Interrogating Phylogenetic Discordance Resolves Deep Splits in the Rapid Radiation of Old World Fruit Bats (Chiroptera: Pteropodidae)

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Abstract.—The family Pteropodidae (Old World fruit bats) comprises >200 species distributed across the Old World tropics and subtropics. Most pteropodids feed on fruit, suggesting an early origin of frugivory, although several lineages have shifted to nectar-based diets. Pteropodids are of exceptional conservation concern with >50% of species considered threatened, yet the systematics of this group has long been debated, with uncertainty surrounding early splits attributed to an ancient rapid diversification. Resolving the relationships among the main pteropodid lineages is essential if we are to fully understand their evolutionary distinctiveness, and the extent to which these bats have transitioned to nectar-feeding. Here we generated orthologous sequences for >1400 nuclear protein-coding genes (2.8 million base pairs) across 114 species from 43 genera of Old World fruit bats (53% and 96% of extant species- and genus-level diversity, respectively), and combined phylogenomic inference with filtering by information content to resolve systematic relationships among the major lineages. Concatenation and coalescent-based methods recovered three distinct backbone topologies that were not able to be reconciled by filtering via phylogenetic information content. Concordance analysis and gene genealogy interrogation show that one topology is consistently the best-supported, and that observed phylogenetic conflicts arise from both gene tree error and deep incomplete lineage sorting. In addition to resolving long-standing inconsistencies in the reported relationships among major lineages, we show that Old World fruit bats have likely undergone at least seven independent dietary transitions from frugivory to nectarivory. Finally, we use this phylogeny to identify and describe one new genus. [Chiroptera; coalescence; discordance; incomplete lineage sorting; nectar feeder; species tree; target enrichment.]

The family Pteropodidae (Old World fruit bats) contains 45 genera with >200 extant species distributed across the Old World tropics and subtropics (Simmons 2005; Simmons and Cirranello 2019). Despite performing key roles in forest ecosystems as agents of seed dispersal and pollination (e.g., Hodgkinson et al. 2003), these bats face widespread threats that include habitat loss, hunting, and persecution (e.g., Fujita and Tuttle 1991). Over half of the recognized species are of conservation concern, with 13 (6.5%) listed as near threatened, 37 (18.5%) as vulnerable, 16 (8%) as endangered, 8 (4%) as critically endangered, and 4 (2%) as recently extinct (IUCN Red List 2019). Past population declines have undoubtedly contributed to a scarcity of data for some species, several of which are known from just a few specimens.

Molecular phylogenies of pteropodids have revealed an early rapid radiation, with modern lineages arising in the late Oligocene ca 26-25 Ma (Teeling et al. 2005; Meredith et al. 2011; Amador et al. 2018). Although phylogenetic analyses have clarified many relationships, and also overturned earlier morphology-based systematics (Bergmans 1997) in favour of newly proposed higher taxa (e.g., Giannini and Simmons 2005; Almeida et al. 2009, 2020; Nesí et al. 2013), many basal nodes remain unresolved. As a result, the relationships among the subfamilies and their relative timing of divergence remain poorly understood (Giannini and Simmons 2003, 2005; Almeida et al. 2011, 2020).

Unlike their closest relatives, all pteropodids are herbivorous, with most species feeding on flowers. Their early transition to a carbohydrate-rich plant-based diet is expected to have posed physiological and metabolic challenges, and several studies have uncovered molecular adaptations linked to diet (Liu et al. 2012; Sharma et al. 2018; Jiao et al. 2019; Wang et al. 2020). In this context, of particular interest is the
increase topological concordance between inference of summary species coalescence methods as well as "PI" sites has been shown to improve the efficiency of loci based on proportion of parsimony-informative steps; for example, to reduce the effects of filtering steps. For large multigene data sets can be further enhanced (2017). Large multigene data sets can be further enhanced and branch lengths (Yang and Rannala 2012; "SRH")—can lead to incorrect inference of topologies due to ancestral polymorphisms and incomplete lineage sorting. Failure of phylogenetic models to capture historical signal, such that gene genealogies often differ due to model misspecification, by removing systematic biases due to model mis specification, by removing model partitions that violate SRH assumptions (Degnan and Rosenberg 2002). Hypothetical topologies selected by a majority ranks competing topologies based on likelihood scores using the approximately unbiased (AU) test (Shimodaira 2002). Failure of phylogenetic models to capture true evolutionary processes—which are typically assumed to be stationary, reversible, and homogeneous—"SRH"—can lead to incorrect inference of topologies and branch lengths (Yang and Rannala 2012; Jermiin et al. 2017).

