicate that around half of all UCE loci conflict with the estimated species tree. Several factors can drive
29	such conflict, including incomplete lineage sorting, introgressive hybridization, or even phylogenetic error.
30	Despite the degree of discordance between nuclear UCE loci and the mitochondrial genome and among
31	UCE loci themselves, the most common nuclear topology is recovered in one quarter of all analyses with

- 32 strong nodal support. Based on these results, we re-examine the evolutionary history of *Myotis* to better
- 33 understand the phenomena driving their unique nuclear, mitochondrial, and biogeographic histories.
- 34

# 35 Keywords

- 36 incomplete lineage sorting, summary tree methods, concatenation, Vespertilionidae, phylogenomics,
- 37 UCE, ultraconserved elements, reticulation
- 38

#### 39 Introduction

The genus Myotis (Order Chiroptera, Family Vespertilionidae) contains more than 100 species that 40 41 originated during the last 10-15 million years (Stadelmann, et al. 2007), making it one of the most 42 successful, extant, mammalian species radiations. Members of *Myotis* are distributed worldwide, 43 excluding polar regions, and generally share a similar ecological niche: aerial insectivory. Myotis species 44 often exhibit little morphological differentiation and, as a result, the rate of cryptic speciation within the 45 genus is thought to be high. For example, specimens identified as M. nigricans and M. albescens form 46 multiple paraphyletic lineages distributed throughout the phylogeny of Neotropical Myotis (Larsen, et al. 2012). 47

48 Confounding matters, the morphological variation that exists is often a poor indicator of species-49 level relationships. Early classifications of Myotis identified three major morphotypes (Findley 1972); each 50 were assumed to be monophyletic and were recognized at the subgeneric level (Simmons 2005). 51 Subsequent phylogenetic analyses of the mitochondrial cytochrome-b (cytb) gene recovered paraphyletic 52 origins of the morphologically defined subgenera, suggesting convergent evolution in Myotis (Ruedi and 53 Mayer 2001). These same analyses demonstrated that geography was a better predictor of phylogenetic 54 relationship than morphology (Ruedi and Mayer 2001; Stadelmann, et al. 2007). Generally, Myotis 55 phylogenies from mitochondrial data contain a single bifurcation at the base of the tree that splits Old 56 World from New World species. An additional bifurcation within New World species separates Nearctic 57 (NA) from Neotropical (NT) species. The NA/NT bifurcation is not absolute, with at least five NA species 58 located in the Neotropics and vice versa. The Old World/New World bifurcation is stricter, with only two 59 Old World species, *M. brandtii* and *M. gracilis*, present in the New World clade.

60 The ability of mitochondrial markers to resolve a well-supported topology does not guarantee 61 that the mitochondrial tree represents the species tree (for examples see Willis, et al. 2014; Li, et al. 2016; 62 Leavitt, et al. 2017). Despite containing 37 genes, the lack of recombination and uniparental inheritance 63 of the mitochondrion means that it is transmitted as a single genetic unit. This makes mitochondria susceptible to evolutionary processes that may cause its history to diverge from the history of the species 64 (Edwards and Bensch 2009). The most widely accepted phylogenies of Myotis rely heavily on 65 mitochondrial data and even phylogenies containing nuclear data demonstrate an over reliance on 66 67 mitochondrial markers for resolution. For example alignments of the nuclear RAG2 and mitochondrial 68 cytb contained 162 and 560 variable characters respectively (Stadelmann, et al. 2007). Phylogenetic 69 analyses of *RAG2* in *Myotis* results in a tree primarily composed of polytomies (Stadelmann, et al. 2007). 70 Combining these two markers increases phylogenetic resolution, but the results are heavily influenced by

71 larger numbers of mitochondrial characters, potentially masking signal form the nuclear marker
72 (Stadelmann, et al. 2007; Larsen, et al. 2012; Ruedi, et al. 2013; Haynie, et al. 2016).

It is difficult to draw major conclusions from studies limited in the number of characters (Ruedi and Mayer 2001; Stadelmann, et al. 2007; Larsen, et al. 2012; Ruedi, et al. 2013; Haynie, et al. 2016) or taxa (Platt, et al. 2015). Current data seem to indicate that nuclear and mitochondrial markers recover similar topologies. Platt et al. (2015) generated a phylogeny based entirely on nuclear data using 85,028 shared transposable element insertions. Their results generally confirmed the mitochondrial phylogenies of *Myotis*, but only included seven taxa in their analysis. In order to fully resolve relationships and understand the *Myotis* radiation it is necessary to increase character and taxon sampling.

80 Recently, targeted sequencing methods have been developed that utilize baits to enrich and 81 sequence ultraconserved elements (UCEs; Faircloth, et al. 2012), and this method has resolved a number 82 of difficult phylogenetic problems (for examples see Crawford, et al. 2012; McCormack, et al. 2013; Green, 83 et al. 2014; Faircloth, et al. 2015; McGee, et al. 2016). Generally speaking, the conserved "core" of UCE 84 regions allows thousands of homologous loci to be enriched from divergent organismal genomes while the sequence that flanks the core UCE region contains a majority of phylogenetically informative sites – 85 allowing researchers to collect a large number of phylogenetically informative, homologous loci from 86 87 throughout the genome in a cost-effective and efficient manner. Broad sampling of the nuclear genome 88 should help to resolve a phylogeny without an over reliance on mitochondrial loci. In addition, increasing 89 the number of nuclear loci sampled from a handful of genes to thousands can recover accurate trees 90 despite high levels of incomplete lineage sorting (Maddison and Knowles 2006)

91 Here, we used targeted sequencing of UCEs to collect ~1.4 Mbp from  $\geq$ 3,600 nuclear loci in 37 92 taxa, primarily representing New World Myotis. Combinations of the UCE data were analyzed using 93 concatenation and tree summary methods to estimate the *Myotis* phylogeny. Analysis of the nuclear UCE 94 data recovered 295 trees representing 175 distinct topologies. The nuclear topologies were compared to 95 trees generated from full mitochondrial genomes to test for conflict between the two types of makers. Our results show that, despite the range of trees recovered from the nuclear data, nuclear and 96 97 mitochondrial markers always depict conflicting phylogenies. Given that the nuclear and mitochondrial 98 trees are distinct from one another it is necessary to reinvestigate conclusions made based solely on the 99 mitochondrial phylogeny.

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101 Results

We used targeted sequencing of UCEs to collect sequence data from 3,648 nuclear loci which we assembled into concatenated alignments as large as 1.37 Mb. In addition, we assembled mitochondrial genomes for most taxa. We then used the data to infer the phylogenetic history of New World *Myotis* in three phases: UCE and mitogenome assembly, initial phylogenetic analysis, extended phylogenetic analyses.

UCE and mitochondrial assembly and alignment – We averaged 3.29 million reads per sample 107 108 after demultiplexing. These reads were assembled into an average of 5,778 contigs per sample (min = 109 1,562 M. martiniquensis, max = 11,784 M. nigricans 3). Recovery of UCE loci varied across taxa. Of the 5,500 loci in the Amniote probe set, we successfully recovered 3,898 UCE loci, 3,648 loci from five or more 110 111 samples, 212 loci in all 37 samples (Table 2). On average, 3,332 UCE loci were recovered per sample, 112 ranging from 1,106 (M. martiniquensis) 4,008 (M. keaysi). Repetitive sequences, identified via 113 RepeatMasker searches, were minimal occupying less than 0.02% of sites across all UCE alignments. 114 Sequence coverage of the mitochondrial genomes averaged 58x (range >1x - 297x). Mitochondrial 115 genome assemblies varied in quality. Some were almost entirely complete while others were missing sections. We found three premature stop codons in mtDNA protein coding genes. Subsequent manual 116 117 alignment and validation suggested that these regions were miscalled by MitoBim, and we corrected the 118 errors prior to analysis.

119 Initial phylogenetic analyses – Initial analysis of the nuclear data used loci that were present in 20 120 or more taxa. This resulted in an alignment of 1,144,471 bp from 2,890 nuclear loci containing 35,284 121 parsimony informative characters. The 2,890 loci were split into 27 partitions as recommended by 122 PartitionFinder (Lanfear, et al. 2012). Maximum likelihood and Bayesian analyses recovered the same 123 topology and found similar support for most nodes (Figure 1A). Maximum likelihood analysis recovered 124 100% support for 31 of 35 bipartitions, and 33 bipartitions were present in  $\geq$ 98% bootstrap replicates. 125 Nodes with the least support were still present in 86% - 88% of bootstrap replicates. After 50 bootstrap 126 replicates the average weighted Robinson-Foulds distance between replicate sets was less than 0.23% 127 (Pattengale, et al. 2009). Bayesian analysis recovered an identical topology, with the only difference being 128 that all bipartitions were supported with a clade probability value of >0.99. Visual inspection of the 129 parameter files in Tracer v1.6 showed good sampling with a likelihood score of -2.419 X 10<sup>-6</sup> and an effective sample size (ESS) of 637 for the likelihood parameter. All other parameters had effective sample 130 131 sizes greater than 500. The average standard deviation of split frequencies (ASDF) across all runs was less 132 than 1% after 5,000 generations.

