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RESEARCH ARTICLE



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Phylogenetic analyses reveal bat communities in Northwestern Mexico harbor a high diversity of novel cryptic ectoparasite species

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Abstract

Parasites are integral parts of ecosystem function and important drivers of evolutionary processes. Characterizing ectoparasite diversity is fundamental to studies of host-parasite interactions, evolution, and conservation, and also for understanding emerging disease threats for some vector borne pathogens. With more than 1400 species, bats represent the second most speciose mammalian clade, but their ectoparasite fauna are poorly known for most species. We sequenced mitochondrial Cytochrome Oxidase C subunit I and nuclear 18S ribosomal gene fragments, and used Bayesian phylogenetic analyses to characterize ectoparasite taxon identity and diversity for 17 species of parasitized bats sampled along the Baja California peninsula and in Northwestern Mexico. The sequence data revealed multiple novel lineages of bat bugs (Cimicidae), flies (Nycteribiidae and Streblidae), and ticks (Argasidae). Within families, the new linages showed more than 10% sequence divergence, which is consistent with separation at least at the species level. Both families of bat flies showed host specificity, particularly on Myotis species. We also identified new records for the Baja peninsula of one tick (Carios kelleyi), and of five Streblid bat fly species. One Nycteribiid bat fly haplotype from Pallid bat (Antrozous pallidus) hosts was found throughout the peninsula, suggesting potential long distance co-dispersal with hosts. Different bat bug and tick communities were found in the north and south of the peninsula. This study is the first systematic survey of bat ectoparasites in the Baja California peninsula, revealing novel lineages that are highly genetically differentiated from other parts of North America. For some ectoparasite species, haplotype distributions may reflect patterns of bat migration. This work is a first step in characterizing ectoparasite diversity over the Baja California peninsula, and understanding how ecological and evolutionary interactions shape bat ectoparasite communities among host species in different parts of their ranges.

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KEYWORDS

18S, Argasidae, Chiroptera, Cimicidae, COI, DNA barcoding, ectoparasites, Nycteribiidae, phylogeny, Streblidae

TAXONOMY CLASSIFICATION

Biodiversity ecology, Ecological genetics, Genetics, Parasitology, Phylogenetics, Taxonomy

1 | INTRODUCTION

Characterizing ectoparasite diversity is fundamental to studies of host-parasite interactions, parasite evolution and conservation, and for understanding emerging disease threats for some vector borne pathogens (Morand et al., 2006; Poulin, 2014; Spencer & Zuk, 2016). Bat ectoparasites are of particular interest as model systems for host-parasite co-evolution and parasite community structure studies (Gómez & Nichols, 2013; Spencer & Zuk, 2016). Bat-ectoparasite relationships are also important for understanding bat dispersal, pathogen transmission, and zoonotic disease risks (Klimpel & Mehlhorn, 2014; Speer et al., 2019; Wilder et al., 2015). Despite the widely recognized need to increase sampling effort for pathogen/parasite discovery in bats, bat ectoparasites are still understudied in most parts of the world (Gay et al., 2014; Reinhardt & Siva-Jothy, 2007). Bat-associated ectoparasites include Insecta, comprising bat bugs (Hemiptera), fleas (Siphonaptera), and flies (Diptera); and Arachnida comprising ticks (Ixodida) and mites (orders Mesostigmata and Trombidiformes) (Seneviratne et al., 2009; Ter Hofstede & Fenton, 2005). All these ectoparasites are hematophagous (blood feeding) organisms, creating potential zoonotic disease transmission risks for humans and domestic animals (Poulin, 2014).

Bat bugs (family Cimicidae), have a worldwide distribution and comprise approximately 110 known species within 24 genera, the majority of which are ecologically and biologically associated with bats (Hornok et al., 2017; Ossa et al., 2019). Figure 1 shows a *Parastrellus hesperus* individual sampled for this study in northwestern Mexico, infested by bat bugs. A second, less studied family, Polyctenidae also includes bat-associated bugs, with tropical and subtropical distributions (Klimpel & Mehlhorn, 2014).

Bat flies comprise two main families, Nycteribiidae and Streblidae, which have a common origin from a single lineage coevolving with bats (Dittmar et al., 2006; Wenzel & Tipton, 1966). As of 2006, there were approximately 520 species described (Dittmar et al., 2006). With new records added regularly (Saldaña-Vázquez et al., 2019; Szentiványi et al., 2019), taxonomy at the species level is often ambiguous, and an updated global review and synthesis is yet to be produced.

Ticks are grouped in three families: Argasidae (soft ticks), Ixodidae (hard ticks), and Nuttalliellidae. The latter family comprises a single known species, *Nuttalliella namaqua* (Burger et al., 2014; Mans et al., 2012), which has been suggested as the basal tick lineage (Mans et al., 2012). There are approximately 894 known species of ticks worldwide, with 32 species of Argasidae and 68 species of Ixodidae in Mexico (Guglielmone et al., 2010; Pérez et al., 2014). Bat tick phylogenetic studies have mainly focused on old world



FIGURE 1 Specimen of canyon bat (*Parastrellus hesperus*) infested by bat bugs, sampled for this study in northwestern Mexico. Image credits: The authors, University of Leeds.

species (Hornok et al., 2017), with a limited number for the Americas and other parts of the world (Black et al., 1997; Burger et al., 2014), and suggest bat-associated ticks commonly exhibit host-specificity (Sándor et al., 2019). Classification of taxonomic relationships among soft ticks are controversial (Burger et al., 2014), and more studies are needed to accurately describe their taxonomic status (Burger et al., 2014; Estrada-Pena et al., 2010).

Previous bat ectoparasite studies in North America have described the distribution and taxonomic status of bat flies, bugs, and ticks (Bradshaw & Ross, 1961; Graciolli et al., 2007; Jobling, 1938; Usinger, 1966), and their medical importance (Dick et al., 2003; Gill et al., 2004; Steinlein et al., 2001), from a limited number of bat host species. In Mexico, publications have focused on describing species diversity of flies, ticks, and mites (Bolaños-García et al., 2018; Guzmán-Cornejo et al., 2017; Pérez et al., 2014); new records of bat flies (Cuxim-Koyoc et al., 2015; Ramírez-Martínez et al., 2016; Trujillo-Pahua & Ibáñez-Bernal, 2018); ecological relationships between bats and bat flies (Saldaña-Vázquez et al., 2019; Salinas-Ramos et al., 2018; Zamora-Mejías et al., 2020); and *Rickettsia* presence in soft ticks (Sánchez-Montes et al., 2016). To our knowledge, there have been no studies of bat bugs in Mexico.

Baja California (Figure 2) is an isolated peninsula, 1300 km long, in northwestern Mexico. Its complex geological history means it has a mosaic of habitats, with high levels of biodiversity and endemism FIGURE 2 Map of sampling locations for bats and bat ectoparasites reported in this study.



TABLE 1 Left section: List of sites sampled showing number, name, abbreviation (site label) and geographic coordinates. Right section: List of bat species sampled and their abbreviated species labels. *Myotis* sp. refers to any unidentified *Myotis* morphologically resembling *M. velifer*, found at Ures site.

Site number	Site name	Site label	Latitude	Longitude	Bat species	Species label
Baja Caliifornia Penir	nsula					
0	San Diego	SanDi	32.927	-117.176	Antrozous pallidus	ANPA
1	Chabacanos	Chaba	32.566	-116.493	Artibeus hirsutus	ARHI
2	Mosqueda	Mosq	32.156	-115.279	Choeronycteris mexicana	CHME
3	Ensenada	Ense	31.770	-116.520	Eptesicus fuscus	EPFU
4	Meling	Meli	30.972	-115.744	Glossophaga soricina	GLSO
5	Jolla	Jolla	30.920	-115.601	Leptonycteris yerbabuenae	LEYE
6	San Fernando	SanFe	29.971	-115.237	Macrotus californicus	MACA
7	Rosarito	Rosa	28.613	-114.047	Mormoops megalophylla	MOME
8	San Ignacio	Sanlg	27.297	-112.898	Myotis californicus	MYCA
9	San Basilio	SanBa	26.371	-111.429	Myotis peninsularis	MYPE
10	Pocitas	Poza	24.403	-111.104	Myotis sp	MYsp
11	La Paz	LaPaz	24.103	-110.306	Myotis velifer	MYVE
12	Testera	Teste	23.764	-110.055	Myotis vivesi	MYVI
13	Faro	Faro	23.427	-110.233	Myotis volans	MYVO
14	Tesos	Teso	23.175	-109.611	Myotis yumanensis	MYYU
Continental					Parastrellus hesperus	PAHE
15	Ures	Ures	29.433	-110.376	Sturnira parvidens	STPA
16	Tucson	Tucs	20.705	-103.336		
17	Primavera	Prima	20.679	-103.602		

(González-Abraham et al., 2010; Nájera-Cortazar et al., 2015; Riddle et al., 2000), with approximately 25 species of bats recorded (Álvarez-Castañeda et al., 2015; Medellín et al., 2007). While there are several studies describing Baja bat diversity, distributions, and ecology (Álvarez-Castañeda et al., 2015; Frick et al., 2008; Nájera-Cortazar et al., 2015), there are no studies describing the bat ectoparasite fauna.

Here, we use a phylogenetic approach to characterize ectoparasite (bat bugs, flies, and ticks) identity and diversity for 17 species of bat resident along the Baja California peninsula, and northwestern Mexico, based on mitochondrial Cytochrome Oxidase C subunit I (COI) and nuclear 18S ribosomal (18S) DNA amplicon sequences. Ectoparasites are assigned as known or novel lineages within Baja and the rest of the Americas, and we evaluate potential distributions with respect to Baja, and their hosts' wider ranges. The results are relevant for increasing knowledge of bat and ectoparasite distributions across western North America, and to provide a basis for understanding how ecological and evolutionary interactions shape parasite community structure along environmental gradients such as those found in northwestern Mexico.