With advances in sequencing technologies, a common strategy to resolve long-standing problematic nodes has been to analyze larger data sets (e.g., Jarvis et al. 2014; Prum et al. 2015; McGowen et al. 2020), typically using concatenation (Gatesy and Springer 2014) and/or multispecies coalescent (MSC)-based approaches (Edwards 2009; Liu et al. 2015; Jiang et al. 2020). The relative merits and disadvantages of both approaches have been strongly debated (Bryant and Hahn 2020). Generally, concatenation appears to perform poorly when ILS is high (Degnan and Rosenberg 2006; Mendes and Hahn 2018), whereas summary tree inferences based on the MSC are prone to errors and higher levels of uncertainty in the presence of uninformative loci (Gatesy and Springer 2014; Xi et al. 2014; Springer and Gatesy 2016; Brown and Thomson 2017). Large multigene data sets can be further enhanced by filtering steps; for example, to reduce the effects of uninformative loci and other sources of error. Removing loci based on proportion of parsimony-informative (PI) sites has been shown to improve the efficiency of summary species coalescence methods as well as increase topological concordance between inference methods (Hosner et al. 2016; Manthey et al. 2016; Blom et al. 2017). Filtering can also be used to reduce systematic error that, unlike stochastic error, cannot be remedied simply by increasing the size of a data set (Ho and Jermiin 2004; Jermiin et al. 2004; Philippe et al. 2005; Kumar et al. 2012; Duchene et al. 2017). Where filtering steps fail to resolve recalcitrant nodes, the gene genealogy interrogation (GGI) method (Arcila et al. 2017; Betancur-R et al. 2019) ranks competing topologies based on likelihood scores using the approximately unbiased (AU) test (Shimodaira 2002). Hypothetical topologies selected by a majority of genes are identified as the preferred resolution of the species tree for that node. This method is being recognized as a powerful tool for disentangling the relative contributions of information content, ILS, and data error in driving discordance.

Here we use target enrichment to capture >1400 nuclear protein-coding genes and the mitogenome (see Lemmon et al. 2012; Li et al. 2013) to perform the first phylogenomic reconstruction of Old World fruit bats. By applying node interrogation, we aim to resolve early splits in the context of a rapid radiation, and, in doing so, determine the higher-level systematic groups. As well as addressing long-standing inconsistencies in the reported relationships among multilocus genealogies, we aim to infer the number of dietary transitions between frugivory and nectarivory in this distinctively herbivorous family.
using DNeasy Blood and Tissue kits (Qiagen, USA) and Norgen’s FFPE DNA Purification kit (Norgen) for one formalin-fixed sample (Cynopterus horsfieldii). Genomic library preparation, target enrichment, and sequencing preparation were performed by Arbor Biosciences. The libraries were prepared using the MyBaits-3 custom Kit (Arbor Biosciences, USA) and sequencing conducted using 150 bp paired-end reads on Illumina HiSeq 4000 platform.

Data Assembly and Sequence Alignment
For each nuclear protein-coding gene, Illumina reads were processed to construct multiple sequence alignments using HyPhyMaker 1.6.4 (Fér and Schmickl 2018). Off-target intronic sequences were trimmed to reduce alignment errors; however, off-target mitochondrial (mt) reads were recovered (see Picardi and Pesole 2012) using the pipeline HyPhIPiper 1.3.1 (Johnson et al. 2016). We added published sequences for four mt loci (COI, CYTB, 12S, 16S) from five additional species (Aproteles balmerae, Casingtisrnis camponaenensis, Eponophorus anelli, Eponops buettikoferi, and Mirimiri acrodena) and two species (Eponophorus minimus and Hyysignathus monstrosus) for which we were unable to recover mitochondrial data sets due to sequence capture (Supplementary Table S9 available on Dryad). Each nuclear and mitochondrial locus was individually aligned using MAFFT (Katoh and Standley 2013). We excluded nuclear loci with <50% species completeness and >60% missing data. We integrated orthologous sequences (NCBI GenBank) of two outgroup species, Hipposideros armiger and Rhinolophus sinicus, representatives of the closely related families Hipposideridae and Rhinolophidae, respectively. We visually inspected and corrected alignments for indels, incorrect reading frames, and premature stop codons in BioEdit 7.0.5 (Hall 1999), and retained 1455 CDSs as our baseline nuclear data matrix (referred to as “all loci”) for downstream analyses.