Thirty-seven mitochondrial protein coding, rRNA and tRNA genes were concatenated into single 133 alignment of 15,520 bp containing 5,007 informative characters. Alignments for 30 samples were ≥ 90% 134 135 complete, and alignments for five samples were 68-84% complete. Only 21% and 50% of nucleotide positions were present in the *M. albescens*<sup>3</sup> (TK 61766) and *Myotis levis* alignments. Maximum likelihood 136 137 and Bayesian analyses of the mitochondrial data recovered similar topologies (Figure 1B), varying in the 138 M. thysanodes, M. evotis, and M. keeni relationships. Neither method recovered significant support for 139 these relationships. Bootstrap replicates of the maximum likelihood analysis meet the stopping criterion after the first 50 of 10,000 replicates (average weighted Robinson-Foulds value = 2.28%). The RAXML 140 141 mitochondrial phylogeny was well supported with 29 of 35 nodes present in  $\geq$  96% of bootstrap replicates. 142 The remaining six nodes were present in 47% to 70% of bootstrap replicates. Bayesian analysis of the 143 mitochondrial data reached convergence, defined as an ASDF of <1%, after the first 424,500 of one million 144 generations. The final ASDF, after discarding 25% of samples, was 0.49%. The trace files from all four 145 independent runs shows proper mixing of samples and the effective sample size for all parameters was greater than 200. The log likelihood score for the Bayesian mitochondrial tree was -1.205x10<sup>-5</sup> with an ESS 146 of 1,446. In all, posterior probabilities were lower than the bound established as significant ( $\geq 0.95$ ) for 147 148 five nodes.

149 Mitochondrial and nuclear analyses recovered different topologies (Figure 1). We stripped branch 150 lengths from all trees and compared the topologies using an approximately unbiased test to determine 151 whether differences in the tree represented conflicting signals in the marker sets. When the nuclear data 152 is constrained to the mitochondrial tree (*p*-value =  $1 \times 10^{-66}$ ) or *vice versa* (*p*-value =  $2 \times 10^{-5}$ ), likelihood 153 scores are significantly worse than expected given similar evolutionary histories. These results reject the 154 hypothesis that the mitochondrial and nuclear UCE phylogenies reflect similar evolutionary histories.

155 Extended phylogenetic analyses - Many factors can bias phylogenetic analyses resulting in inaccurate trees 156 (Sanderson and Shaffer 2002). Rather than assuming our initial nuclear UCE tree was an accurate estimate 157 of the phylogenetic relationships of *Myotis*, we wanted to build a range of plausible topologies from the 158 nuclear UCE data. To do this we reanalyzed the nuclear UCE data set with minor deviations in locus 159 sampling, partitioning, inference method, etc. In all this effort resulted in 291 unique phylogenetic 160 analyses. Individual results or topologies are not the focus of these analyses. Rather, the goal was to 161 recover as many, reasonable, nuclear UCE topologies as possible in an effort to account for phylogenetic 162 uncertainty not present in the initial analysis and to compare the range of nuclear UCE trees to the 163 mitochondrial genome tree.

164 We investigated the effects of matrix composition (or completeness) on our phylogenetic 165 inference by generating 10 different alignments having levels of matrix completeness spanning 15-95% at 166 10% intervals and including a final matrix of 100% completeness. Loci in these alignments were partitioned 167 using three separate schemes: all loci were partitioned individually, loci were unpartitioned, or loci were 168 combined into optimum partitions using PartitionFinder. The result was 10 different alignments with three 169 partitioning schemes each. These were analyzed using Bayesian and maximum likelihood methods. Due 170 to computational limits we abandoned the fully partitioned, Bayesian analyses. The length, number of 171 loci, and optimum number of partitions per alignment is shown in Table 2. Bootstrap topologies stabilized 172 in 9 of 10 alignments after 50 replicates and all Bayesian runs converged in less than ten thousand 173 generations. In general, the same alignment produced the same topology regardless of inference method 174 or partitioning scheme with the only exception being the terminal relationships M. levis/M. albescens 175 clade in the optimum vs. unpartitioned Bayesian analysis of the 100% complete data matrix.

176 Trees were generated from data matrices incorporating loci of differing lengths (Hosner, et al. 177 2016). All 3,648 loci were ordered based on their length and split into nine bins of 365 loci and 1 bin of 363 loci, so that the first bin contained the 365 shortest loci, the second bin contained the 366<sup>th</sup> to the 178 179 731<sup>st</sup>, and so on. The number of informative characters per bin ranged from 1,115 to 6,995 and the 180 number of informative characters was correlated with average locus length (Supplemental Figure 2). On 181 average, only 2.6% of characters in each bin were parsimony-informative. Each of the ten length-based 182 alignments recovered slightly different topologies. Terminal relationships were generally stable across 183 analyses with the majority of differences between topologies found in the early bifurcations of the ingroup 184 (Myotis).

185 From the above analyses combining different matrix composition, inference method, partitioning, 186 and locus-length variants we observed that, in general, larger alignments produced well resolved 187 topologies with significant nodal support regardless of the phylogenetic method or partitioning scheme 188 used. On the other hand, re-analyses of smaller portions of the data were more likely to recover unique 189 topologies. Given that the overall goal of the extended analyses was to generate as many reasonable 190 nuclear UCE based topologies for comparison with the mitochondrial tree, we decided to randomly 191 sample small portions of the nuclear UCE loci to create alignments that are more likely to result in unique 192 topologies. We randomly subsampled the 3,648 enriched loci to create 100 unique data sets. Loci were 193 concatenated in each replicate data set and analyzed using maximum likelihood in RAxML. Of the 100 194 alignments analyzed, 80 unique topologies were generated (mean Robinson-Foulds distance = 4.3).

195 In addition to concatenated analyses, three summary-based species tree programs were used on 196 datasets of varying matrix-completeness (ASTRAL-II, ASTRID, SVDquartets). Normalized quartet scores 197 from ASTRAL-II (Mirarab and Warnow 2015) analyses were quite consistent with scores ranging from 198 0.540 to 0.553, and between 7,745,739 (100% complete 212 gene trees) and 63,042,410 (15% 3648 loci) 199 induced quartet gene trees. SVDquartets (Chifman and Kubatko 2014) sampled all 66,045 quartets. On 200 average, the total weight of incompatible guartets was 2.84%. Similar to the concatenated analysis, we 201 inferred coalescent-based species from the same 100 subsamples of 365 loci described above. Despite 202 being generated from the same underlying data, summary and concatenation methods only recovered 203 the same tree in one of 100 attempts.

204 Finally, we used weighted and unweighted statistical binning to combine individual gene trees 205 into supergenes, estimate the supergene phylogeny, and then infer the species tree from the supergene 206 trees. The 3,648 loci were combined into 528 binned loci with 480 bins containing seven loci each and 48 207 bins containing six loci each. Normalized quartet scores were 0.672 for the binned-unweighted and 0.673 208 for the binned-weighted ASTRAL-II analysis. Given the relative even distribution of loci into bins the 209 negligible difference in quartet/species tree discordance is expected. Both binning methods recovered 210 the same topology which was the same tree recovered in the initial nuclear UCE analyses and was the 211 most common topology observed across all analyses.

212 Topology comparisons – After rejecting topological congruence between the initial nuclear UCE and mitochondrial phylogenies we used various methods to re-analyze the nuclear UCE data in an effort 213 214 to identify alternative nuclear topologies. Topological congruence between the mitochondrial sequence 215 data and nuclear topologies resulting from the extended analyses were tested to see if any were 216 statistically congruent with the mitochondrial phylogeny of Myotis. Site-log likelihood scores for the 217 mitochondrial alignments when constrained to all 175 unique nuclear UCE topologies were generated in 218 RAxML and analyzed in Consel using the Shimodaira-Hasegawa (Shimodaira and Hasegawa 1999) and 219 approximately unbiased tests (Shimodaira 2002). In each case, the mitochondrial data produced 220 significantly worse likelihood scores, rejecting congruence between the nuclear UCE and mitochondrial 221 phylogenies (Supplemental Table 1).