2 | METHODS

2.1 | Sample collection

Bat sampling and collection of ectoparasites were conducted at 14 sites along the Baja California peninsula, and at three sites in western continental Mexico during 2016-2018 (Figure 2). Bats were identified to species level in the field following published identification guides (Álvarez-Castañeda et al., 2015; Medellín et al., 2007), and subsequently, Myotis bats were identified by molecular assays (Najera-Cortazar, 2020). A list of sites and bat species sampled is given in Table 1. Ectoparasites were collected manually from bats using forceps and transferred to labeled Eppendorf tubes with 96% ethanol for storage. Ectoparasites collected from the same bat but from different taxonomic families were stored in separate tubes for each family, with corresponding labels and appropriate specimen source identifiers. All samples were kept on ice during fieldwork, until their arrival to the laboratory where they were stored at -20°C. Ectoparasites were photographed during fieldwork using a portable Maozua 5MP 20-300X USB microscope, taking ventral and dorsal views. Preliminary classifications of ectoparasites were assigned according to morphological characters specified in keys by McDaniel (1979), Usinger (1966), Knee and Proctor (2006) and Dick and Miller (2010), and adapted for North American ectoparasites by this study. In addition, 10 specimens of bat bugs from Parastrellus hesperus hosts, captured at Ramona, California, U.S.A. (Site 0), were donated from the San Diego Natural History Museum, California U.S.A. Since the focus of this study was a molecular barcoding and phylogenetic approach, morphologies are not described in detail, and will be presented elsewhere.

2.2 | DNA extraction, primer selection, and PCR amplification

Individual ectoparasite bodies were crushed in Eppendorf tubes using sterile pestles, followed by digestion with Proteinase K at 56°C for 18h. DNA was extracted using either a Thermo-Fisher DNA extraction and purification kit (Thermo Fisher Scientific Inc.) or QuickExtract kit (Epicenter, Illumina), following the manufacturers' protocols. PCR conditions were optimized separately for each ectoparasite group, and according to each primer set. Primers LCO1490 (5'-GGTCAACAAATCATAAAGATATTGG-3) and HCO2198 (5'-TAAACTTCAGGGTGACCAAAAAATCA-3') (Folmer et al., 1994) were used to amplify an approximately 700 bp fragment of the mitochondrial COI gene in a 25μ l reaction containing: 1 U of Flexi GoTaq Taq Polymerase, 5× GoTaq reaction buffer, 50mM MgCl₂ (1.5-2.5 mM final concentration), 0.5 µl PCR nucleotide Mix (0.2 mM each), 0.5 µl of each set of primers (1 µM final concentration), 15.8 µl ddH₂O, and 8 µl template DNA (extractions diluted at 1:10 with sterile distilled water). Thermal cycling parameters were as follows on a TECHNE thermocycler model TC-512: initial denaturation step at 95°C for 5 min, followed by 40 cycles of denaturation at 94°C for 40s, annealing at 53°C to 56°C for 1 min and extension at 72°C for 1 min. Final extension was performed at 72°C for 10 min.

For the 18S rDNA gene, approximately 800 bp amplicons were amplified using the primers 18S-1F (5' CTGGTTGATCCTGCCAG TAGT 3') and 18S-3R (5' GGTTAGAACTAGGGCGGTATCT 3') for bat bugs (Campbell et al., 1995); a0.7-F (5' ATTAAAGTTGTTGC GGTT 3') and 7R (5' GCATCACAGACCTGTTATTGC 3') for flies (Whiting, 2002); and D-F (5' GGCCCCGTAATTGGAATGAGTA 3') and C-R (5' CTGAGATCCAACTACGAGCTT 3') for ticks (Mangold et al., 1997). The same reaction mix quantities were used as for the COI gene, with MgCl₂ at a final concentration of 1.5 mM for all primer sets. Thermal cycling parameters were: initial denaturation at 95°C for 5 min, followed by 30 cycles of denaturation at 94°C for 40s, annealing at 53°C for 1 min and extension at 72°C for 1 min. Final extension was performed at 72°C for 10 min.

PCR products were visualized by gel electrophoresis using 1% agarose with GelRed® (Biotium Ltd) staining, and then sent to Genewiz Inc. (Azenta Life Sciences), for PCR product purification and Sanger sequencing, with each amplicon sequenced in both forward and reverse directions.

2.3 | Quality control and reference sequences

Sequence quality for both the forward and reverse strands of each amplicon was evaluated in BioEdit 7.2.5 (Hall, 2005). Trimmed forward and reverse sequences were combined to generate a consensus sequence for each amplicon, and then analyzed in BLAST (The National Library of Medicine, 2018) to generate initial taxon identities and identify reference sequences. Additionally, COI sequences were also analyzed in the BOLD system platform (Ratnasingham & Hebert, 2007). Reference sequences for phylogenetic analyses were compiled from

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previous studies on each ectoparasite group (Burger et al., 2014; Dittmar et al., 2006; Mans et al., 2012; Tortosa et al., 2013), and by performing a systematic search using *AnnotationBustR* 1.2 package (Borstein, 2018) in RStudio 1.1.456 (RStudio Team, 2021), searching for the closest genus for the sequences generated in this study and for those obtained by BLAST. Alignments combining reference sequences and those from this study were generated using CLUSTAL W (Thompson et al., 1994), implemented in BioEdit, and were reviewed by eye, with manual correction of potential errors as required.

2.4 | Sequence summary statistics and phylogenetic analyses

Genetic distance estimates among haplotypes and best fit sequence evolution models for the COI and 18S datasets were evaluated using MEGA 10.1.7 (Sudhir et al., 2018). The Barcode Index Number system was followed to delimit a lineage, where intraspecific variation at COI is generally considered as groups of sequences with less than 2% divergence, and exhibiting more than 4% divergence from neighboring lineages (Hebert et al., 2003; Ratnasingham & Hebert, 2013; Salinas-Ramos et al., 2018).

Bayesian phylogenetic analyses were performed using the best fit evolution model identified for each group and each marker implemented with BEAST 1.10.4 (Suchard et al., 2018). A MCMC chain length of 10,000,000 was used and priors specific to each parasite group and sequence evolution model were selected using the program BEAUti 1.10.4 (Suchard et al., 2018). Phylogenetic analyses were performed separately for each ectoparasite group and each marker. Nycteribiidae and Streblidae families of bat flies were analyzed separately in order to improve sequence alignment quality and phylogenetic resolution. For each family-marker combination, two separate runs using the same Bayesian settings file generated by BEAUti were run in BEAST, with a burn in of the 10% of the total number of iterations. After, stationarity of BEAST results were assessed in Tracer 1.7.1 (Rambaut et al., 2018). Both files for each ectoparasite family-marker combination were combined using LogCombiner 1.10.4 (Suchard et al., 2018), generating a single .log file and a single .tree file. A majority-rule consensus tree was inferred using TreeAnnotator 1.10.4 (Suchard et al., 2018) with the combined .tree file, and using a posterior probability limit of 0.6, median nodes heights were summarized. Phylogenetic trees were annotated, edited, and visualized using iTOL 5.6.2 (Letunic & Bork, 2016).

Following phylogenetic analysis, haplotypes for each family were grouped by lineage, and haplotype diversity and genetic summary statistics were calculated in DNAsp 5.10.01 (Librado & Rozas, 2009). Outgroups were chosen based on previous phylogenetic studies of each group, using *Bucimex chilensis*, *Primicimex cavernis*, and *Anthocoris flavipes* for bat bugs (Ossa et al., 2019); Drosophila melanogaster, Chrysops niger, Musca domestica, and Sarcophaga bullata (Dittmar et al., 2006), for both bat fly families, plus Ornithomya avicularia for the family Streblidae only; and Nuttalliella namaqua (Mans et al., 2012) for ticks.

2.5 | Ethics approval

All bat capture, handling and sampling was carried out with the approval of the ethics committee of the Faculty of Biological Sciences, University of Leeds (AWCNRW170615); and following the Guidelines of the American Society of Mammalogists (Sikes et al., 2016). Sampling was carried out under the permits SEMARNAT-DGVS-008972-16 and SEMARNAT-DGVS-001642-18 issued by Secretaría del Medio Ambiente y Recursos Naturales (SEMARNAT) in Mexico. The latter included two Myotis bat species listed in the Mexican Official Norm for the protection of native species of flora and fauna in Mexico [NOM-059-SEMARNAT-2010, (Secretaría de Medio Ambiente y Recursos Naturales, 2010)], under the Pr (under special protection) and P (in danger of extinction) categories (M. evotis and M. vivesi, respectively); and sampling on protected reserves. When required, permission was also solicited and granted from private land owners. All samples were imported into the UK under permission of the Department for Environment, Food and Rural Affairs (DEFRA) from the Animal and Plant Health Agency (permit ITIMP19.0036).

3 | RESULTS

3.1 | Sampling and field identifications

A total of 1988 ectoparasites were collected within the orders Diptera (flies), Hemiptera (bugs), Ixodida (ticks), Mesostigmata (mites), Siphonaptera (fleas), and Trombidiformes (chiggers). Fleas, chiggers, mites, and any unclassified specimens (e.g., where it was not possible to differentiate small tick larvae and mites) were excluded for this study, to be evaluated separately. The remaining samples comprised 90 bat bugs, 213 bat flies, and 126 ticks, collected from 138 individual bat hosts of 17 species (Figure 3).