Phylogenetic Analyses and Tests for Model Misspecification
We inferred phylogenetic trees using maximum likelihood (ML) with IQ-TREE (Nguyen et al. 2015; Chernomor et al. 2016; Hoang et al. 2018) on our concatenated nuclear “all loci” matrix as well as our complete mitochondrial data set. We then reduced the “all loci” data set using a filtering strategy based on the parsimony informativeness content within each locus (Meiklejohn et al. 2016). This resulted in three additional modified CDS data sets including loci with ≥18% (PI18, n = 1088 loci), 22% (PI22, n = 728 loci), and 27% (PI27, n = 341 loci) of PI variation in our baseline data set. Thus, four nuclear CDS data sets and one concatenated mtDNA data set were used in subsequent analyses. Details of loci assembly, alignment, and filtering strategy are available in the Supplementary Materials and Methods available on Dryad.

We also reduced our “all loci” data set by removing partitions that violated SRH assumptions. First, best-fit partitioning schemes and models of molecular evolution were determined with PartitionFinder 2.1.1 (Lanfear et al. 2016) using a partition-by-locus scheme for nuclear data sets and partition by locus and codon position for the mt data set. Then, for each partition in each nuclear data set (“all loci,” PI18, PI22, PI27), we tested for violation of the SRH assumptions. Although new non-SRH models have been recently developed to account for data that have evolved under non-SRH conditions (e.g., Zou et al. 2012; Woodhams et al. 2015), they are not easy to use and require large computation power (e.g., Betancur-R et al. 2013). An alternative method is to identify data that have evolved under non-SRH conditions before performing phylogenetic reconstruction. We used matched-pairs tests of homogeneity (Jeremim et al. 2017) implemented in IQ-TREE 2.0-rc1 (Naser-Khdour et al. 2019). Partitions rejecting the SRH assumptions (P < 0.05) were discarded before performing ML phylogenetic inferences.

In addition to concatenation analysis, coalescent-based species tree summary methods were performed with ASTRAL-III (Mirarab et al. 2014; Zhang et al. 2018) and MP-EST (Liu et al. 2010) on all gene trees, as well as the reduced PI data sets. Individual nuclear ML gene trees, based on the full CDS, inferred with IQ-TREE were used as input Brown posterior probabilities (Sayayri and Mirarab 2016), and quartet support for the main topology and alternative topologies were calculated with ASTRAL. For MP-EST analysis, 10 independent searches for the maximum pseudo-likelihood tree were performed. We estimated topological concordances among our concatenated and MSC trees by performing weighted Shimodaira-Hasegawa and AU tests in IQ-TREE with 1000 REL replicates (Kishino et al. 1990). Topologies were imported in Dendroscope 3.5.10 (Huson and Scornavacca 2012) and compared with each other as a tanglegram.

Tree Discordance and GGI
To summarize discordance among ML gene trees, we estimated bipartitions based on topology using PhyParts (Smith et al. 2015) with the function phypartsiecharts.py (https://github.com/mossmatters). Internode Certainty All (ICA) values were calculated by comparing the individual ML gene trees to both the estimated concatenated “all loci” tree and the ASTRAL species tree. We also quantified the gene concordance factor (gCF) and site concordance factor (sCF) in IQ-TREE 1.7-beta7 (Minh et al. 2020). The concatenated “all loci” tree was used as the reference tree with 100 quartets computed to estimate sCF values. Individual genes in concordance with specific branches of the reference tree were identified using the –cf-verbose option. Conflict was visualized using a cloumdogram generated via the DensiTree function in
the R package Phangorn v2.5.5 (Schliep 2011), with CDS alignments reduced to 15 species from the main subfamilies of the Pteropodidae.

To further interrogate discordance among recalcitrant backbone relationships, we tested whether observed differences among gene tree topologies were likely due to gene estimation error (limited phylogenetic signal content), or to biological conflict arising from ILS. For this, we applied a GGI approach to the key backbone nodes identified in our different analyses (nodes B–D; Fig. 1). We tested 105 alternative rooted topologies that only differed with respect to the relationships among the five main lineages of Pteropodidae: 1) Rousettinae + Eidolinae, 2) Cynopterinae, 3) Harpyionycterinae, 4) Macroglossusinae, and 5) Nycitimeninae + Pteropodinae + Notopterisinae (Bergmans 1997; Almeida et al. 2020). We constrained the ancestral node of each of these five clades in all analyses. In addition, we repeated this analysis for a topology in which these lineages formed a polytomy. We ran ML searches on each of our 1455 loci for these 105 hypotheses (152,775 constrained ML searches) (Fig. 2). To assess topology support, we compared site-wise likelihood scores across all 105 trees with the AU test in IQ-TREE, and ranked trees by $P$ value. Finally, we conducted summary species-tree analyses with ASTRAL using as input all rank 1 constrained trees (1455 gene trees), as well as with only the set of “rank 1” trees significantly better than the alternatives ($P < 0.05$; 54 gene trees).