When visualizing all topologies in tree space, nuclear trees co-localized and were distinct from mitochondrial topologies (Figure 2A). Pairwise comparisons of Robinson-Foulds symmetrical differences show that 98.75% of nuclear UCE vs. nuclear UCE (Figure 2B) trees are more similar to each other than the mitochondrial trees are to even the most similar nuclear UCE tree (Figure 2C). The most frequently observed topology was recovered in 45 of the 294 nuclear analyses and was identical to the tree recovered 227 in the initial nuclear UCE analysis (Figure 1a). Of the 45 analyses that recovered the most frequently 228 observed topology, 38 were Bayesian and RAxML searches that varied by matrix completeness and 229 partitioning scheme. The fact that these analyses recovered the same topology is expected given that they 230 are not independent. For example, a RAXML analysis of the 15% complete data set uses 1.26 Mb of the 231 1.38 Mb of data from the 25% complete dataset. These two alignments are 91% identical. Analyses that 232 directly varied the alignments and/or sampled less data (e.g. randomly sampling loci) were more likely to 233 generate unique topologies than the nested analyses described above. Of the 200 analyses that randomly 234 sampled UCE loci 164 unique topologies were observed. This implies that when analyses of large data sets 235 produce well-resolved trees with significant nodal support, sampling smaller portions of the data, may 236 provide a mechanism for creating phylogenetic uncertainty not represented by typical tree scoring 237 metrics. To account for the phylogenetic uncertainty present in our dataset, we generated a consensus 238 tree from all nuclear topologies using an 85% threshold to resolve bipartitions (Figure 3).

#### 239 Discussion

240 We generated phylogenies from 3,648 UCE loci and mitochondrial genomes of 35 Myotis bats. 241 Initial analyses of the mitochondrial and nuclear UCE phylogenies recovered distinct topologies (Figure 1). 242 Rather than rejecting concordance between the two data types from a single analysis we took steps to re-243 analyze the nuclear UCE data in an effort to generate as many viable nuclear topologies as possible. We 244 recovered 175 unique nuclear topologies using multiple methodologies, sampling strategies, and 245 parameters. None of these nuclear topologies were similar to the topology produced from the 246 mitochondrial data suggesting that nuclear UCE loci and the mitochondrial genomes of Myotis have 247 distinct evolutionary histories. The conflict between the mitochondrial and nuclear data may be driven by 248 error in phylogenetic estimation or may reflect genuine conflict between the two marker types (Degnan 249 and Rosenberg 2009; Huang, et al. 2010). We relied on multiple tree-inference methods (e.g. summary 250 vs. concatenation), manipulated phylogenetic parameters (e.g. partitioning strategy), and sampling 251 criteria (e.g. loci sampled in all taxa) to minimize the impacts of phylogenetic error on the data set. In most 252 cases, varying parameter or methodologies generated unique topologies, often due to rearrangements of 253 a few terminal taxa. M. volans and M. brandtii were often placed as either sister to the remaining NW 254 Myotis or as an early bifurcation between the NA and NT clades. M. vivesi was often found as sister to the 255 clade containing M. lucifugus, occultus and fortidens or as sister to the clade containing the NT Myotis.

Around 98.3% of all nuclear tree vs. nuclear tree comparisons contain fewer than 30 symmetric differences (Figure 2b) but there are no mitochondrial vs. nuclear tree comparisons with less than 30 symmetric differences (Figure 2c). Interestingly, the most common nuclear topology most often recovered 259 by concatenation analyses (Figure 1A). Summary methods failed to recover the most common nuclear 260 topology except when loci were binned together prior to gene tree estimation. Summary methods also 261 tended to recover more unique topologies than concatenation methods when analyzing data from the 262 same gene(s). For example, the random sample analyses recovered 80 unique topologies using 263 concatenation (RAxML) and 89 unique topologies with summary methods (ASTRAL-II). This likely has to 264 do with the limited number of informative characters per locus and by extension limited phylogenetic 265 signal per gene tree (Supplemental Figure 2). In these instances, limited phylogenetic signal per gene 266 would likely lead to increased opportunity for phylogenetic error in gene tree estimation. Further 267 supporting this idea, binning of compatible UCE loci may have indirectly increased phylogenetic signal 268 resulting in the same topology that many of the concatenation analyses recovered. No other 269 summary/coalescent method recovered this topology.

270 Previous work with UCE loci demonstrated that support for deep divergences varied based on the 271 number of loci examined (McCormack, et al. 2013). Further, bootstrap replicates and clade probability 272 values can be inaccurate metrics of nodal support (Douady, et al. 2003; Hedtke, et al. 2006). Varying the 273 input data and phylogenetic parameters can produce a range of reasonable nuclear topologies that may 274 be more useful than overreliance on a tree resulting from a one or two analyses. Here, by considering the 275 different topologies that result from various analyses (e.g. partitioning strategies, inference methods, 276 etc.), we can account for phylogenetic uncertainty better than considering a single nuclear or 277 mitochondrial topology alone.

The mitochondrial alignment constrained to any of the 175 nuclear topologies generated significantly worse likelihood scores than expected by chance (Supplemental Table 2) and a comparison of topologies in tree space shows that the mitochondrial topologies are unique from all nuclear topologies (Figure 2A). Pairwise tree distances demonstrate that all but the most divergent nuclear topologies are more similar to each other (Figure 2B) than any nuclear vs. mitochondrial comparison (Figure 2C). Despite the number of different analyses, the nuclear data never recover a topology that is similar, much less identical, to the mitochondrial topology.

285 Multiple studies have recovered effectively the same relationships among *Myotis* using 286 mitochondrial markers to the one presented here (Ruedi and Mayer 2001; Stadelmann, et al. 2007; 287 Roehrs, et al. 2010; Larsen, et al. 2012; Ruedi, et al. 2013; Haynie, et al. 2016). Thus, we are confident that 288 the mitochondrial phylogeny we recovered here, and by others, reflects the true mitochondrial tree. 289 However, the mitochondrial topology may not adequately reflect the species history, particularly when 290 considering the factors that cause incongruence between nuclear and mitochondrial gene trees. Possible causes of conflicting gene trees are horizontal transfer, gene duplication, introgressive hybridization, and
 incomplete lineage sorting. Some of these phenomena are more likely to have influenced the *Myotis* radiation than others.

294 Horizontal transfer of genes is thought to be rare in eukaryotes, but, vespertilionids in general 295 (Thomas, et al. 2011; Platt, et al. 2014), and Myotis (Pritham and Feschotte 2007; Ray, et al. 2007; Ray, et 296 al. 2008) in particular, have experienced horizontal transfer of DNA transposons. These events would not 297 be reflected in our phylogeny since repetitive sequences were removed prior to phylogenetic analyses. 298 More generally, gene duplications could create conflicting signal among individual UCE markers (ex. 299 comparing non-orthologous UCE loci), but the number of gene duplication events would have to be very 300 high to impact enough of the 3,648 UCE loci to confound the mitochondrial and nuclear phylogenies. 301 Further ruling out gene duplication events as the dominant cause of conflicting phylogenetic signal is the 302 fact that such events are likely depressed in *Myotis* as evidenced by their smaller genome size (~2.2 Gb) 303 and trend towards DNA loss (Kapusta, et al. 2017) combined with low rates of paralogy in UCEs general 304 (Derti, et al. 2006).

305 Introgressive hybridization and reticulation could significantly influence the phylogenies of *Myotis* 306 in a way that leads to conflicting signal between the nuclear and mitochondrial genomes (Sota 2002; 307 Good, et al. 2015). Hybridization in bats may be relatively common given their propensity to swarm at 308 cave entrances for breeding purposes. In European Myotis, swarming has allowed for high degrees of 309 hybridization between M. brandtii, M. mystacinus, and M. alcathoe (Bogdanowicz, et al. 2012). Further, 310 M. evotis, thysanodes, and keeni all experienced historical gene flow during their divergence (Carstens 311 and Dewey 2010; Morales, et al. 2016). It is also possible to explain the differences between the 312 mitochondrial and nuclear UCE phylogenies if *Myotis* experienced extensive incomplete lineage sorting 313 during their radiation. Two factors can influence the rate of lineage sorting, the fixation rate and the 314 speciation rate (Hudson, et al. 2002). Increasing the time to fixation and/or decreasing the amount of time 315 between cladogenic events will increase the likelihood of incomplete lineage sorting. *Myotis* are generally 316 long lived species (Dzeverin 2008) and underwent a rapid radiation between 5-10 MYA (Lack, et al. 2010), 317 suggesting that Myotis species are likely to experience higher levels of lineage sorting. The importance of these events -introgressive hybridization and incomplete lineage sorting- in driving the differences 318 319 between the mitochondrial and nuclear phylogenies cannot be determined with the current data.