Flies were the predominant ectoparasite type, found in 13 of 17 sites, followed by ticks and bugs, present in eight sites each. Initial field morphological evaluations suggested most bug specimens were related to *Cimex pilosellus* (Usinger, 1966). Most bat flies could be identified to the genus level, which were later corroborated with molecular data. On the basis of morphology, all bat ticks were identified as family Argasidae (soft ticks), with at least six different morphotypes present. Most of these were tentatively attributed to the genus *Ornithodoros*.

3.2 | BLAST and BOLD sequence matches

Amplicon sequences were generated from 145 specimens for mitochondrial COI and from 147 specimens for 18S rDNA across the three groups: bugs n = 30/30 (COI/18S, respectively); flies n = 76/73; and ticks n = 39/44. When there was more than one ectoparasite specimen available per ectoparasite family per bat, one specimen for sequencing was selected based on morphological similarity, sequencing one individual of each morphotype of each family. No haplotypes from



FIGURE 3 Number of bat bugs, flies, and ticks captured per site in 2016–2018 fieldwork.

NAJERA-CORTAZAR ET AL.

bugs and most tick specimens matched existing sequences deposited in GenBank or BOLD at the species level (divergence >4%), with the exception of one of the tick lineages showing similarity to *Carios kelleyi* (96.95% in GenBank, 97.07% in BOLD). Similarly for nycteribild bat flies, there were no species level matches for neither COI nor 18S markers. In the case of streblid bat flies, 14 sequences matched (~99%-100% in GenBank-BOLD databases) with five known species for COI (see Table 2); but there were no matches for 18S. These marker differences are likely attributable to database coverage, since where reference sequences were available for COI and 18S from the same specimen/ study, BLAST results for our sequences were consistent for both markers. For brevity and to provide comparability with larger numbers of reference sequences further reporting of diversity, divergence statistics, and taxonomic identity will be focused on COI results.

3.3 | Genetic diversity

Genetic diversity statistics for each COI lineage in each ectoparasite family are summarized in Table 2. Excluding four lineages represented by single individuals (e.g., *Cimex* 1), and three lineages including only individuals with the same haplotype (e.g., *Basilia* 1), nucleotide diversity (N_d) ranged from 0.001 to 0.006, with the highest value 0.071, presented by the Tick 4 lineage, with three haplotypes (*H*) in three sequences (Table 2). In general, the number of haplotypes were close to or the same as the number of sequences tested per lineage.

3.4 | Phylogenetic assessment of Baja peninsula bat bug sequences

The best fit evolution model for the COI gene set was GTR+G+I, and K2+G+I for 18S. For each marker, 30 sequences were generated, forming four novel lineages with respect to reference sequences (Figure 4). Genetic divergence between the four peninsular lineages ranged from 9.9% to 17.1% (Appendix 1), and between 7.2% and 20.9% against reference sequences, with *C. latipennis*, *C. antennatus*, and *C. adjuntus* presenting the lowest divergence values. Phylogenetically, all the lineages sit within the *Pilosellus* complex, of North American members of the *Cimex* genus.

For COI, Cimex 1 is represented by a single specimen (EBCO155) obtained from a Myotis yumanensis host at Mosqueda, and forms a sister lineage to Cimex 2, represented by two haplotypes from two bugs parasitizing M. californicus hosts, which were also sampled from northern sites. The 18S analysis also placed EB18S155 in a separate clade, but Cimex 2 is paraphyletic. Both markers placed Cimex latipennis, which is distributed from Canada to the north western USA (Usinger, 1966), as the closest named molecular reference species to these 2 lineages. Cimex 3 is represented by a monophyletic clade for COI, consisting of two haplotypes, sampled from specimens collected from four Antrozous pallidus and one M. californicus hosts, all distributed in the southern half of the peninsula (sites Rosarito, San Ignacio and Tesos). For 18S, a C. cf antennatus reference sequence nests within the clade with greater than 60% of posterior probability support. For COI, reference sequences for C. pilosellus and C. brevis formed a clade which appeared to be ancestral to Cimex 3, with 13% divergence (Appendix 1). There is no molecular reference for C. brevis in the 18S analysis, and Cimex 4 is the closest sister lineage.

The *Cimex* 4 COI lineage includes 17 haplotypes derived from 22 specimens, where 21 of the bugs were found parasitizing *Parastrellus hesperus* individuals, and one from *A. pallidus* (EBCO201/EB18S201), all from northern sites (Figure 2), with *C. antennatus* as the closet sequence match (9.3%; Figure 4), followed by *C. adjunctus*. Sequences of *C. adjunctus* in the 18S topology clustered in a clade with other *Cimex* species with mixed origins, including some non-Palearctic species. This may represent

-C(ORTAZAI	R et A	.L.											Ecolo	ogy and	Evo	olution	Open Ac	-W	/11	LEY	7
	Known congeneric ectoparasite species for host(s) when no molecular match	C. incrassatus, C. pilosellus ¹	Cimex adjunctus, C. antennatus, C. brevis, C. pilosellus, C. latippenis ¹	C. incrassatus, Paracimex cavernis ¹	C. antennatus, C. incrassatus, C. pilosellus ^{1.8}	Basilia antrozoi, B. corynborhini, B. forcipata, B. rondani ^{6,8}	Basilia antrozoi, B. corynborhini, B. forcipata, B. jellisoni, B. rondani ^{5,6,8}	Basilia forcipata, B. jellisoni, B. rondani ^{5, 6}	Basilia plaumanni, B. pynzonix, B. producto ⁶	Basilia antrozoi, B. rondani ⁶		T	Aspidoptera phyllostomatis, Paratrichobius Iongicrus ⁸		Aspidoptera phyllostomatis, Paratrichobius Iongicrus ³	Trichobius adamsi, Nycterophilia coxata ⁸	Trichobius sphaeronotus, T. galei, T. leionotus, Nycterophilia mormoopsis ⁷	Trichobius mixtus, Paratrichobius longicrus ⁸	I	I	1	(Continues)
	Min. % div.	5.1	3.2	5.9	4.4	9.4	9.4	5.0	9.5	8.6	2.3	0.8	3.0	0.0	4.3	10.3	6.0	5.6	3.1	0.2	0.0	
	Closest reference species	Cimex latipennis	Cimex latippenis	Cimex adjuntus	Cimex antennatus	Nycteribia pygmaea	Nycteribia pygmaea	Basilia boardmani	Basilia ortizi	Basilia ortizi.	Aspidoptera phyllostomatis	Megistopoda aranea	Megistopoda aranea	Nycterophilia coxata	Paratrichobius longicrus	Trichobius sparsus	Trichobius sphaeronotus	Trichobius dugesii	Trichobius intermedius	Trichobius dugesii	Trichobius sphaeronotus	
III tills study	Host(s)	Antrozous pallidus	Myotis californicus	Antrozous pallidus, Myotis californicus	Parastrellus hesperus, Antrozous pallidus	Myotis sp Myotis californicus Parastrellus hesperus	Myotis volans Antrozous pallidus Myotis californicus Myotis yumanensis	Myotis yumanensis	Myotis vivesi	Antrozous pallidus	Sturnira parvidens	Sturnira parvidens	Artibeus hirsutus	Leptonycteris yerbabuenae Macrotus californicus	Artibeus hirsutus	Macrotus californicus	Mormoops megalophylla	Choeronycteris mexicana	Artibeus hirsutus	Glossophaga soricina	Leptonycteris yerbabuenae	
א ומבוורוו ובמ	P	0	0.006	0.001	0.005	0.003	0.007	0	0.0009	0.001	0	0	0.0015	0.006	0	0.011	0.008	0	0	0	0	
	۳	0	4	0.4	0.97	4	0.89	0	0.67	0.58	0	0	4	0.75	0	0.83	4	0	0	0	0	
iyai asi	п	1	7	7	17	2	œ	1	2	5	-	1	7	4	4	ო	7	1	4	1	4	
וום ברור	S	0	4	7	14	9	22	0	1	4	0	0	4	6	0	15	9	0	0	0	0	
	z	1	7	2	22	2	12	ო	ო	26	2	1	2	6	4	4	7	2	4	1	7	
lidi summary statisti	snus	Cimex 1	Cimex 2	Cimex 3	Cimex 4	Nycteribiid 1	Nycteribiid 2	Basilia 1	Basilia 2a	Basilia 2b	Aspidoptera phyllostomatis	Megistopoda aranea	Megistopoda sp	Nycterophilia coxata	Paratrichobius 1	Streblid 1	Trichobius 1	Trichobius 2	Trichobius intermedius	Trichobius dugesii	Trichobius sphaeronotus	
ADEE 2 MORCH	Lineage/putative ge	Bat bugs: Cimicidae				Bat flies: Nycteribiidae					Bat flies: Streblidae											

res identified in this study Molecular summary statistics for the ectoparasites linea TABLE 2

Lineage/putative	genus	z	S	н	ΡH	م ع	Host(s)	Closest reference species	Min. % div.	Known congeneric ectoparasite species for host(s) when no molecular match
Bat Ticks:	Antricola 1	1	0	1	0	0	Macrotus californicus Myotis vivesi	Antricola marginatus	10.7	Ornithodoros sp. ⁸
Argasidae	Carios kelleyi	10	33	6	0.98	0.011	Antrozous pallidus	Carios kelleyi	1.9	1
	Tick 1	с	7	с	1	0.002	Antrozous pallidus Myotis peninsularis	Ornithodoros turicata	12.3	Ornithodoros sp., O. kelleyi, O. rossi ^{8,9}
	Tick 2	7	4	7	1	0.001	Antrozous pallidus Myotis yumanensis	Ornithodoro faccinii	13.2	Ornithodoros kelleyi, O. yumatensis, O. rossi ^{5.8.9}
	Tick 3	9	15	4	0.8	0.008	Myotis yumanensis	Carios vespertilionis	13.1	Ornithodoros kelleyi, O. yumatensis ^{5,8}
	Tick 4	ო	72	ო	7	0.071	Antrozous pallidus Eptesicus fuscus	Ornithodoro faccinii	11.6	Ornithodoros sp., O. kelleyi, O. rossi ^{8,9}
	Tick 5	ო	16	œ	0.93	0.005	Myotis yumanensis Antrozous pallidus Parastrellus hesperus Eptesicus fuscus	Antricola marginatus	11.8	Ornithodoros sp., O. kelleyi, O. rossi, O. yumatensis ^{8,9}
	Tick 6	ო	1	2	0.67	0.001	Myotis peninsularis	Ornithodoros yumatensis	9.9	No previous record to our knowledge
<i>Note:</i> N, number of divergence from clc	sequences within each sest reference sequer	h lineag(nce. 1. U	e; S, nur. Isinger (nber of : (1966), 2	segregating Cuxim-Ko	sites; H, nu voc et al. (2	mber of haplotypes; H _d , haplotyp 015), 3. Trujillo-Pahua and Ibáñez	e diversity; N _d , nucleotide -Bernal (2018), 4. Zamora-	diversity, a Meiías et a	nd Min. % Div., minimum percentage sequence 1. (2020), 5. Braun et al. (2015), 6. Graciolli

Bradshaw and Ross (1961), 9. Steinlein et al. (2001).