Ancestral State Reconstruction of Diet

To reconstruct the evolutionary history of nectar-feeding, we performed ancestral state reconstructions of diet (frugivory vs. nectarivory) based on Topology 1 in which long-standing problematic nodes were resolved (see Result Fig. 1). For classifying nectar-feeding lineages, we followed Freeman (1995) supplemented with Bergmans (1989), Sheherazade et al. (2019), and Almeida et al. (2020). We used a Bayesian stochastic character mapping approach in the R package phytools version 0.7-47 (Revell 2012) with the function make.simmap (Bollback 2006). Akaike information criteria were used to assess which of the three available hierarchical models (ER, SYM, and ARD) was the most appropriate for our data. Analyses were then run using the best model for 10,000 simulation replicates, with all other options set as default.

RESULTS AND DISCUSSION

Using sequence capture, we successfully recovered an average of 1536.9 loci (99.4%) per sample ($n = 135$). Following manual examination of our alignments, we discarded 90 problematic loci, yielding a final data set of 1455 CDSs spanning 2,791,426 base pairs (“all loci” data set). Filtering the “all loci” data set on parsimony informativeness reduced the numbers of CDSs to 1088, 728, and 341 based on PI thresholds of $\geq 18\%$, $22\%$, and $26\%$, respectively. Using PartitionFinder 2, we also divided the 1455 loci into 1028 partitions, of which 473 partitions met SRH assumptions (555 failed) based on a matched-pairs test of symmetry (MPTS) (Table 1) and we included in our “SRH loci” data set. Finally, we recovered sequences from up to 15 mitochondrial loci, totaling 13,950 bp, with a mean of 106 taxa (min 65 and max 131) (Table 1; Supplementary Tables S1 and S4–S6 available on Dryad).

Resolution of the Basal Pteropodid Relationships

All phylogenetic trees inferred using concatenation recovered the majority of recognized subfamilies and tribes with high bootstrap support (>90), with the exception of the tribe Rousettini (see Implications for Systematics). At the same time, we found that the topology reconstructed from the “SRH loci” data set (Topology 1) differed from that of the “all loci” data set (Topology 2) with respect to the position of Macroglossusinae, suggesting that SRH violation may affect phylogenetic inference (Figs. 1 and 2; Supplementary Fig. S1 available on Dryad). However, one alternative explanation for this difference is that removing loci that failed the MPTS may lead to improved model adequacy and accuracy. PI filtering schemes applied to both data sets also supported Topologies 1 and 2 (Fig. 2; Supplementary Figs. S2–S4 available on Dryad). In contrast to concatenation, coalescent-based summary analyses of the complete data set using both ASTRAL and MP-EST recovered another distinct topology (Topology 3), although Topology 2 was recovered with ASTRAL under the PH18 and PH27 filtering schemes (Fig. 2; Table 1; Supplementary Figs. S5–S12 available on Dryad). Finally, phylogenetic reconstruction based on concatenated mitochondrial genes produced a fourth topology with most major subfamilies retained (Fig. 2, Topology 4; Supplementary Fig. S13 available on Dryad). To assess the relative statistical support for these four alternative topologies, we performed an AU test using the complete concatenated data set and found that Topology 1 was significantly most likely ($P < 0.05$, AU test) (Supplementary Table S7 available on Dryad). A potential driver of the observed topological discordance at the base of the tree is the presence of ILS. Thus, our finding that concatenation outperformed coalescent summary methods was somewhat unexpected given that the latter are often considered more robust than concatenation in the presence of high rates of ILS. However, coalescent methods have also been shown to be susceptible to the effects of gene tree estimation error, and simulations indicate that filtering to reduce this error is only effective in cases of low to moderate ILS (Molloy and Warnow 2018).