Evolutionary history of *Myotis* – Our previous understanding of relationships within *Myotis* is heavily biased with mitochondrial data because nuclear markers were harder to collect and produced fewer informative sites (Ruedi and Mayer 2001; Stadelmann, et al. 2007; Lack, et al. 2010; Roehrs, et al. 2010; Larsen, et al. 2012; Ruedi, et al. 2013; Haynie, et al. 2016). Our UCE-based results indicate that nuclear trees vary substantially from the mitochondrial tree. Given that the nuclear and mitochondrial trees are different, we find it necessary to re-evaluate *Myotis* in the context of the nuclear data.

326 Paraphyly of *M. nigricans* and *M. albescens* was inferred from previous mitochondrial phylogenies 327 and confirmed in the UCE tree (Larsen, et al. 2012). Larsen et al (Larsen, et al. 2012) identified a minimum 328 of four and potentially twelve lineages in *M. albescens* and *M. nigricans*. Our sampling included four *M.* 329 albescens and three M. nigricans, compared to Larsen's 17 and 29 samples. Despite different 330 mitochondrial and nuclear topologies overall, our mitochondrial and nuclear phylogeny recovered the 331 same paraphyletic clade of three *M. albescens* samples and *M. levis*. Close relationships between these 332 taxa was found in previous work and expected. More importantly we did not find that M. albescens was 333 paraphyletic across much of NT Myotis. We also found that *M. nigricans* is monphlyletic in the nuclear 334 tree, but paraphyletic in the mitochondrial tree. These results from M. nigricans and M. albescens are 335 interesting but further inference is limited due to low sample sizes for these taxa.

336 The original subgeneric taxonomy of Myotis was based on three morphotypes that were later 337 shown to be the result of convergent evolution (Ruedi and Mayer 2001). If lineage-sorting affected the 338 mitochondrial phylogeny, it is possible that the morphotypes truly are monophyletic. However, 339 superimposing the previous subgeneric/morphological classification onto the species tree shows 340 interspersed distribution of morphotypes throughout even the most conservative nuclear tree (Figure 3). Many strongly supported terminal relationships link species with different morphotypes. Based on these 341 342 results, it appears that the three major morphotypes in Myotis are indeed a result of convergent 343 evolution, as suggested by previous work (Ruedi and Mayer 2001; Stadelmann, et al. 2007).

344 Among the more dramatic differences between the nuclear and mitochondrial topologies is the 345 placement of *M. volans* and *M. brandtii* as sister to all New World taxa by the nuclear data. Our 346 mitochondrial analyses place *M. volans* within a Nearctic clade and *M. brandtii* directly in-between the 347 Nearctic and Neotropical bifurcations as has been previously reported (Stadelmann, et al. 2007). Clade 348 probability values and bootstrap frequencies support these placements in trees from both data types. Our placement of *M. brandtii* as sister to all other New World *Myotis* more closely affiliates it with Old 349 World taxa. This make sense given that the *M. brandtii* distribution is also Old World. On the other hand, 350 351 a placement of *M. volans* sister to all New World taxa (and *M. brandtii*) in the nuclear tree is a significant 352 departure from previous work and, at first glance, does not make as much sense in a biogeographic 353 framework. M. volans is distributed across western and northwestern North America as far as far north

as Alaska. *M. brandtii* is distributed across much of Northern Europe. The key may lie in understanding a
third species, *M. gracilis*.

356 M. gracilis, along with M. brandtii, are the only two Myotis geographically distributed in the Old 357 World, but phylogenetically affiliated with the New World (Stadelmann, et al. 2007). If the sister 358 relationship between M. brandtii and M. gracilis (not sampled here) holds when nuclear data are 359 examined, then we can envision a scenario where M. gracilis, M. brandtii, and M. volans represent 360 speciation events that occurred during the transition of *Myotis* from the Old World to the New World. It 361 is important to remember that this interpretation relies on a fairly dramatic departure from the currently 362 accepted mitochondrial relationships of *M. volans* (represented here by a single sample) to other *Myotis* 363 species, and this hypothesis should be viewed as highly speculative. Increasing the number of Myotis 364 lineages sampled will shed additional light on this hypothesis.

Other taxa with conflicting positions between datasets include *M. lucifugus* + *M. occultus*, *M. fortidens*, and *M. vivesi*. In general, these relationships are characterized by very short branches and are the most likely to be affected by incomplete lineage sorting or limited phylogenetic information. This could explain the strong support with the mitochondrial tree compared to the nuclear species tree, while allowing for a number of nuclear loci to disagree with the species tree, as well.

370 There are a number of monophyletic groups identified with nuclear data (Fig. 1A) that exhibit 371 distinct biological characteristics. For example all of the long eared bats (septentrionalis, auriculus, evotis, thysanodes and keenii) represent a monophyletic group of higher elevation, forest-dwelling species that 372 373 glean insects off of surfaces (Fitch and Shump 1979; O'Farrell and Studier 1980; Warner 1982; Manning 374 and Jones 1989; Caceres and Barclay 2000). The group represented by fortidens, lucifuqus and occultus 375 represent a relatively long-haired form of Myotis. While having a distinct dental formula, fortidens was 376 historically described as a subspecies of M. lucifugus (Miller Jr and Allen 1928) and occultus has alternately 377 represented its own species or been considered a subspecies of *lucifuqus* (Hollister 1909; Valdez, et al. 378 1999; Piaggio, et al. 2002). The clade consisting of keavsi, oxyotus, ruber, simus, riparius, albescens and 379 diminutus represents a NT group of primarily woolly-haired bats (LaVal 1973). If the mitochondrial 380 genome has been subjected to phenomena that obscure the true species tree then these species groups, 381 along with their synapomorphic morphological features, can be reevaluated.

Conclusion - Relationships within *Myotis*, which until now have relied heavily on mitochondrial data, have served as the basis for species identification (Puechmaille, et al. 2012), evolutionary hypotheses (Simões, et al. 2007), and even conservation recommendations (Boyles and Storm 2007). Previous studies using nuclear data have largely been uninformative or utilized too few samples to draw 386 definitive conclusions. Trees estimated from ~3,650 nuclear loci and 295 different phylogenetic analyses 387 recovered 175 topologies, none of which are congruent with the mitochondrial phylogeny of Myotis. 388 Conflict between the mitochondrial and nuclear trees as well as among individual nuclear loci suggest that 389 the *Myotis* radiation may have been accompanied by high levels of incomplete lineage sorting and 390 possible hybridization. Rather than placing emphasis on the mitochondrial tree, it may be more 391 appropriate to consider it for what it really is: a single gene on par with a single UCE locus, albeit one with 392 many more phylogenetically informative characters. If true, then the mitochondrial genome is as likely to 393 reflect the true species tree as any UCE locus chosen at random. Large amounts of lineage sorting make 394 phylogenetic inference difficult and potentially impossible. Other phenomena such as reticulation, 395 hybridization, and introgression have likely influenced the genomes of Myotis and should be accounted 396 for in subsequent work. It is possible that the Myotis radiation is more accurately reflected as a hard 397 polytomy or a phylogenetic network rather than a strictly bifurcating phylogeny.

398

#### 399 Materials and Methods:

400 <u>Taxon Selection</u> - Taxa were selected to span the major phylogenetic break points with emphasis 401 on the Nearctic and Neotropical bifurcation as recovered in previous mitochondrial phylogenies 402 (Stadelmann, et al. 2007; Ruedi, et al. 2013) (Table 1). In addition, multiple individuals morphologically 403 identified as *M. nigricans* and *M. albescens* were included to test paraphyly as demonstrated by Larsen et 404 al. (2012). Three Old World species of *Myotis* and the outgroup, *E. fuscus*, were included to root 405 phylogenetic analyses. All field identifications were confirmed from voucher specimens. Information for 406 all specimens examined is available in Table 1.

407 UCE preparation, sequencing, and processing - Genomic DNA was extracted from 33 samples 408 using either a Qiagen DNEasy extraction kit or a phenol-chloroform/ethanol precipitation. DNA was 409 fragmented using the Bioruptor UCD-300 sonication device (Diagenode, Denville, NJ, USA). Libraries were 410 prepared using the Kapa Library Preparation Kit KR0453-v2.13 (Kapa Biosystems, Wilmington, MA, USA) following the manufacturer's instructions with five minor modifications. First, we used half volume 411 reactions. Second, subsequent to end repair, we added Sera-Mag Speedbeads (Thermo-Scientific, 412 413 Waltham, MA, USA; prepared according to (Glenn, et al. 2016)) at a ratio of 2.86:1 for end repair cleanup. Third, we ligated universal iTru y-yoke adapters (Glenn, et al. 2016) onto the genomic DNA. Fourth, 414 415 following adapter ligation, we performed one post-ligation cleanup followed by Dual-SPRI size selection 416 using 55 µL of speedbead buffer (22.5mM PEG, 1M NaCl) and 25 µL of Speedbeads. Finally, we performed 417 a PCR at 95 °C for 45 sec, then 14 cycles of 98 °C for 30 sec, 60 °C for 30 sec, 72 °C for 30 sec, then 72 °C for a 5 minute final extension and a 12 °C hold using iTru5 and iTru7 primers to produce Illumina TruSeqHT
compatible libraries (Glenn, et al. 2016).