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(2007), 7. Ramírez-Martínez et al.

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3.5 | Phylogenetic assessment of bat fly sequences

The best fit sequence evolution models were GTR+G for the COI gene set, and T92+G+I for the 18S data. In total, 77 bat flies were sequenced, yielding 76 sequences for COI and 73 for 18S amplicons respectively. Individual sequences for two specimens for COI, and three for 18S did not pass sequence quality thresholds and were discarded. In the final sequence set for the COI marker, there were 49 sequences from the Nycteribiidae family (wingless bat flies) and 27 for the Streblidae family (winged bat flies), representing 10 novel lineages and six new species records for Baja. For the 18S marker, 46 sequences were generated for the Nycteribiidae family, and 26 sequences for the Streblidae family. For this marker, nine novel lineages were observed (18S sequences corresponding to the Basilia 2a COI lineage could not be amplified), along with six new species records for Baja.

In the COI phylogenetic analysis, the 49 Nycteribiidae sequences formed five lineages, all of which appeared to be novel with respect to GenBank references. Genetic divergence among Baja lineages ranged from 2.9% to 14.5%, and up to 16% against reference sequences (Appendix 2). For COI, Nycteribiid 1 and 2, formed sister clades with 4.4% divergence, and 10% divergence from the closest reference haplotypes derived from Asian Nycteribia, species. For 18S, the closest references to Nycteribiid 1 and 2 were North American species not available for COI, *Basilia corynorhini* and *Basilia forcipata*, respectively. Nycteribiid 1 and 2 lineages were primarily associated with *Myotis* bat hosts, but with one Nycteribiid 2 haplotype recovered from a *Parastrellus hesperus* host in the 18S dataset (specimen EN18S514, Figure 5).

The three other Baja lineages formed clades associated with *Basilia* reference sequences from species recorded in Madagascar, USA, and Panama. *Basilia* 1 has 4.9% divergence from *Basilia boardmani*, a bat fly distributed throughout the United States parasitizing *Myotis* bats (Graciolli et al., 2007). Specimens with *Basilia* 1 haplotypes were sampled at mid-peninsula, parasitizing *Myotis yumanensis*, which is widely distributed throughout western North America.

Genetic divergence among lineages *Basilia* 2a and 2b was 2.9% (Table 2), representing the threshold for intra/inter interspecific values (Hebert et al., 2003; Ratnasingham & Hebert, 2013). We assign them as distinct lineages due to their different host species, where *Basilia* 2a parasitized *M. vivesi*, and *Basilia* 2b appeared to be restricted to *Antrozous pallidus* (Figure 5). The spatial distribution of *Basilia* 2b haplotypes across the peninsula, suggests potential long distance co-dispersal with their hosts *A. pallidus* (Figure 5), providing evidence supporting bat movement along Baja. *Basilia* 2b may

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FIGURE 4 Bayesian phylogenetic trees for bat bugs of the family Cimicidae, obtained using mitochondrial COI (left) and the ribosomal 18S (right) sequences. Posterior probability support is indicated by the size of black circles at tree nodes. Where available, information for location and host is written next to each reference sequence label. To improve clarity of the tree, collapsed clades of reference sequences are shown as gray triangles. GenBank accession numbers for reference sequences are given in the sequence labels.

represent *B. antrozoi*, which has been previously reported to parasitize *A. pallidus* (Graciolli et al., 2007).

The 11 Streblidae lineages had genetic divergence for COI of 0.01%–18.9% for sequences sampled within this study, and up to 21.6% against their reference sequences (Appendix 3). For the five novel lineages from this family four were named according to the closest genus in the phylogenetic analysis (*Trichobius* 1, *Trichobius* 2, *Paratrichobius* 1, and *Megistopoda* sp., Figure 6), with genetic divergence values ranging from 3% to 8.4% from each of the closest corresponding clades. The Streblid 1 lineage presented 10.8% divergence against its closest reference, *T. sphaeronotus* (Appendix 3, Figure 6), and is therefore not attributed to an existing genus.

The other streblid clades matched sequences from six known species: Aspidoptera phyllostomatis, Megistopoda aranea, Nycterophilia coxata, Trichobius dugesii, T. intermedius, and T. sphaeronotus (Figure 6), representing new records for these species in Baja and western Mexico. For the specimens classified as N. coxata, T. dugesii, T. intermedius, and T. sphaeronotus (Figure 6, Appendix 3), there was less than 1.5% divergence against the GenBank references. The fly lineages A. phyllostomatis matched with two reference sequences classified as A. delatorrei (Figure 6). However, A. phyllostomatis and A. delatorrei reference sequences only differed by 2.1%-2.3%, which falls at the threshold for species level differentiation. Without further information on the source A. delatorrei specimens, we decided to retain the A. phyllostomatis classification for our specimens. In the case of the M. aranea and Megistopoda sp lineages, there were two separate clades of M. aranea reference sequences present. The Megistopoda sp specimens (ESCO606 and ESCO610, dark blue clade, Figure 6) were grouped closer to M. aranea EF531219 reference sequence, with 2.9% and 3.1% divergence, suggesting a different species from the M. aranea EF531219 reference, as well as from the other M. aranea clade (bright blue clade, Figure 6), with 4.4% genetic divergence. Supporting this, it is noted that each lineage had different host species, Sturnira parvidens and Artibeus hirsutus, respectively (Table 2). Most streblid lineages were parasitizing fruit-nectar feeding bats (Phyllostomidae) over the mid and northern peninsula, with the exception of Trichobius 1 found on Mormoops megalophylla hosts, (family Mormoopidae), an insectivore. Trichobius 1 was the only streblid fly lineage found in the south of Baja. The other fly lineages were distributed in the southern continental sites (Tucson

and Primavera, Figure 2, Table 1). Outside of the closest sequence matches, none of the other species reported as parasitizing hosts in Baja (Table 2) appear to have reference sequences in GenBank. Genetic divergence within groups from the lineages in this study showed levels between 0.0% and 1.1%, with the exception of *A. phyllostomatis*, which showed a 3.3% divergence within its group (Appendix 3).

3.6 | Phylogenetic assessment of Baja peninsula bat tick sequences

The best fit sequence evolution models were GTR+G+I for the COI gene set, and K2+G+I for the 18S gene set. There were 45 ticks

sequenced from the Argasidae family (soft ticks), with 39 sequences generated for COI, and 44 for 18S, where six specimens failed to amplify for COI (specimens ETCO157, ETCO306, ETCO338, ETCO480b, ETCO480c, and ETCO485), and one specimen failed to amplify for 18S (specimen ET18S457). One new Baja species record and seven potential novel lineages were obtained (Figure 7). The only match with GenBank and BOLD COI records (95.95% and 97.07%, respectively) was the soft tick *Carios kelleyi*, for 11 specimens (parasitizing A. *pallidus* hosts) with intra-clade divergence of 1.94%-2.47% (Figure 7, Appendix 4). For COI data, the Antricola 1 lineage had 10.6% divergence from Antricola marginatus, and 14.6% from A. mexicanus (Appendix 4). Ornithodoros faccini had a divergence of around 11% for clades Carios kelleyi, Tick 1 to 5, and O. yumatensis showed divergence of around 9.9% with respect to Tick



FIGURE 5 Bayesian phylogenetic trees for bat flies of the family Nycteribiidae obtained using mitochondrial COI (left) and the ribosomal 18S (right) sequences. Posterior probability support is indicated by the size of black circles at tree nodes. Where available, information on location and host is written next to each reference sequence label. To improve clarity of the tree, collapsed clades of reference sequences are shown as gray triangles. GenBank accession numbers for reference sequences are given in the sequence labels.

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6 lineage. Ticks of lineage Tick 6 were recovered from *M. peninsularis* from which there are no previous records of bat ticks to our knowledge.

All sequences from this study generated the same lineages for both COI and 18S genes (Figure 7). Lineages *C. kelleyi* and Tick 1 to 5 formed a monophyletic group with respect to the reference sequences with posterior support greater than 0.6 for both markers, while *Antricola* 1 and Tick 6 lineages were positioned in separate clades. The topology of sister clade relationships varied slightly between markers, particularly around deeper nodes which had posterior probability support less than 0.85. This suggests more data is required to resolve deeper taxonomic relationships among species. *Antricola* 1 and Tick 6 were separated around shallow deep nodes in both analyses, where the absence or presence of reference sequences influenced their topological proximity (Figure 7). None of the reference sequences from *Antricola*, *Carios*, and *Ornithodoros* genera, form monophyletic groupings with respect to genus nomenclature.