In severe cases of ILS, short branches can produce gene trees that conflict with the species tree more frequently than they agree with it, leading to a so-called “anomaly zone” (Degnan and Rosenberg 2006).
FIGURE 1. Phylogram based on maximum likelihood inference of the complete CDS data set (1455 loci). Backbone nodes are indicated by the letters A–F. Node bootstrap support values are ≥90% unless indicated otherwise. Pie charts on backbone nodes show the proportion of genes trees that support that clade (blue), the proportion that support the main alternative for that clade (green), the proportion that support the remaining alternatives (red), and the proportion of uninformative gene trees for that node (grey). Numbers next to the pie charts correspond to ICA, gCF, and sCF values for backbone nodes. For two species, A. bulmerae and M. acrodonta, the putative placement here using dashed lines is based on our mitochondrial tree (Supplementary Fig. S13 available on Dryad). The tribes, following Bergmans (1997) and Almeida et al. (2020), are listed to the right, and are depicted by a solid line if recovered as monophyletic in our analysis, or by a dashed line if not monophyletic. Green branches correspond to nectar-feeding pteropodid lineages. Representatives species from subfamilies and tribes across the family from top to bottom are: Eumops frenatus (Gambian epauletted fruit bat), Pteropus anschietae (Anchieta’s broad-faced fruit bat), Myonycteris brachycephala (São Tomé collared fruit bat), Strepsirhinius longicaudus (long-tailed fruit bat), R. aegyptiacus (Egyptian rousette), Hapalopterus balu merae (Bulmer’s fruit bat), Harpyionycteris celebensis (Sulawesi harpy fruit bat), Pteropus samoensis (Samoan flying fox), Pteropus pudicus (montane monkey-faced fruit bat), Melonycteris maculata (black-faced fruit bat), Nyctimene robinsoni (Queensland tube-nosed fruit bat), and Nyctimene carolinana (Halmaheran blossom bat). Illustrations by I. Velikov from Wilson and Mittermeier (2019), reproduced by permission of Lynx Edicions.
FIGURE 2. a) Phylogenetic relationships of the family Pteropodidae inferred using concatenation and species-tree coalescent-based methods on the complete CDS, the PI filtered CDS, and the concatenated mtDNA data sets. Backbone nodes are named as in Figure 1. Colored clades correspond to: Rousettinae, turquoise; Cynopterinae, orange; Harpiyonycterinae, green; Pteropodinae, yellow; Nyctimeninae, blue; and Macroglossinae, red. b) On the left, the DensiTree cloudogram inferred from nuclear genes for the reduced 16-taxon data set. For contrast, Topology 1 is shown in black. On the right, GGI results testing alternative hypotheses of the backbone relationships within the Pteropodidae. Plot: lines correspond to the cumulative number of genes (x-axis) supporting topology hypothesis with highest probability and their P values (y-axis) from the AU topology tests. Values above the dashed line indicate topology hypotheses that are significantly better than the alternatives (P < 0.05).

TABLE 1. Summary of data sets and methods of analyses used in the study

| Data set          | Nb of taxa | Nb of loci | Size (bp) | Nb partitions pass SRH | Concatenation approach | Species tree coalescent-based |
|-------------------|------------|------------|-----------|------------------------|------------------------|-----------------------------|
| Nuclear CDSs      | 137        | 1455       | 2,791,426 | 473                    | Topology 1             | Topology 2 |
| Complete          |            |            |           |                        |                        |                             |
| Parsimony informative filtering | 137       | 1086       | 2,100,753 | 314                    | Topology 1             | Topology 3 |
| PI ≥ 18%          |            |            |           |                        |                        |                             |
| PI ≥ 22%          | 137        | 728        | 1,405,743 | 201                    | Topology 2             | Topology 3 |
| PI ≥ 27%          | 137        | 341        | 631,544   | 103                    | Topology 2             | Topology 3 |
| Mitochondrial markers | 142       | 15         | 13,950    | NA                     | Topology 4             | NA |
| Complete          |            |            |           |                        |                        |                             |

Note: Number of taxa, loci, size for each data set, and number of partitions following the SRH assumptions are indicated, as well as the recovered topology by each method. Topology code follows Figure 2. NA = not applicable; Nb = number; ML = maximum likelihood.

To quantify the degree of topological discordance with respect to the backbone nodes A–F (Fig. 2), we applied two analyses of concordance based on Topology 1 and found comparable patterns of weak support. Specifically, concordance analysis using PhyParts showed low percentages of gene trees recovering nodes A–F (ICA: 0.09–0.10; Fig. 1; Supplementary Fig. S14 and Table S8 available on Dryad). We repeated the PhyParts concordance analysis using Topologies 2 and 3, and found that a similar number of genes supported these
backbone relationships (Fig. 2; Supplementary Figs. S15 and S16 available on Dryad). Concordance analyses performed on Topology 1 using IQ-TREE also indicated that nodes A–F were supported by few gene trees (gCF = 10.1; 25.57; mean gCF = 15.95) and a low proportion of parsimony-informative sites (sCF = 31.14–40.41; mean sCF = 35.68), despite very high bootstrap support (mean BS = 100) (Fig. 1). Furthermore, no alternative to Topology 1 emerged from the inferred gene trees, likely reflecting limited phylogenetic signal in individual loci (i.e. low gDF1 and gDF2 values; Supplementary Table S9 available on Dryad).