420 Libraries were quantified on a Qubit 2.0 (Life Technologies) and 83 ng from each library was added 421 to create 5 pools of 6 or 7 libraries each. We then split the pools in two. One subsample was enriched for 422 UCE loci, the other was not. UCE loci in the enriched library pools were captured using Tetrapods 5K 423 version 1 baits from MYcroarray (Ann Arbor, MI, USA) following their MYbaits protocol v. 2.3.1 with 424 overnight incubations (Faircloth, et al. 2012). Enriched libraries were quantified with a Qubit and pooled 425 with other unrelated samples prior to sequencing on an Illumina HiSeq 3000 to produce paired-end reads 426 of  $\leq$  151 bases. The unenriched samples were sequenced on a separate run using a single lane of Illumina 427 HiSeq 2500. All samples were demultiplexed with Illumina software fastq2bcl. Reads were quality filtered 428 by removing any potential adapter sequence and trimming read ends once the average Phred quality over 429 a four base window score dropped below 20 using the Fastx toolkit (Gordon and Hannon 2010).

430 Quality filtered raw sequence reads were assembled into contigs using the Trinity assembler 431 (Grabherr, et al. 2011) and a minimum kmer coverage of 2, and we used Phyluce to identify those 432 assembled contigs that were UCE loci. We also harvested UCE loci from Eptesicus fuscus (GCA 000308155.1), Myotis brandtii (GCA 000412655.1), M. davidii (GCA 000327345.1), and M. 433 434 lucifuqus (GCF 000147115.1) genome assemblies using the Phyluce package (Faircloth 2016). Once 435 extracted from Trinity and genome assemblies, we aligned all UCE loci MAFFT (Katoh, et al. 2002), 436 trimmed the aligned data with gBlocks (Castresana 2000). Repetitive sequences (i. e. transposable 437 elements) in each alignment were identified with RepeatMasker and trimmed where found.

438 Mitochondrial genome assembly and annotation – Raw reads from the unenriched libraries were 439 used to generate mitochondrial genomes via MitoBim (Hahn, et al. 2013) in most cases. This program 440 used MIRA (B, et al. 1999) to map reads to a M. brandtii reference genome (Genbank accession number 441 KT210199.1). Alternative methods of mitochondrial genome assembly were used when MitoBim assembly 442 failed. These taxa include M. albescens (TK61766), M. albescens (TK 101723), M. albescens (RDS 7889), M. fortidens, M. keeni, M. melanorhinus, M. nigricans (QCAZ 9601), M. septentrionalis, M. simus, M. velifer, 443 and M. volans. For these samples, we first identified reads that were mitochondrial in origin using BLAST 444 searches against the *M. brandtii* mitochondrial genome (KT210199.1). Those reads were assembled using 445 Trinity v2.2.0 with the -single option. For taxa where we either could not assemble useable mitochondrial 446 447 genomes, we retrieved proxy data from GenBank as follows: M. brandtii (KT210199.1), E. fuscus 448 (KF111725.1), M. lucifugus (KP273591.1), and M. davidii (KM233172.1).

Once assembled, each mitogenome was annotated via MITOS (Bernt, et al. 2013). Annotated genes were manually validated via BLAST to confirm sequence identity and length. Protein coding genes were checked for stop codons using EMBOSS's transeq program (Rice, et al. 2000). When a stop codon was found, we used the raw reads to verify the sequence. We used BWA v0.7.12 (Li and Durbin 2009) to align the reads to the Mitobim assembled mitogenome to verify base calls from Mitobim. The protein coding rRNA and tRNA genes from each assembly were aligned using MUSCLE and concatenated into a single alignment for phylogenetic analyses, as described below.

456 Initial phylogenetic analyses - Initial phylogenies derived from UCE loci and mitochondrial coding 457 regions were generated using maximum likelihood and Bayesian methodologies. For the first round of 458 phylogenetic analyses all UCE loci present in 20 or more taxa were concatenated into a single alignment. 459 PartitionFinder v1.1.1 (Lanfear, et al. 2012) was used to identify and combine loci into an optimal 460 partitioning scheme using the hcluster heuristic algorithm. We assumed a GTR+F model for all loci (Darriba 461 and Posada 2015). Initial trees were generated using RAxML v7.4.1 (Stamatakis 2006) with linked branch 462 lengths RaxML (v8.1.3) was used to estimate and score the maximum likelihood phylogeny with the rapid 463 bootstrapping option and 1,000 bootstrap replicates. We define strongly supported bipartitions as those 464 present in 95-100% of bootstrap replicates and moderately supported bipartitions are present in 85-95% 465 of bipartitions (Wiens, et al. 2008). A Bayesian phylogeny was generated with the MPI version of ExaBayes 466 (v1.4.1) using two independent runs of 4 chains each. ExaBayes runs were terminated after 1 million 467 generations only if the average standard deviation of split frequencies was less than 0.01. The first 25% of 468 samples were discarded after which every 100<sup>th</sup> generation was sampled. The "-M 3" option was used to 469 reduce the memory footprint of all ExaBayes runs. Proper sampling, post burn-in was inspected via Tracer 470 v1.6. (Rambaut, et al. 2014). Effective sample sizes greater than 200 were considered acceptable. 471 Posterior probability values greater than 95% were considered to be significant.

472 The mitochondrial phylogeny was generated using methods similar to those described above with 473 the following exceptions. All 37 coding regions, including protein coding genes, tRNA and rRNAs, were 474 concatenated into a single alignment. Genes were partitioned individually except in the instances where two genes overlapped. These regions were partitioned separately from the individual genes resulting in 475 476 three partitions for the two genes: a partition for gene A, a partition for gene B, and a partition for the overlapping nucleotides of gene A and B. The fast bootstrapping search in RAxML was run for 10K 477 478 replicates and Bayesian analyses were performed across four independent runs with four chains of ten 479 million generations.

The mitochondrial and nuclear phylogenies were compared to each other to identify discordance between marker types. Site-log likelihood scores were calculated for the mitochondrial alignments when constrained to the nuclear tree while the nuclear alignment was scored against the mitochondrial tree using RAxML. Model parameters were then recalculated for each constrained alignment (-f G). Site-log likelihood scores were analyzed using CONSEL (Shimodaira and Hasegawa 2001) and compared using the approximately unbiased test (Shimodaira 2002). P-values less than 0.05 were used to indicate that the trees produced by mitochondrial or nuclear data differed.

487 *Extended phylogenetic analyses* - Many factors can influence phylogenetic inference. To reduce 488 the likelihood of any single factor influencing the results, we re-examined the nuclear UCE dataset using 489 the strategies described below. A flow chart of analyses is available in Supplemental Figure 1.

490 Aligned UCE loci were binned based on the number of taxa represented in the alignment 491 (phyluce align get only loci with min taxa; Faircloth 2016), or degree of completeness. Groups 492 included loci present in 100% (number of specimens (n) = 37), 95% (n = 35), 85% (n = 31), 75% (n = 27), 493 65% (n = 24), 55% (n = 20), 45% (n = 16), 35% (n = 12), 25% (n = 9), and 15% (n = 5) of specimens examined. 494 These 10 groups were non-exclusive, so a locus that was assembled in all specimens (100% complete) 495 would also be included with loci present in only 55% of specimens. On the other hand, a locus found in 496 only 55% of specimens would not be included in the 100% complete data set. Each set of UCE alignments 497 was concatenated using phyluce align format nexus files for raxml and a nexus character block was 498 created using the phyluce align format nexus files for raxml -charsets option. These datasets then 499 served as the basis for downstream phylogenetic analyses. For example, when a partitioning methodology 500 (discussed below) was tested, it was performed for each of the 100%, 95%, 85%, etc. alignments. In 501 addition to partitioning schemes, the effect of missing data was examined using Bayesian and maximum 502 likelihood methods.

503 Concatenated alignments were analyzed using three different partitioning schemes. 504 Unpartitioned alignments were simply concatenated UCE loci treated as a single genetic unit (No 505 Partitions). Fully partitioned alignments were concatenated alignments of UCE loci that were partitioned 506 by locus (All Partitions). Finally, PartitionFinder v1.1.1 (Lanfear, et al. 2012) was used to combine individual 507 loci into an optimal partitioning scheme (Optimal Partitions). Rather than searching for best-fit 508 substitution models for each UCE locus or partition, the GTR+F model of sequence evolution was assigned 509 to all loci (Darriba and Posada 2015). Initial trees for PartitionFinder were generated using RAxML v7.4.1 510 (Stamatakis 2006) with linked branch lengths. Partitioning schemes were heuristically searched using the 511 hcluster algorithm.