Tick lineages 2 and 6 (gray and orange, Figure 7) were only observed in the south of the peninsula, with hosts A. pallidus and M. peninsularis, and M. peninsularis, respectively. These two species of bats share at least one confirmed roosting site within the study region, at Tesos (Figure 2), suggesting a potential interchange of host species for Tick lineage 2. Evidence of singular host specificity was observed for the lineages C. kelleyi (A. pallidus), Tick 3 (M. yumanensis), and Tick 6 (M. peninsularis), all observed exclusively on the same hosts across sites and field seasons. The Antricola 1 lineage (Figure 7, lilac clade, ETCO301 MYVI) was recovered from one specimen on a M. vivesi host using the COI marker, and grouped with Antricola marginatus (found in the South East of Mexico), followed by A. mexicanus. In the 18S topology, the 18S sequence from this sample formed a clade with five additional sequences from M. vivesi and Macrotus californicus hosts, grouped with A. mexicanus, as no A. marginatus reference was available. The two host species for ticks of this lineage were, sampled mostly at mid-peninsula (Figures 2 and 3). Tick 5 lineage had the most diverse host and spatial distribution, being found on four bat species, and at sites from the north to the south of the peninsula (yellow clade, Figure 7).

4 | DISCUSSION

This study represents the first molecular characterization and phylogenetic analysis of bat bug, bat fly, and bat tick diversity along the Baja California peninsula and northwestern Mexico, a region where the bat ectoparasite fauna is largely undescribed. From a total of 292 ectoparasite specimens, from 17 bat species, evaluated for COI and 18S markers, 21 novel genetic lineages, plus seven new species records for the region, were found. Some of the novel lineages may derive from previously recorded species with no reference sequences available, while others are likely to represent new species. Overall, the work demonstrates that the northwestern region of Mexico hosts a high diversity of previously unknown bat ectoparasites.

4.1 | Bat bugs

Four novel lineages of *Cimex* bugs were identified. The lineages showed evidence of host preference, being found primarily on *M. yumanensis*, *M. californicus*, *A. pallidus*, and *P. hesperus*, respectively, for lineages 1–4. A threshold of 4% sequence divergence is typically taken as representing species level differentiation for COI in arthropods (Hebert et al., 2003), indicating *Cimex* 1, 3, and 4, could be classed as novel species under this criterion. Genetic divergence of *Cimex* 2 compared with *C. latippenis* was 3.2%, but *C. latippenis* has not previously been recorded parasitizing *M. yumanensis* (Braun et al., 2015), which might also suggest *Cimex* 2 as a potential new cryptic species.

Cimicid bugs have low inherent dispersal capacity, generally feeding for a few days, before dropping from the bat host to digest the blood in the roost, where they can survive without feeding for approximately 1.5 years (Ossa et al., 2019). The population structure of bat bugs is mainly influenced by bat movements (Ossa et al., 2019; Talbot et al., 2016, 2017; Usinger, 1966). While the current data indicate the new Cimex lineages may have distinct regional distributions within Baja, their host species' ranges extend further through western North America, suggesting these bugs may also have wider distributions, or come into contact with hosts which migrate over long distances at roost sites. For example, M. yumanensis individuals captured in the northern peninsula in our study had mitochondrial haplotypes matching reference sequences from bats sampled in Alaska (Najera-Cortazar, 2020). This raises the possibility of longrange mixing of microbial pathogen communities in bat bugs along the west coast of North America.

4.2 | Bat flies

Ten novel genetic lineages and six new records of bat flies for the study region were found. Nycteribiid flies were more abundant in the northern temperate sites, while streblids were more abundant in the southern and subtropical sites, supporting trends noted by Dittmar et al. (2006). To our knowledge, molecular records of *Basilia* sp. species have not previously been reported from bats with ranges in the peninsula. Streblid flies were present on bats from the Phyllostomidae, which in general are fruit and nectar feeders, with the exception of the insectivorous *Macrotus californicus*; as well as *Mormoops megallophyla*, from the family Mormoopidae. Nycteribiids parasitized only Vespertilionid bats, which include insectivores and omnivores.

In previous studies it was found that host-specificity can vary according to the diversity and geographic distribution of hosts (de Vasconcelos et al., 2016; Graciolli et al., 2007; Saldaña-Vázquez et al., 2019). We found that most of our Nycteribiid lineages exhibited host specificity, despite having hosts with overlapping ranges and which may share roost sites (e.g., *Myotis* species in northern Baja). The Nycteribiid 2 lineage was found on multiple *Myotis* species, which could potentially indicate specificity at a



FIGURE 6 Bayesian phylogenetic trees for bat flies of the family Streblidae obtained using mitochondrial COI (left) and the ribosomal 18S (right) sequences. Posterior probability support is indicated by the size of black circles at tree nodes. Where available, information on location and host is written next to each reference sequence label. To improve clarity of the tree, collapsed clades of reference sequences are shown as gray triangles. Genbank accession numbers for reference sequences are given in the sequence labels.

genus level. Streblid winged flies have previously been described as mostly non host-specific (Dittmar et al., 2006), but more recent studies suggest most species to be host specific (de Vasconcelos et al., 2016; Estrada-Villegas et al., 2018), a change attributable to methodological improvements in sample collection and taxonomic assignments of flies and hosts. In our data *N. coxata* was found parasitizing *Macrotus californicus* and *Leptonycteris yerbabuenae*, both Phyllostomidae species, which are known to share roost site in Baja (Álvarez-Castañeda et al., 2015), implying potential horizontal transmission.

Lineages Basilia 2a and Basilia 2b (COI divergence 2.9%) are restricted to hosts Myotis vivesi and Antrozous pallidus, respectively. M. vivesi is endemic to the Gulf of Cortes, and restricted to coastal habitats because of its piscivorous diet (Blood & Clark, 1998; Herrera-Montalvo et al., 2017). A. pallidus is sympatric with M. vivesi on the Gulf coast but has a wider distribution across western North America. It is primarily an insectivorous feeder, but also includes scorpions and nectar in its diet (Frick et al., 2009). Basilia fly records previously reported for M. vivesi, but without reference sequences, include B. plaumanni, B. pynzonix, and B. producto (Graciolli et al., 2007), while flies parasitizing A. pallidus have been described as B. antrozoi (Table 2). The threshold level of divergence between the *Basilia* 2a and *Basilia* 2b lineages may indicate recent divergence from a common ancestor, and potential incipient speciation driven by association with sympatric but ecologically differentiated hosts.

4.3 | Bat ticks

A new record of *Carios kelleyi* and seven novel tick lineages belonging to the Argasidae family were found. For both COI and 18S tick lineages 1-5 and *Carios kelleyi* formed a clade with more than 0.6 posterior support, suggesting they form a Baja or western North America endemic lineage of bat ticks derived from a common ancestor.

The genera Antricola, Carios, Nothoaspis, and Ornithodoros are associated with bats and their roosting sites in Mexico (Sánchez-Montes et al., 2016). Compared with Ixodidae, the family Argasidae has few molecular studies and reference sequences (Porter & Hajibabaei, 2018). Classifications at genus level are controversial, with many genera being paraphyletic in existing phylogenies, and debates over synonymous use of genus names such as *Carios* and *Ornithodros* (Burger et al., 2014). Such



FIGURE 7 Bayesian phylogenetic trees for bat ticks of the family Argasidae obtained using mitochondrial COI (left) and the ribosomal 18S (right) sequences. Posterior probability support is indicated by the size of black circles at tree nodes. Where available, information on location and host is written next to each reference sequence label. To improve clarity of the tree, collapsed clades of reference sequences are shown as gray triangles. GenBank accession numbers for reference sequences are given in the sequence labels.

Argas perisicus L76353 Argas peringueyi KY457465 Argas strictus KY457493 rgas africolumbae K.1133632 Argas brumpti KR907227 Argas brumpti KY457462 Amblyomma amer Dermacentor

m AF29187 soni L7634 Ixodes

ambiguities are also reflected in our phylogenies. Furthermore, while many key internal nodes had posterior support greater than 0.6, differences in reference sequence availability made it challenging to interpret the consistency of inter-clade placement between makers.

Argasid ticks have previously been reported for the bat species in this study, primarily *Ornithodoros* species (Table 2), but with the exception of *Carios* (*Ornithodros*) *kelleyi*, none are close molecular matches for our sequences. In ticks, for COI, the threshold for between genus divergence is considered to be above 10% (Hebert et al., 2003). The observed COI divergence among lineages of this study (6.1% up to 19.3%), and to reference sequences (9.9%–22.8%), suggests that the Baja lineages could represent novel species and potentially novel genera.

Nuttalliella namagua KR907240

We found apparent host-specific and generalist lineages for the ticks reported here. Lineages that appeared host-specific were restricted to single sites, while generalists were found across multiple sampling locations. For example, Tick 3 parasitizing *M. yumanensis* WILEY_Ecology and Evolution _

NAJERA-CORTAZAR ET AL.

was found only in San Basilio, and Tick 6 was found only on *M. peninsularis* sampled in Teste. *Carios kelleyi* was found to only parasitise *A. pallidus* in this study, and was found at three sites in the mid- and north peninsula. However, *C. kelleyi* is known to parasitise multiple species across North America and Cuba (Gill et al., 2004), and the reference COI sequence used here derives from an *Eptesicus fuscus* bat sampled in New Jersey, eastern USA (GenBank accession code: MT780277). Argasid ticks show a continuum of hosts-specificity (Cumming, 1998; Esser et al., 2016), but tick stage-cycle must be considered, with immature ticks being more generalist than adult conspecifics (Esser et al., 2016; Nava & Guglielmone, 2013). In this study, life stage was not assessed while collecting ticks; therefore, it is possible that there are gaps in host range regarding unidentified larvae that were not sequenced.