To evaluate the alternative topologies in greater depth, we reordered the branching pattern of the conflicting nodes B–D relative to the other early splits, and identified and compared support for 105 possible topologies using GGI (Supplementary Fig. S17 available on Dryad). We found that each of the 105 hypotheses was supported by at least one gene, and a large number of alternative topologies (n=36) received reasonable support (>15 loci). Although no topology showed strong support above all others, Topology 1 was consistently the best supported based on rank of P values (46 of 1455 loci; 3.36%). In contrast, Topologies 2 and 3 were supported by 39 (2.68%) and 31 (2.13%) loci, respectively, although both of these were outperformed by alternatives. For each of these three topologies, only a subset of loci was significantly better than the second-ranked locus (5, 4, and 3 loci, respectively). We also compared each of these topologies to a null hypothesis in which the five major lineages were placed as a polytomy at the crown node of the Pteropodidae (Fig. 2b; also see Alda et al. 2019), and found that none of the loci supported a resolved topology over the polytomy. On the other hand, more than a third of loci supported the polytomy over one of the alternatives (36%; 524 out of 1455 loci), with 74 showing significant support (Supplementary Table S10 available on Dryad). Finally, we performed coalescent analysis in ASTRAL using both the set of all gene trees of rank 1 (n=1455), and a subset of these that showed statistical support above rank 2 (P < 0.05; n=54). In both cases, the inferred tree was consistent with Topology 1 with the exception of the relationship of Notopterus, which in the latter analysis was recovered as a sister taxon to the Nyctimeneae (Supplementary Figs. S18 and S19 available on Dryad).

By revealing that multiple evolutionary scenarios receive some degree of support, these results highlight the long-standing difficulties that researchers have faced in resolving the early splits in this group of mammals. Our analyses of concordance and node interrogation suggest that such uncertainty likely stems from two main causes. First, many single genes (around a third of those examined) appear to contain too little information for resolving ancestral nodes separated by short branches, a pattern that characterizes the pteropodid tree. Second, for a larger proportion of genes sampled, we find that incongruence among loci with respect to the positions of backbone nodes falls into several discrete branching patterns, consistent with extensive ILS (Suh et al. 2015; Arcila et al. 2017; Alda et al. 2019). If ILS is indeed a major cause of this discordance, we might expect Topology 3 recovered with ASTRAL to be the best supported tree. Yet when coalescence-based analysis was repeated using only either the top-ranked topologies in GGI, or the set of gene trees that performed statistically better than alternatives, we recovered the relationships among nodes A–F seen in Topology 1. Thus, our findings differ from those of Alda et al. (2019), who found that coalescence-based reanalysis using statistically supported gene trees recovered the main coalescent-based species tree. We suggest that in the case of basal pteropodid lineages, although ILS is a major cause of discordance, gene tree error may have had a greater effect in misleading coalescence-based analyses. In most cases, Topology 1 remains the most highly supported bifurcating tree.

**Diversification and Diet**

Our analyses of a large molecular data set confirm the early rapid divergence of the major fruit bat lineages. The early diversification of this group has previously been attributed to a shift in feeding habits coupled with the potential loss of laryngeal echolocation (Amdor et al. 2018; Almeida et al. 2020), although such scenarios are speculation given the highly depauperate fossil record (e.g., Gunnell and Manfhi 2018). The ultimate drivers of this early diversification remain obscure, particularly since it appears to have occurred ~25 to 20 Ma, long after the radiation of angiosperms. One scenario is that the phenotypic changes associated with a transition to frugivory, combined with their ability to fly, means that early pteropodids would have been able to realize the ecological opportunities presented by an unevenly distributed and ephemeral resource such as fruiting trees. Indeed, frugivory was previously linked to high rates of diversification in primates (Gómez and Verdú 2012) and phyllostomid bats (Rojas et al. 2012), and it is noteworthy that these groups all exhibit visual and/or olfactory adaptations for detecting fruit (Zhao et al. 2009; Hayden et al. 2014; Wang et al. 2020).