512 Maximum likelihood trees were inferred for the concatenated alignments using RAxML v8.1.3 513 (Stamatakis 2014). The three partitioning schemes (described above) were applied to each analysis. The 514 best scoring (lowest -InL) tree from each dataset was identified from 100 random starting trees and 515 bootstrapped 100 times using the GTR+F in both cases. The autoMRE function in RAxML v8.1.3 was used 516 to determine the need for additional bootstrap replicates beyond the initial 100 (Pattengale, et al. 2009). 517 A stopping criterion was set a priori if the weighted Robinson-Foulds distance was less than 5% in 95% of 518 random permutations of computed bootstrap replicates (Pattengale, et al. 2009). If necessary, an 519 additional 100 bootstrap replicates were computed until the convergence stopping criteria were met. 520 Finally, bipartition frequencies of bootstrap replicates were drawn onto the best scoring tree from the 521 initial RAxML searches for each of the respective data sets.

522 Bayesian analyses were conducted using ExaBayes v1.4.1 (Aberer, et al. 2014). For all Bayesian 523 analyses four independent runs of four chains each were run in parallel for a minimum of one hundred 524 thousand generations sampling every thousandth generation and applying a GTR+F substitution model 525 for each partition. Two of the partitioning schemes (described above) were used for each analysis: No 526 Partitions and Optimal Partitions. After one hundred thousand generations, analyses continued until the 527 standard deviation of the split frequency between chains was less than 0.01. An extended majority rule 528 consensus tree was created from all trees after the first 25% of trees were discarded using TreeAnnotator 529 v1.7.0 (Rambaut and Drummond 2013) and parameter estimates across all runs were calculated with 530 Tracer v1.6 (Rambaut, et al. 2014).

531 Sampling loci by number of informative characters – Previous coalescent analyses of UCE data 532 have shown that sub-sampling the most informative loci can result in different topologies (Meiklejohn, et 533 al. 2016). Under these assumptions, UCE loci were sorted into ten groups based on their length and the 534 predicted correlation between length and number of informative characters was confirmed 535 (Supplemental Figure 2). UCE loci in the same size cohort were combined into a single alignment. Rather 536 than using coalescent based analyses we used concatenation of UCE loci to identify different topologies 537 based on length. UCE loci were individually partitioned and the maximum likelihood tree was estimated 538 with the rapid bootstrapping option in RaxML (bootstrap replicates = 100) using the GTR+ $\Gamma$  substitution 539 model.

540 *Random sampling of loci* – In large phylogenetic analyses, systematic error can result in highly 541 supported, but incorrect topologies as a result of compounding non-phylogenetic signal (Rodríguez-542 Ezpeleta, et al. 2007). By randomly reducing the dataset and replicating the ML analyses, we can reduce 543 the potential effects of compounding error. Roughly 10% of the dataset, 365 loci, were randomly sampled and concatenated to create 100 new alignments. ML methods were similar to those used when samplingloci by the number of informative characters.

546 *Summary methods* – Gene trees for individual UCE loci recovered in five or more taxa were 547 inferred using the GTR+Γ substitution model and fast bootstrapping (-f a) option in RAxML (replicates = 548 1,000). In general, gene trees were classified based on the degree of completeness (i.e. number of taxa 549 represented) similar to the way we treated individuals as described above.

550 Species trees were estimated and bootstrapped using three different programs. ASTRAL-II v4.10 551 (Mirarab and Warnow 2015) was used to build a summary tree. Support values for bipartitions in the tree 552 were generated from 100 bootstrap replicates using site as well as site and locus resampling (Seo 2008). 553 Species trees were estimated from ASTRID v1.4 (Vachaspati and Warnow 2015) using bionj and 554 bootstrapped for 100 replicates. SVDquartets (Chifman and Kubatko 2014), as implemented in PAUP 555 v4.0a150 (Swofford 2003), was used to estimate a species trees from a random subset of 200,000 quartets 556 and 1,000 bootstrap replicates.

557 Errors in gene tree estimation may reduce the accuracy of summary methods (Liu, et al. 2009; 558 Leaché and Rannala 2011; DeGiorgio and Degnan 2014; Mirarab, et al. 2016). We used weighted (Bayzid, 559 et al. 2015) and unweighted (Mirarab, et al. 2014) statistical binning to combine gene trees into 560 compatible supergenes using the pipeline available in Bayzid et al. (Bayzid, et al. 2015). The gene trees 561 used for the summary tree methods described above were used rather than re-estimating trees. 562 Bifurcations supported by more than 50% of the bootstrap replicates were retained for each gene tree. 563 Alignments from compatible trees were concatenated into a single supergene alignment. Trees for 564 supergenes were estimated using RAxML. The best trees for each supergene, as defined by log likelihood 565 score, were retained from 500 searches. Bipartition support was estimated from 500 bootstrap replicates. 566 For all analyses, the GTR+F model of substitution was used and each gene in the supergene alignment was 567 partitioned separately. The resulting supertrees were then used for species tree estimation using ASTRAL-568 II. For the unweighted analysis, all supertrees were included in the pool of trees. For the weighted analysis, 569 supertrees were weighted according to the number of genes combined in the supergene alignment. For 570 example, if a supergene was a composite of six genes, the supertree was present 6 times compared to a composite of five genes which would be represented only five times. Support for the weighted and 571 572 unweighted species trees was estimated by site and site and locus re-sampling (Seo 2008) for 100 573 bootstrap replicates in ASTRAL-II.

574 *Meta-analyses* - Trees recovered from all analyses were compared to each other in tree space. 575 Unweighted Robinson-Foulds distances were calculated among all trees. This distance matrix was

transformed into two dimensions using the stochastic CCA algorithm for nonlinear dimension reduction
in TreeScaper v1.09 (Huang, et al. 2016). Coordinates were then visualized in R using hexagonal binning
in the hexbin library v1.27.1 (Lewin-Koh 2011).

579 We compared the mitochondrial data to all nuclear trees. Branch lengths have different meanings based on the type of analysis. For example, ASTRAL-II branch lengths are representative of coalescent 580 581 units. ASTRID doesn't even calculate branch lengths. For accurate tree comparisons, branch lengths were 582 stripped from all trees using regular expressions and Sed. Site log-likelihood scores were calculated for 583 each nuclear tree, without branch lengths, and using the mitochondrial alignment. Model parameters 584 were re-estimated for each tree. Site-log likelihood scores were compared with the approximately 585 unbiased (Shimodaira 2002) and Shimodaira-Hasegawa (Shimodaira and Hasegawa 1999) test in CONSEL 586 and values less than 0.05 were indicative of differences. Finally, we used RAxML to generate a 85% and 587 majority rule consensus trees from all nuclear trees.

588

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604

## 605 Figure and Table Captions

- 606 Figure 1. Comparison of nuclear and mitochondrial phylogenetic trees in *Myotis*. Bayesian trees
- 607 generated from (A) 2,890 nuclear UCE loci and (B) 37 mitochondrial protein, tRNA, and rRNA genes (B)
- 608 Posterior probability values greater than 0.95 are shown as a "\*". Values below branches are
- 609 percentages of maximum likelihood bootstrap replicates supporting that clade. Conflicting tips between
- 610 data types (nuclear vs. mitochondrial) are indicated with lines between the topologies. The
- 611 mitochondrial Bayesian and maximum likelihood trees resolved different relationships among *M*.
- 612 thysanodes, evotis, and keeni as indicated by the dotted lines. Species with more than one sample are
- designated with a superscript that is referenced in Table 1. Specimens derived from whole genome
- 614 alignments are designated with a superscript "G".

**Figure 2.** Differences between mitochondrial and nuclear topologies. (A) All trees recovered from

616 mitochondrial genes and the extended analysis of the nuclear data were visualized in tree space using

- 617 multidimensional scaling. Nuclear trees (green) were distinct from mitochondrial trees forming a large
- 618 cluster. Most nuclear trees were found within a limited region of tree space. Mitochondrial trees (blue)
- 619 were distinct from any of the 294 nuclear trees recovered. Tree vs tree comparisons show that most (B)
- 620 nuclear trees are more similar to each other than they are to (C) mitochondrial trees. Symmetric
- 621 differences are equal to twice the Robinson-Foulds distance between two trees.
- 622 **Figure 3.** Consensus tree. A consensus tree of New World Myotis was generated from 294 nuclear
- topologies with a threshold cutoff of 85%. Values shown above the branches represent the percentage
- of nuclear analyses that support a given bipartion. Previous subgeneric classifications based on
- 625 morphology are listed at each tip (*Myotis* "M", *Selysius* "S", *Leuconoe* "L". Biogeographic regions are
- 626 color coded. Species with more than one sample are designated with a superscript that is referenced in
- Table 1. Specimens derived from whole genome alignments are designated with a superscript "G". This
  consensus tree represents a very conservative estimate of the *Myotis* radiation.
- 629 **Table 1.** Specimens examined.
- 630 Collection abbreviations: Museum of Southwestern Biology (MSB), Museum of Vertebrate Zoology
- 631 (MVZ), Natural History Museum of Geneva (MHNG), Pontificia Universidad Catolica del Ecuador Museo
- de Zoologia (QCAZ), Texas Tech University Natural Science Research Laboratory (TK), University of Alaska
- 633 Museum of the North (UAM), University of Michigan Museum of Zoology (UMMZ)

Table 2. General alignment information. For a subset of analyses a series of alignments were generated
based on the number of taxa per locus. Thirty-seven taxa were examined so an alignment with all 37
taxa was considered 100% complete. Parsimony-informative characters make up a small portion of the
total alignment. The optimum partitioning scheme was calculated with PartitionFinder.