The Tick 5 lineage was found on multiple species in the mid- and north peninsula, but the only specimen recorded for the south peninsula was collected from *Parastrellus hesperus* (specimen 338, Faro site, 18S only). Since *P. hesperus* is rare in southern *Baja*, the presence of this tick indicates potential dispersal of *P. hesperus* from the north to the south peninsula.

4.4 | Limits on ectoparasite sampling and identification

The present study identified 21 novel genetic lineages, plus 7 new ectoparasite species records, from 138 bats of 17 species, sampled across 18 sites and 2 years. This suggests a diverse ectoparasite fauna in this previously unsurveyed region of Mexico, but it is also likely to be an underestimate of the true diversity, due to constraints around sampling effort. Bat sampling was limited to May-August and conducted at relatively accessible locations with water sources to facilitate bat capture. While our sampling sites were chosen to be representative of habitat types across Baja, increasing the spatial and temporal scope of sampling would be likely to increase the number of ectoparasite species discovered. Assessment of ectoparasite fauna found in roosting sites against those found feeding directly from their hosts and expanding seasonal coverage will be important for future work.

Although previous studies report limited data on bat ectoparasites from North Western Mexico and South-Western USA (Bradshaw & Ross, 1961; Braun et al., 2015; Pérez et al., 2014; Usinger, 1966), they do not integrate morphological and molecular information. For many species, no reference sequence is available from voucher specimens, or there are errors in species identifications and incorrect annotation of reference sequences. Therefore, for all the ectoparasite groups in this study, further work is needed to unify molecular and morphological characterization, to fully confirm which lineages represent previously undescribed species, and which are species with morphological descriptions but no previous molecular record. This is particularly important for bat tick lineages where input from expert morphologists is needed to account for different life stages.

5 | CONCLUSIONS

This work presents an initial description of bat ectoparasite diversity relevant to western North America, providing resources useful for future ectoparasite surveys and studies of hostectoparasite ecology, and evaluation of zoonotic disease risks. Future work should focus on expanding spatial and bat host species coverage, integrating morphological and molecular characterization of ectoparasite species, profiling ectoparasite microbiomes and viromes, and understanding the ecological and environmental factors that influence host-parasite community structure and evolution. Bat populations and habitat in Baja California are vulnerable to anthropogenic pressures, and such knowledge will be vital for informing assessments of population status and extinction risks of both hosts and parasites.

AUTHOR CONTRIBUTIONS

Laura A. Najera-Cortazar: Conceptualization (equal); data curation (lead); formal analysis (lead); funding acquisition (equal); investigation (lead); methodology (lead); project administration (lead); resources (equal); software (lead); supervision (supporting); validation (equal); visualization (lead); writing - original draft (lead); writing review and editing (equal). Alex Keen: Data curation (supporting); formal analysis (supporting); methodology (supporting); writing review and editing (supporting). Thomas Kitching: Data curation (supporting); validation (supporting); writing - review and editing (supporting). Drew Stokes: Resources (supporting); writing - review and editing (supporting). Simon J. Goodman: Conceptualization (equal); data curation (supporting); funding acquisition (equal); investigation (supporting); methodology (supporting); project administration (equal); resources (equal); supervision (lead); validation (equal); visualization (supporting); writing - original draft (supporting); writing - review and editing (equal).

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Ecology and Evolution

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20(3), 175-194. https://doi.org/10.1111/j.1365-3113.1995.tb000 90.x Cumming, G. S. (1998). Host preference in African ticks (Acari: Ixodida): A quantitative data set. Bulletin of the Entomological Research, 88(4), 379-406 Cuxim-Koyoc, A., Reyes-Novelo, E., Morales-Malacara, J. B., Bolívar-Cimé, B., & Laborde, J. (2015). Streblidae (Diptera: Hippoboscoidea) from Yucatan and updated species list for Mexico. Journal of Medical Entomology, 52(5), 947-961. https://doi.org/10.1093/jme/tjv117 de Vasconcelos, P. F., Falcão, L. A. D., Graciolli, G., & Borges, M. A. Z. (2016). Parasite-host interactions of bat flies (Diptera: Hippoboscoidea) in Brazilian tropical dry forests. Parasitology Research, 115(1), 367-377. https://doi.org/10.1007/s0043 6-015-4757-8 Dick, C. W., Gannon, M. R., Little, W. E., & Patrick, M. J. (2003). Ectoparasite associations of bats from Central Pennsylvania. Journal of Medical Entomology, 40(6), 813-819. https://doi. org/10.1603/0022-2585-40.6.813 Dick, C. W., & Miller, J. A. (2010). Streblidae (bat flies). In B. V. Brown, A. Borkent, J. M. Cumming, D. M. Wood, N. E. Woodley, & M. Zumbado (Eds.), Manual of central American Diptera. (Vol. 2, pp. 1249-1260). NRC Research Press. Dittmar, K., Porter, M. L., Murray, S., & Whiting, M. F. (2006). Molecular phylogenetic analysis of nycteribiid and streblid bat flies (Diptera: Brachycera, Calyptratae): Implications for host associations and phylogeographic origins. Molecular Phylogenetics and Evolution, 38(1), 155–170. https:// doi.org/10.1016/j.ympev.2005.06.008 Esser, H. J., Herre, E. A., Blüthgen, N., Loaiza, J. R., Bermúdez, S. E., & Jansen, P. A. (2016). Host specificity in a diverse Neotropical tick community: An assessment using quantitative network analysis and host phylogeny. Parasites and Vectors, 9(1), 1-14. https://doi. org/10.1186/s13071-016-1655-6 Estrada-Pena, A., Mangold, A. J., Nava, S., Venzal, J. M., Labruna, M., & Guglielmone, A. A. (2010). A review of the systematics of the tick family argasidae (Ixodida). Acarologia, 50(3), 317-333. https://doi. org/10.1051/acarologia/20101975

- Estrada-Villegas, S., Halczok, T. K., Tschapka, M., Page, R. A., Brändel, S. D., & Hiller, T. (2018). Bats and their bat flies: Community composition and host specificity on a Pacific Island archipelago. Acta Chiropterologica, 20(1), 161-176. https://doi.org/10.3161/15081 109ACC2018.20.1.012
- Folmer, O., Black, M., Hoeh, W., Lutz, R., & Vrijenhoek, R. (1994). DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Molecular Marine Biology and Biotechnology, 3(5), 294-299. https://doi.org/10.1371/ journal.pone.0013102
 - Frick, W. F., Hayes, J. P., & Heady, P. A. (2008). Island biogeography of bats in Baja California, Mexico: Patterns of bat species richness in a near-shore archipelago. Journal of Biogeography, 35(2), 353-364. https://doi.org/10.1111/j.1365-2699.2007.01798.x
 - Frick, W. F., Hayes, J. P., & Heady, P. A. (2009). Nestedness of desert bat assemblages: Species composition patterns in insular and terrestrial landscapes. Oecologia, 158(4), 687-697. https://doi.org/10.1007/ s00442-008-1168-x
 - Gay, N., Olival, K. J., Bumrungsri, S., Siriaroonrat, B., Bourgarel, M., & Morand, S. (2014). Parasite and viral species richness of Southeast Asian bats: Fragmentation of area distribution matters. International Journal for Parasitology: Parasites and Wildlife, 3(2), 161–170. https:// doi.org/10.1016/j.ijppaw.2014.06.003
 - Gill, J., Rowley, W., Bush, P., Viner, J., & Gilchrist, M. (2004). Detection of human blood in the bat tick Carios (Ornithodoros) kelleyi (Acari: Argasidae) in Iowa. Journal of Medical Entomology, 41(6), 1179-1181. https://doi.org/10.1603/0022-2585-41.6.1179
 - Gómez, A., & Nichols, E. (2013). Neglected wild life: Parasitic biodiversity as a conservation target. International Journal for Parasitology:

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CONFLICT OF INTEREST

The authors declare no competing interests.

DATA AVAILABILITY STATEMENT

Sequences for the haplotypes reported in this study are available under GenBank accession numbers OP963052-OP963192 for the COI set, and OP941218-OP941364 for the 18S set.

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REFERENCES

- Álvarez-Castañeda, S. T., Álvarez, T., & González-Ruíz, N. (2015). Guía para la identificación de los mamíferos de México en campo y laboratorio. Centro de Investigaciones Biológicas del Noroeste, S. C. y Asociacion Mexicana de Mastozoología, A. C.
- Black, W. C., IV, Klompen, J. S. H., & Keirans, J. E. (1997). Phylogenetic relationships among tick subfamilies (Ixodida: Ixodidae: Argasidae) based on the 18S nuclear rDNA gene. Molecular Phylogenetics and Evolution, 7(1), 129-144.
- Blood, B. R., & Clark, M. K. (1998). Myotis vivesi. Mammalian Species, 588, 1-5.
- Bolaños-García, R., Rodríguez-Estrella, R., & Guzmán-Cornejo, C. (2018). Ectoparásitos asociados a polluelos del Búho Cornudo (Aves: Strigidae) en un paisaje fragmentado de la península de Baja California, México. Acta Zoológica Mexicana, 34(1), 1-15. https:// doi.org/10.21829/azm.2018.3412142
- Borstein, S. (2018). AnnotationBustR: An R package to extract subsequences from GenBank annotations. https://cran.r-project.org/ web/packages/AnnotationBustR/vignettes/AnnotationBustR -vignette.html
- Bradshaw, G. V. R., & Ross, A. (1961). Ectoparasites of Arizona bats. Journal of the Arizona Academy of Science, 1(4), 109. https://doi. org/10.2307/40026556
- Braun, J. K., Yang, B., González-Pérez, S. B., & Mares, M. A. (2015). Myotis yumanensis (Chiroptera: Vespertilionidae). Mammalian Species, 47(918), 1-14. https://doi.org/10.1093/mspecies/sev001
- Burger, T. D., Shao, R., Labruna, M. B., & Barker, S. C. (2014). Molecular phylogeny of soft ticks (Ixodida: Argasidae) inferred from mitochondrial genome and nuclear rRNA sequences. Ticks and Tick-Borne Diseases, 5(2), 195–207. https://doi.org/10.1016/j. ttbdis.2013.10.009
- Campbell, B. C., Steffen-Campbell, J. D., Sorensen, J. T., & Gill, R. J. (1995). Paraphyly of Homoptera and Auchenorrhyncha inferred from 18S rDNA nucleotide sequences. Systematic Entomology,