Within the Pteropodidae, a best-fitting model (ER) of diet evolution supported seven independent transitions from frugivory to nectariivy (Fig. 3; Supplementary Fig. S21 available on Dryad). Nectar-feeding is relatively rare in vertebrates, and our understanding of its evolution in pteropodids has changed markedly with improved resolution of systematic relationships (e.g., Hollar and Springer 1997; Kirsch and Lapointe 1997; Alvarez et al. 1999; Juste et al. 1999). Historically, Andersen (1912) grouped most nectarivorous Old World fruit bats into the distinct subfamily Macroglossinae, comprising five genera (Eonycteris, Macroglossus, Meloemycteris, Notopterus, and Syconycteris) from Indo-Australasia plus a sixth genus (Megaloporus) endemic to Africa. Bergmans (1989) subsequently proposed that the little-known African genus Plenotes feeds on nectar based on evaluation of its palate and dentition, and
FIGURE 3. Tanglegram of the Topology 1 recovered in this study (see Table 1) on the left and the pteropodids topology presented in Almeida et al. 2011 (based on maximum likelihood tree, Fig. 2) on the right. Branches are colored by major clades: Rousettinae, turquoise; Eidolinae, dark purple; Cynopterinae, orange; Harpiyoncterinae, green; Pteropodinae, yellow; Nyctimeninae, blue; Notopterisinae, purple; and Macroglossinae, red. Grey lines connect taxa between trees. Genera only present in our study are indicated in bold and nectar feeder genera are colored in green. Bootstrap support is reported for nodes of the Almeida et al. 2011 topology. Dashed branches correspond to relationships with node support value $<50$. $^*$ indicates Micropteropus synonymized to Epomophorus. 

Almeida et al. (2020) split Nesonycteris from Melonycteris. Finally, the Australian species Pteropus scapulatus was proposed to be nectarivorous based on behavioral data, and other Pteropus species have also been observed to either feed on nectar occasionally (e.g., Aziz et al. 2017; Sheherazade et al. 2019) or to possess morphological features that are consistent with flower-visiting and nectarivory (Almeida et al. 2014).

In light of molecular phylogenies, the shared characters that led nectar-feeding bats to be considered a single clade—including reduced cheek teeth, elongated rostra, and elongated brushy tongues (also see Marshall 1983; Courts 1998)—are now recognized as convergent traits (e.g., Hollar and Springer 1997; Almeida et al. 2011). Yet while nectarivory is often considered to be a highly derived specialization, we find that several lineages of nectar-feeding pteropodids appear to have long evolutionary histories, branching off early in their respective subfamilies. Moreover, nectar-feeding bats are also seen in four of the eight major pteropodid subfamilies in our phylogeny, with species affiliations implying at least two origins in Africa, and up to five in Asia. Currently, it is not known whether such independent origins have involved convergent changes at the molecular level, however, the well-supported topology presented here will provide the requisite phylogenetic framework for determining the likely evolutionary position of molecular adaptations that might have contributed to the transitions to fruit- and nectar-based diets, and subsequent diversification of the Pteropodidae.

Implications for Systematics

The inferred relationships among the main pteropodid lineages differ substantially from published phylogenetic trees (e.g., Giannini and Simmons 2005; Almeida et al. 2011, 2020; Fig. 3). Given that these earlier analyses were based on eight loci ($\sim8000$ bp), representing $<1\%$ of the data set generated here ($\sim2,791,500$ bp), we attribute the greater resolution and support for basal nodes in our trees to the much larger volume of data.

We found that the tribes Pteralopini and Melonycterini grouped together (BS = 100, Fig. 1; Topologies 1–3) and not with other genera of the Pteropodini with which they have been traditionally allied (Bergmans 1997; Almeida et al. 2020). Of particular interest was the phylogenetic position of the genus Eidolon, which has long been problematic and unstable, even across molecular studies (e.g., Giannini and Simmons 2005;
Almeida et al. 2011, 2016; Hassanin et al. 2016). Eidolon is superficially similar to the other large members of the Pteropodini, yet a sister relationship between Eidolon and Pteropodini has never been strongly supported by either morphological characters (e.g., Giannini and Simmons 2005) or mtDNA (e.g., Hassanin et al. 2016). Our results indicate that Eidolon is in fact a sister taxon to the subfamily Rousettinae, with strong bootstrap support in all ML concatenated analyses based on nuclear DNA data (node A, Fig. 1), albeit with low concordance support across individual gene trees (n = 201, ICA = 0.99; Fig. 1). The exception was the mitochondrial analysis, which placed Eidolon at the base of the pteropodid tree, likely reflecting the lack of power with this marker (Supplementary Fig. S20 available on Dryad). Eidolon was previously included in Rousettinae by Bergmans (1997) based on morphological similarity, but Almeida et al. (2016) subsequently reclassified Eidolon in a new subfamily Eidolinae, which we also retain due to this taxon’s distinctiveness. By moving Eidolon to a grouping with Rousettinae, we place all continental African pteropodids in one well-supported clade (Eidolinae + Rousettinae), although several species within the Rousettinae (Myonycteris, species within Rousettus) are currently restricted to the Asian tropics.