638 Sup Fig 1. Analytical flow chart. In the initial analysis (A) mitochondrial and nuclear data were analyzed

using Bayesian and maximum likelihood methods. Trees were compared using the approximately

640 unbiased and the Shimodaira-Hasegawa tests and determined to be conflicting. Extended analysis (B) of

641 the data used multiple methods and sampling strategies to generate 292 different phylogenetic

642 inferences. (C) All nuclear and mitochondrial trees were compared in tree space and with topological

tests. An 85% meta-consensus tree from all analyses was used to represent a conservative estimate of

644 the *Myotis* radiation. Trees from the extended analysis were compared to the mitochondrial trees using

645 the approximately unbiased and the Shimodaira-Hasegawa tests and determined to be conflicting.

646 **Sup Fig 2.** UCE loci sorted by length. The length of a UCE locus is correlated with the number of

647 phylogenetically informative characters. UCE loci were sorted by length and ten bins of alignments were

648 created so that the shortest loci were combined into one alignment, the next shortest set of loci were

649 combined ... etc. Parsimony informative characters made up a minor part of each alignment.

650 **Sup Table 1.** Tree topology tests. Trees were compared to each other using the approximately unbiased

and the Shimodaira-Hasegawa tests. Likelihood scores were calculated for the mitochondrial alignment

when constrained to all topologies (mitochondrial and nuclear) recovered herein. Alignment and tree

topology incompatibility were identified as p-values < 0.01.

654 **Sup File** -AllTrees.nex – Trees generated from all analyses in nexus format.

655 **Sup File** -uceLociAlignments.tgz – Fasta alignments of all UCE loci recovered in at least four taxa.

656 Sup File - UceGeneTrees.tgz – UCE gene trees for all loci recovered in at least four taxa. Trees were

657 inferred from 100 ML searches using RAxML.

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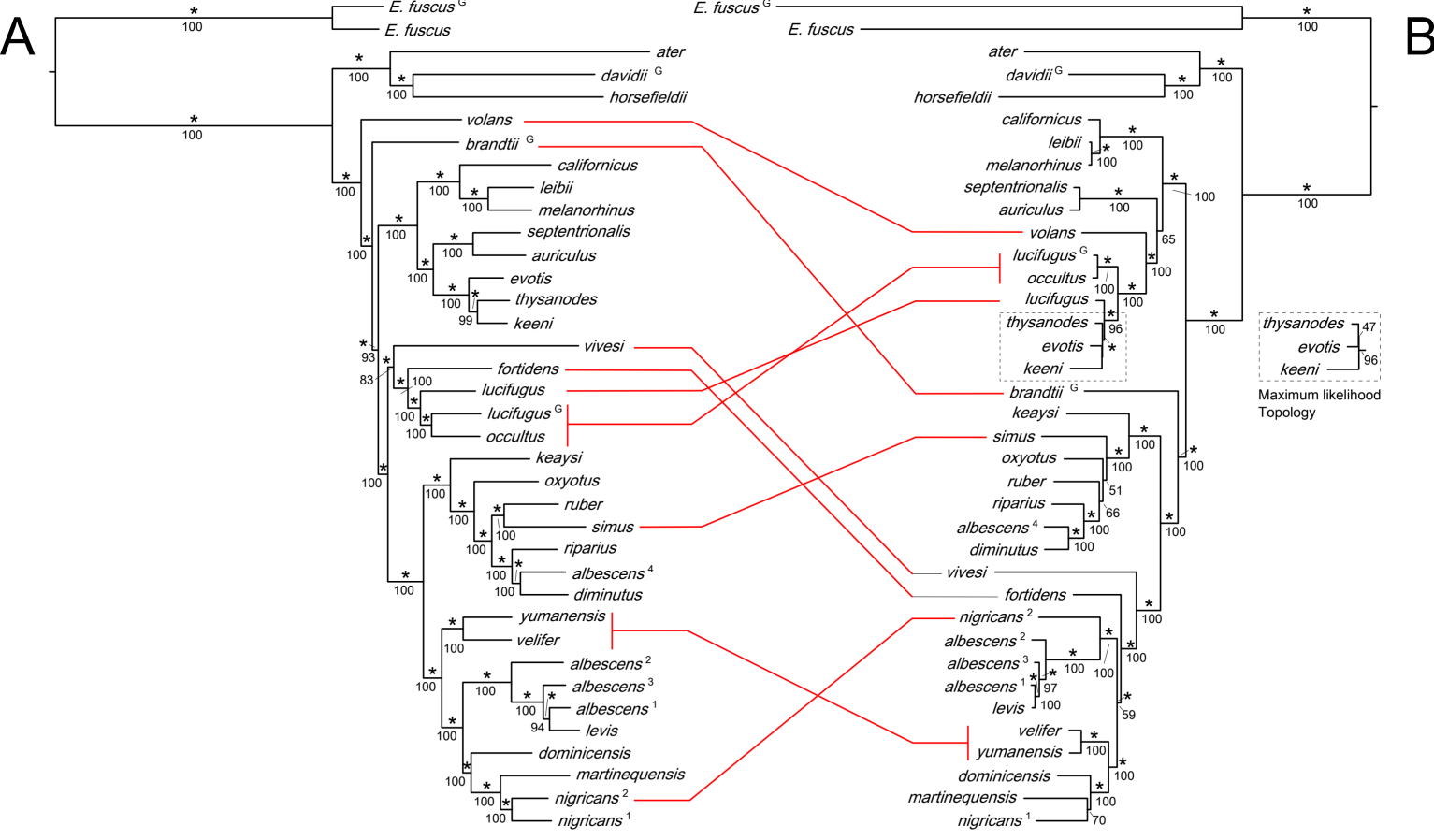
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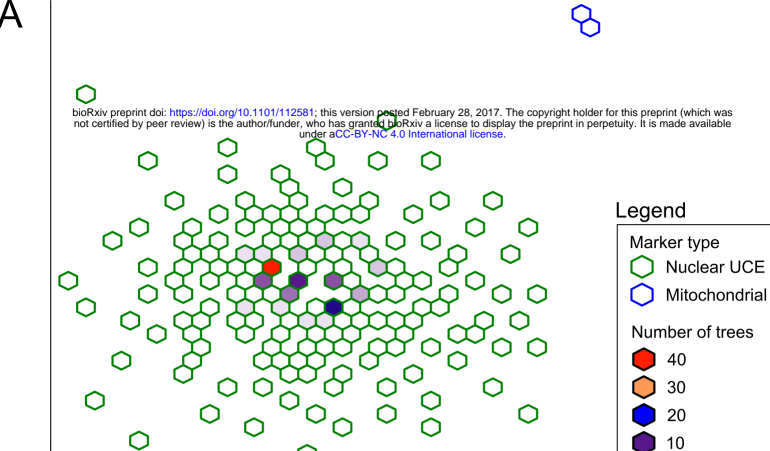
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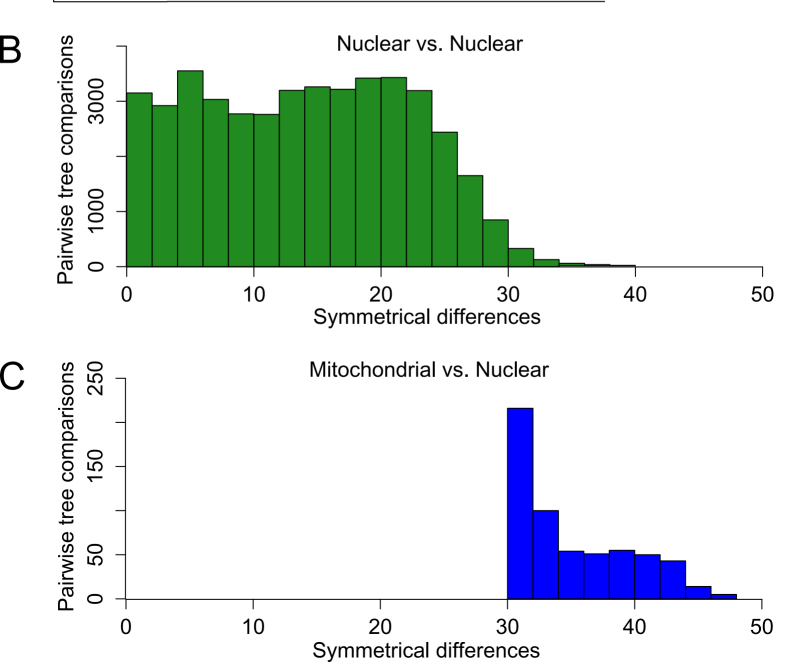
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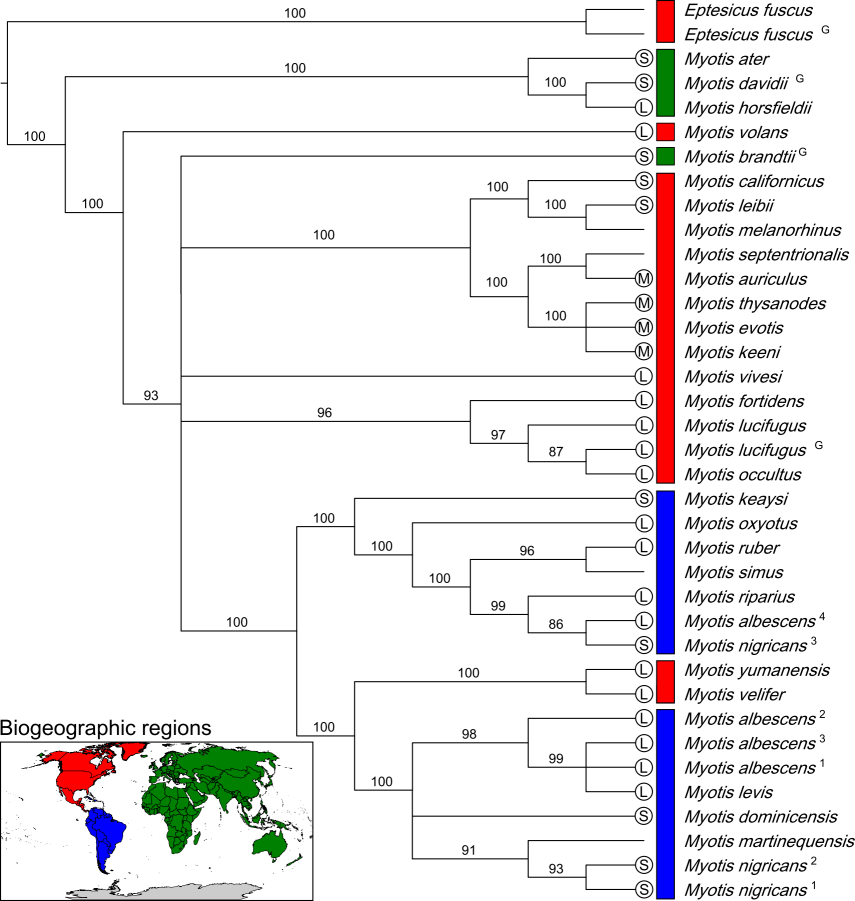
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### **Table 1 - Specimens Examined**

			Museum identification	UCE contig		Mitochondrial	
Genus	Specific epithet	Name herein	num.	Num. UCE Loci accession		genome accession	
Eptesicus	fuscus	fuscus	TK 178736	2,849	Pending	Pendin	
Eptesicus	fuscus	fuscus <sup>G</sup>	GCA_000308155.1	2,467	Pending	Pendin	
Myotis	albescens	albescens <sup>1</sup>	RDS 7889	2,764	Pending	Pendin	
Myotis	albescens	albescens <sup>2</sup>	QCAZ 9157	1,185	Pending	Pendin	
Myotis	albescens	albescens <sup>3</sup>	TK 61766	2,872	Pending	Pendin	
Myotis	albescens	albescens <sup>4</sup>	TK 101723	2,990	Pending	Pendin	
Myotis	atacamensis	atacamensis	M4430	2,774	Pending	Pendin	
Myotis	auriculus	auriculus	MSB 40883	2,229	Pending	Pendin	
, Myotis	brandtii	brandtii <sup>G</sup>	GCA_000412655.1	2,446	Pending		
, Myotis	septentrionalis	septentrionalis	 RDS 7705	2,916	Pending		
Myotis	californicus	californicus	UMMZ 175828	2,948	Pending		
Myotis	davidii	davidii <sup>G</sup>	GCA_000327345.1	2,450	Pending	Pendin	
Myotis	dominicensis	dominicensis	TK 15624	2,576	Pending	Pendin	
Myotis	evotis	evotis	MSB 47323	2,586	Pending	Pendin	
Nyotis	fortidens	fortidens	MSB 54941	2,791	Pending	Pendin	
Myotis	horsfieldii	horsefeldi	MHNG 1926.039	3,017	Pending	Pendin	
Nyotis	keaysi	keaysi	TK 13525	3,195	Pending	Pendin	
Myotis	keenii	keenii	UAM 113849	2,723	Pending	Pendin	
Nyotis	leibii	leibii	TK 24872	3,119	Pending	Pendin	
Myotis	levis	levis	RDS 7781	2,538	Pending	Pendin	
Myotis	lucifugus	lucifugus	MSB 46679	2,736	Pending	Pendin	
Myotis	lucifugus	lucifugus <sup>G</sup>	GCA_000147115.1	2,429	Pending	Pendin	
Myotis	martiniquensis	martiniquensis	TK 151413	856	Pending	Pendin	
Myotis	melanorhinus	melanorhinus	M8944	3,177	Pending	Pendin	
Myotis	nigricans	nigricans <sup>1</sup>	QCAZ 9601	2,854	Pending	Pendin	
Nyotis	nigricans	nigricans <sup>2</sup>	RDS 7791	3,159	Pending	Pendin	
Nyotis	nigricans or diminutus	nigricans <sup>3</sup>	QCAZ 9168	3,078	Pending	Pendin	
Myotis	occultus	occultus	MSB 121995	2,957	Pending	Pendin	
Myotis	oxyotus	oxyotus	UMMZ RCO1013	3,106	Pending	Pendin	
Nyotis	riparius	riparius	TK 145199	2,890	Pending	Pendin	
Nyotis	ruber	ruber	MVZ 185692	2,757	Pending	Pendin	
Nyotis	simus	simus	TK 22688	2,924	Pending	Pendin	
Nyotis	thysanodes	thysanodes	07LEP	2,821	Pending	Pendin	
Nyotis	velifer	velifer	MSB 70877	2,704	Pending	Pendin	
Myotis	vivesi	vivesi	MSB 42658	2,469	Pending	Pendin	
Myotis	volans	volans	MSB 40886	2,819	Pending	Pendin	
Myotis	yumanensis	yumanensis	RDS 7734	2,589	Pending	Pendin	

Collection abbreviations: Museum of Southwestern Biology (MSB), Museum of Vertebrate Zoology (MVZ), Natural History Museum of Geneva (MHNG), Pontificia Universidad Catolica del Ecuador Museo de Zoologia (QCAZ), Texas Tech University Natural Science Research Laboratory (TK), University of Alaska Museum of the North (UAM), University of Michigan Museum of Zoology (UMMZ). Samples beginning with "GCA\_" represent genome assemblies available through NCBI.<sup>®</sup>

P	Percent of		Parsimony-			
taxa per			informative	Variable	Alignment	Optimum
Min. num. of taxa	locus	Number of Loci	characters	uninformative	length	Partitions
37	100	212	3,480	4,778	112,125	6
35	95	1,193	18,288	24,189	575,321	15
31	85	2,034	29,179	41,711	903,903	16
27	75	2,481	33,031	48,732	1,041,099	20
24	65	2,668	34,373	51,148	1,091,620	27
20	55	2,890	35,284	53,200	1,144,471	27
16	45	3,064	36,539	54,782	1,187,492	31
12	35	3,232	37,259	56,453	1,227,093	34
9	25	3,379	37,894	57,672	1,260,248	37
5	15	3,648	38,718	62,588	1,377,262	31

### Table 2. Character information

General alignment information. For a subset of analyses a series of alignments were generated based on the number of taxa per locus. Thirty-seven taxa were examined so an alignment with all 37 taxa was considered 100% complete. Parsimony-informative characters make up a small portion of the total alignment. The optimum partitioning scheme was calculated with PartitionFinder.