Unexpectedly, and in disagreement with previous systematic assessments, the tribe Rousettini (sensu Almeida et al. 2016) was found to be paraphyletic. Specifically, we found that Rousettus celebensis—previously included here in a molecular analysis for the first time—did not group with other Rousettus species, but instead appeared as a sister taxon to a clade containing the tribes Stenonycterini, Myonycterini, Plerotini, and Epomophorini. This arrangement, which was strongly supported and recovered in the majority of individual gene trees (BS = 100, ICA = 0.38; Fig. 1), is intriguing because it suggests that R. celebensis represents a distinct lineage from Wallacea that has strong affiliations with African lineages. This is the second time that an endemic Sulawesi species has been removed from Rousettus, with Boneia previously elevated to a separate genus. Although Wallacea is a hotspot of endemism for fruit bats, with at least three other genera known only from Sulawesi and its offshore islands (Boneia, Neopteryx, and Thootherus), these genera are not sister taxa (also see Almeida et al. 2020), implying that pteropodids have likely crossed Wallace’s line several times in their evolutionary history.

The systematics of Rousettus has undergone several revisions, largely following genetic analyses. Boneia, Lissonycteris (now synonymized to Myonycteris; Nesi et al. 2013), and Stenonycteris were all previously classified as subgenera of Rousettus on the basis of specific morphological characters (Andersen 1912; Bergmans and Rozendaal 1988; Koopman 1994), but were removed subsequently (Bergmans 1997; Nesi et al. 2013). To our knowledge, specimens of R. celebensis described to date do not show any morphological characters that distinguish this taxon from the other Rousettus species (Andersen 1912; Bergmans and Rozendaal 1988), and thus in-depth morphological re-examination is warranted in light of our results. In this respect, the future inclusion of R. linduensis—another Sulawesi endemic that was recently described (Maryanto and Yani 2003)—and the only Rousettus missing from our study—will be important.

**Taxonomy**

The results of this study have implications for taxonomy, with R. celebensis specimen MSB 93154 (Supplementary Table S1 available on Dryad) found to represent a new monotypic genus of bat, described below for the first time.

**Pilonycteris gen. nov.** Nesi, Tsang, Simmons, McGowen & Rossiter

*Pilonycteris* Gray 1821

**Etymology.**—Named from the combination of the Latin pilo, hairy, from its distinctly longer fur, and the Greek nycteris, bat. The gender of the genus is masculine.

**Type species.**—*Pilonycteris celebensis* Andersen 1907 (holotype BMNH 1897.2.8; type locality: Mt Masarang, 3500 ft, Celebes).

**Diagnosis.**—Andersen (1912) noted that *Pilonycteris celebensis* is distinguishable from sympatric species *Rousettus amplexicaudatus*, *R. brachyotis* (now *R. amplexicaudatus brachyotis*), and *R. leschenaultii* by having a bony palate that is narrower posteriorly, last upper and lower premolar and molars that are unusually narrow, fur that is brighter, longer, and richer in color, a notopatagium (= unopatagium) that is partly or wholly hairy, and a generally smaller size but with digits proportionally longer than other sympatric *Rousettus* species. Rookmaaker and Bergmans (1981) further indicated that *P. celebensis* differs in hairiness, especially the absence of reduction of hair in the neck region, presence of a hairy notopatagium, and the greater length of its fur, which differentiates it from both *Rousettus amplexicaudatus* and *R. leschenaultii*. *Pilonycteris celebensis* also has a relatively longer rostrum, longer lower and upper tooth rows, and a smaller average distance between the two M2s. Particularly distinctive characters involve dimensions of the cheek teeth; *P. celebensis* has a longer P4, narrower M1, shorter and narrower M2, longer and narrower M3, narrower M2, and an M3 that is sometimes shorter and narrower than seen in *Rousettus amplexicaudatus* spp. (see Table 6 in Rookmaaker and Bergmans 1981).

**Contents.**—Monotypic, comprising only *P. celebensis* (Andersen 1907). May also contain *Rousettus linduensis*, but further analyses will be needed to confirm this.
AUTHOR'S CONTRIBUTIONS
N.N. and S.J.R. conceived of the study; V.N., A.L., A.T.S., S.A.R., S.W., A.T.H., J.J., C.A.P., F.J.B., C.M.T., B.K.L., and N.B.S supplied tissues; N.N. and G.T. compiled genes and designed baits with the assistance of MYcroarray (now Arbor Biosciences); N.N. performed laboratory work and all bioinformatics and phylogenetic analyses; N.N., M.R.M., and S.J.R. wrote the paper with contributions from the other authors.

SUPPLEMENTARY MATERIAL
Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.7mkc8pq5

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CONFLICT OF INTEREST
All authors declare no conflict of interest.

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