Parasites and Wildlife, 2(1), 222-227. https://doi.org/10.1016/j. ijppaw.2013.07.002

- González-Abraham, C. E., Garcillán, P., & Ezcurra, E. (2010). Ecorregiones de la península de Baja California: una síntesis. Boletín de la Sociedad Botánica de México, 87, 69–82.
- Graciolli, G., Autino, A. G., & Claps, G. L. (2007). Catalogue of American Nycteribiidae (Diptera, Hippoboscoidea). Revista Brasileira de Entomologia, 51(2), 142–159. https://doi.org/10.1590/S0085 -56262007000200004
- Guglielmone, A. A., Robbins, R. G., Apanaskevich, D. A., Petney, T. N., Estrada-Peña, A., Horak, I. G., Shao, R., & Barker, S. C. (2010). The Argasidae, Ixodidae and Nuttalliellidae (Acari: Ixodida) of the world: A list of valid species names. *Zootaxa*, 2528, 1–28. https://doi. org/10.1023/A:1025381712339
- Guzmán-Cornejo, C., García-Prieto, L., Rebollo-Hernández, A., Venzal, J. M., Nava, S., & Sánchez-Montes, S. (2017). Molecular evidence and additional morphological characters to distinguish Ornithodoros brodyi and Ornithodoros yumatensis (Ixodida: Argasidae) in their different developmental stages. Acta Parasitologica, 62(2), 432-448. https://doi.org/10.1515/ap-2017-0051
- Hall, T. (2005). BioEdit. Biological sequence editor. http://www.mbio.ncsu. edu/BioEdit/bioedit.html
- Hebert, P. D. N., Ratnasingham, S., & DeWaard, J. R. (2003). Barcoding animal life: Cytochrome c oxidase subunit 1 divergences among closely related species. Proceedings of the Royal Society B: Biological Sciences, 270, 96–99. https://doi.org/10.1098/rsbl.2003.0025
- Herrera-Montalvo, L. G., Flores-Martínez, J. J., & Sánchez-Cordero, V. (2017). Geographical distribution and conservation status of an endemic insular mammal: The vulnerable fish-eating bat *Myotis vivesi*. *Oryx*, 53(2), 388–393. https://doi.org/10.1017/S0030605317000874
- Hornok, S., Szoke, K., Boldogh, S. A., Sándor, A. D., Kontschán, J., Tu, V. T., Halajian, A., Takács, N., Görföl, T., & Estók, P. (2017). Phylogenetic analyses of bat-associated bugs (Hemiptera: Cimicidae: Cimicinae and Cacodminae) indicate two new species close to *Cimex lectularius. Parasites and Vectors*, 10(1), 1–10. https://doi.org/10.1186/ s13071-017-2376-1
- Jobling, B. (1938). A revision of the species of the genus *Trichobius*: (Diptera Acalypterae, Streblidae). *Parasitology*, 30(3), 358–387. https://doi.org/10.1017/S0031182000025932
- Klimpel, S., & Mehlhorn, H. (Eds.). (2014). Bats (Chiroptera) as vectors of diseases and parasites. *Parasitology Research Monographs*, 5 (pp. 187). Springer-Verlag. https://doi.org/10.1007/978-3-642-39333-4
- Knee, W., & Proctor, H. (2006). Keys to the families and genera of blood and tissue feeding mites associated with albertan birds. *Canadian Journal of Arthropod Identification*, 2(2), 1–18. https://doi. org/10.3752/cjai.2006.02
- Letunic, I., & Bork, P. (2016). Interactive tree of life (iTOL) v3: An online tool for the display and annotation of phylogenetic and other trees. *Nucleic Acids Research*, *28*(3), 1–4. https://doi.org/10.1093/ nar/gkw290
- Librado, P., & Rozas, J. (2009). DnaSP v5: A software for comprehensive analysis of DNA polymorphism data. *Bioinformatics*, 25(11), 1451-1452.
- Mangold, A. J., Bargues, M. D., & Mas-Coma, S. (1997). 18S rRNA gene sequences and phylogenetic relationships of European hardtick species (Acari: Ixodidae). *Parasitology Research*, 84(1), 31–37. https://doi.org/10.1007/s004360050352
- Mans, B. J., de Klerk, D., Pienaar, R., de Castro, M. H., & Latif, A. A. (2012). The mitochondrial genomes of Nuttalliella namaqua (Ixodoidea: Nuttalliellidae) and Argas africolumbae (Ixodoidae: Argasidae): Estimation of divergence dates for the major tick lineages and reconstruction of ancestral blood-feeding characters. PLoS One, 7(11), e49461. https://doi.org/10.1371/journal.pone.0049461
- McDaniel, B. (1979). How to know the mites and ticks. W.C. Brown Co.

- Medellín, R. A., Arita, H. T., & Sánchez, O. (2007). *Identificación de los murciélagos de México. Claves de Campo* (2nd ed.). Universidad Nacional Autonóma de México, Instituto Nacional de Ecología.
- Morand, S., Krasnov, B. R., & Poulin, R. (2006). Micromammals and macroparasites: From evolutionary ecology to management. Springer Publishing.
- Najera-Cortazar, L. A. (2020). Ecological genomics of Myotis bats and their ectoparasites in the Baja California peninsula. PhD thesis, University of Leeds, UK.
- Nájera-Cortazar, L. A., Álvarez-Castañeda, S. T., & De Luna, E. (2015). An analysis of Myotis peninsularis (Vespertilionidae) blending morphometric and genetic datasets. Acta Chiropterologica, 17(1), 37–47. https://doi.org/10.3161/15081109ACC2015.17.1.003
- Nava, S., & Guglielmone, A. A. (2013). A meta-analysis of host specificity in Neotropical hard ticks (Acari: Ixodidae). Bulletin of the Entomological Research, 103(2), 216–224.
- Ossa, G., Johnson, J. S., Puisto, A. I. E., Rinne, V., Sääksjärvi, I. E., Waag, A., Vesterinen, E. J., & Lilley, T. M. (2019). The Klingon batbugs: Morphological adaptations in the primitive bat bugs, *Bucimex chilensis* and *Primicimex cavernis*, including updated phylogeny of Cimicidae. *Ecology and Evolution*, 9(4), 1736–1749. https://doi. org/10.1002/ece3.4846
- Pérez, T. M., Guzmán-Cornejo, C., Montiel-Parra, G., Paredes-León, R., & Rivas, G. (2014). Biodiversidad de ácaros en México. Revista Mexicana de Biodiversidad, 85(Suppl), 399-407. https://doi. org/10.7550/rmb.36160
- Porter, T. M., & Hajibabaei, M. (2018). Over 2.5 million COI sequences in GenBank and growing. PLoS One, 13(9), e0200177. https://doi. org/10.1371/journal.pone.0200177
- Poulin, R. (2014). Parasite biodiversity revisited: Frontiers and constraints. International Journal for Parasitology, 44(9), 581–589. https://doi.org/10.1016/j.ijpara.2014.02.003
- Rambaut, A., Drummond, A. J., Xie, D., Baele, G., & Suchard, M. A. (2018). Posterior summarization in Bayesian phylogenetics using Tracer 1.7. Systematic Biology, 67(5), 901–904. https://doi.org/10.1093/ sysbio/syy032
- Ramírez-Martínez, M., Ibarra-López, M. P., Iniguez-Dávalos, L. I., Yuill, T., Orlova, M. V., & Reeves, W. K. (2016). New records of ectoparasitic Acari (Arachnida) and Streblidae (Diptera) from bats in Jalisco, Mexico. *Journal of Vector Ecology*, 41(2), 309–313. https:// doi.org/10.1111/jvec.12228
- Ratnasingham, S., & Hebert, P. D. N. (2007). BOLD: The barcode of life data system (www.barcodinglife.org). Molecular Ecology Notes, 7, 355-364. https://doi.org/10.1111/j.1471-8286.2006.01678.x
- Ratnasingham, S., & Hebert, P. D. N. (2013). A DNA-based registry for all animal species: The Barcode Index Number (BIN) system. *PLoS* One, 8(8), e66213. https://doi.org/10.1371/journal.pone.0066213
- Reinhardt, K., & Siva-Jothy, M. T. (2007). Biology of the bed bugs (Cimicidae). Annual Review of Entomology, 52(1), 351–374. https:// doi.org/10.1146/annurev.ento.52.040306.133913
- Riddle, B. R., Hafner, D. J., Alexander, L. F., & Jaeger, J. R. (2000). Cryptic vicariance in the historical assembly of a Baja California peninsular desert biota. Proceedings of the National Academy of Sciences of the United States of America, 97(26), 14438-14443. https://doi. org/10.1073/pnas.250413397
- RStudio Team. (2021). RStudio: Integrated development for R. RStudio, PBC http://www.rstudio.com/
- Saldaña-Vázquez, R. A., Sandoval-Ruiz, C. A., Veloz-Maldonado, O. S., Durán, A. A., & Ramírez-Martínez, M. M. (2019). Host ecology moderates the specialization of Neotropical bat-fly interaction networks. *Parasitology Research*, 118(10), 2919–2924. https://doi. org/10.1007/s00436-019-06452-1
- Salinas-Ramos, V. B., Zaldívar-Riverón, A., Rebollo-Hernández, A., & Herrera-M, L. G. (2018). Seasonal variation of bat-flies (Diptera:

16 of 20

Streblidae) in four bat species from a tropical dry forest. *Mammalia*, 82(2), 133–143. https://doi.org/10.1515/mammalia-2016-0176

- Sánchez-Montes, S., Guzmán-Cornejo, C., Martínez-Nájera, Y., Becker, I., Venzal, J. M., & Labruna, M. B. (2016). *Rickettsia lusitaniae* associated with Ornithodoros yumatensis (Acari: Argasidae) from two caves in Yucatan, Mexico. *Ticks and Tick-Borne Diseases*, 7(6), 1097– 1101. https://doi.org/10.1016/j.ttbdis.2016.09.003
- Sándor, A. D., Corduneanu, A., Péter, Á., Mihalca, A. D., Barti, L., Csősz, I., Szőke, K., & Hornok, S. (2019). Bats and ticks: Host selection and seasonality of bat-specialist ticks in eastern Europe. *Parasites & Vectors*, 12(1), 605. https://doi.org/10.1186/s13071-019-3861-5
- Secretaría de Medio Ambiente y Recursos Naturales (SEMARNAT). (2010). Norma Oficial Mexicana NOM-059-SEMARNAT-2010. Protección ambiental-especies nativas de México de flora y fauna silvestres-Categorías de riesgo y especificaciones para su inclusión, exclusión o cambio-lista de especies en riesgo.
- Seneviratne, S. S., Fernando, H. C., & Udagama-Randeniya, P. V. (2009). Host specificity in bat ectoparasites: A natural experiment. International Journal for Parasitology, 39(9), 995–1002. https://doi. org/10.1016/j.ijpara.2008.12.009
- Speer, K. A., Luetke, E., Bush, E., Sheth, B., Gerace, A., Quicksall, Z., Miyamoto, M., Dick, C. W., Dittmar, K., Albury, N., & Reed, D. L. (2019). A fly on the cave wall: Parasite genetics reveal fine-scale dispersal patterns of bats. *Journal of Parasitology*, 105(4), 555. https://doi.org/10.1645/19-20
- Spencer, H. G., & Zuk, M. (2016). For host's sake: The pluses of parasite preservation. Trends in Ecology and Evolution, 31(5), 341–343. https://doi.org/10.1016/j.tree.2016.02.021
- Steinlein, D. B., Durden, L. A., & Cannon, W. L. (2001). Tick (Acari) infestations of bats in New Mexico. *Journal of Medical Entomology*, 38(4), 609–611. https://doi.org/10.1603/0022-2585-38.4.609
- Suchard, M. A., Lemey, P., Baele, G., Ayres, D. L., Drummond, A. J., & Rambaut, A. (2018). Bayesian phylogenetic and phylodynamic data integration using BEAST 1.10. *Virus Evolution*, 4(1), vey016.
- Sudhir, K., Stecher, G., Li, M., Knyaz, C., & Tamura, K. (2018). MEGA X: Molecular evolutionary genetics analysis across computing platforms. *Molecular Biology and Evolution*, 35, 1547–1549.
- Szentiványi, T., Christe, P., & Glaizot, O. (2019). Bat flies and their microparasites: Current knowledge and distribution. Frontiers in Veterinary Science, 6, 1–12. https://doi.org/10.3389/fvets.2019.00115
- Talbot, B., Vonhof, M. J., Broders, H. G., Fenton, B., & Keyghobadi, N. (2016). Range-wide genetic structure and demographic history in the bat ectoparasite *Cimex adjunctus*. *BMC Evolutionary Biology*, 16(1), 1–13. https://doi.org/10.1186/s12862-016-0839-1
- Talbot, B., Vonhof, M. J., Broders, H. G., Fenton, M. B., & Keyghobadi, N. (2017). Population structure in two geographically sympatric and congeneric ectoparasites: *Cimex adjuntus* and *Cimex lectularius*, in the North American Great Lakes region. *Canadian Journal of Zoology*, 95, 901–907.
- Ter Hofstede, H. M., & Fenton, M. B. (2005). Relationships between roost preferences, ectoparasite density, and grooming behaviour of

neotropical bats. *Journal of Zoology*, 266(4), 333–340. https://doi. org/10.1017/S095283690500693X

The National Library of Medicine. (2018). The Basic Local Alignment Search Tool (BLAST). The National Center for Biotechnology Information website: http://blast.ncbi.nlm.nih.gov/Blast.cgi

Ecology and Evolution

- Thompson, J. D., Higgins, D. G., & Gibson, T. J. (1994). CLUSTAL W: Improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Research*, 22, 4673–4680.
- Tortosa, P., Dsouli, N., Gomard, Y., Ramasindrazana, B., Dick, C. W., & Goodman, S. M. (2013). Evolutionary history of Indian Ocean Nycteribiid bat flies mirroring the ecology of their hosts. *PLoS One*, 8(9), e75215. https://doi.org/10.1371/journal.pone.0075215
- Trujillo-Pahua, L., & Ibáñez-Bernal, S. (2018). New geographical records of bat flies (Diptera: Streblidae) associated with phyllostomid bats (Chiroptera: Phyllostomidae) in the West Highlands of Mexico. *Journal of Medical Entomology*, 56, 18–28. https://doi.org/10.1093/ jme/tjy166
- Usinger, R. L. (1966). Monograph of Cimicidae (Hemiptera-Heteroptera). Entomological Society of America, VII, 572. http://apps.webofknowl edge.com/full_record.do?product=UA&search_mode=GeneralSea rch&qid=32&SID=Y2m6OyGYg7txy2Oc2bE&page=1&doc=3
- Wenzel, R., & Tipton, V. J. (1966). Some relationships between mammal hosts and their ectoparasites. In R. Wenzel & V. J. Tipton (Eds.), *Ectoparasites of Panama* (pp. 677–723). Field Museum of Natural History.
- Whiting, M. F. (2002). Mecoptera is paraphyletic: Multiple genes and phylogeny of Mecoptera and Siphonaptera. *Zoologica Scripta*, 31(1), 93–104. https://doi.org/10.1046/j.0300-3256.2001.00095.x
- Wilder, A. P., Kunz, T. H., & Sorenson, M. D. (2015). Population genetic structure of a common host predicts the spread of white-nose syndrome, an emerging infectious disease in bats. *Molecular Ecology*, 24, 5495–5506. https://doi.org/10.1111/mec.13396
- Zamora-Mejías, D., Morales-Malacara, J. B., Rodríguez-Herrera, B., Ojeda, M., & Medellín, R. A. (2020). Does latitudinal migration represent an advantage in the decrease of ectoparasite load in *Leptonycteris yerbabuenae* (Chiroptera)? *Journal of Mammalogy*, 101, 979-989. https://doi.org/10.1093/jmammal/gyaa075

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	I xəmiƏ	2 xəmiƏ	Cimex 3	4 xəmiƏ	suinuį́bo .C	sutonnotno.C	C. brevis	C. hemipterus	C. latipennis	C. lectularius	C. pipistrelle	sizA xəmiƏ	Oeciacus
Cimex 1													
Cimex 2	9.9												0
Cimex 3	16.3	16.0											0
Cimex 4	17.1	16.1	13.5										0
C. adjuntus	19.9	17.1	12.6	10.5									0
2. antennatus	16.9	15.2	12.1	9.3	9.7								L
C. brevis	14.2	13.1	13.5	14.6	15.8	14.3							L
C. hemipterus	19.3	17.9	20.9	18.1	19.6	19.7	17.7						0
C. latipennis	10.6	7.2	13.9	16.2	17.1	15.2	13.7	19.1					
C. lectularius	18.7	17.3	19.8	18.0	19.1	17.9	17.4	18.5	18.6				
C. pipistrelli	19.5	16.8	20.0	17.5	18.1	17.3	16.6	15.3	17.8	15.0			
Cimex Asia	19.1	16.4	20.1	17.8	17.8	17.4	16.6	15.3	17.5	15.2	2.3		L
Jeciacus	20.1	17.7	20.7	18.2	18.1	17.8	17.0	15.2	18.7	15.7	6.9	7.2	v
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Nycteribiid 1													0.35
Nycteribiid 2	4	1.4											0.69
Vycteribia	1	10.7	10.5										5.91
Penicillidia	1	11.7	11.4	9.8									5.75
3asilia 1	1	14.1	13.0	13.3	13.6								0.0
3asilia 2a	1	13.5	11.4	12.9	14.6	9.8							0.10
3asilia 2b	1	12.9	12.7	13.6	15.1	9.2	2.9						0.09
3asilia boardmani	1	14.5	13.9	13.0	13.2	4.9	10.3	9.9					0.00
3 <i>asilia</i> America	1	13.4	12.7	11.9	12.6	8.5	10.5	10.6	8.8				6.54
													2

18 of 20 WILEY_Ecology and Evolution

Estimates of percentage divergence (COI) over sequence-pairs between groups (diagonal left matrix), and within groups (rightmost column) of bat flies (family Streblidae), showing the lineages obtained in this study (row label shaded) versus closely related reference sequences from GenBank. Names in the first column have the genus abbreviated. n/c - not calculated as only a single sequence available.

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squorg nithiW	3.3	n/c	0.2	0.5	n/c	1.1	0.9	0.0	0.3	1.1	0.0	n/c	0.2	0.6	4.7	0.2	1.2	9.1	0.0	13.3	0.2	
Trichobius urodermae																						
qs suidon7riT																					13.1	
Trichobius major																				15.1	15.1	
Strebla sp.																			16.1	14.6	14.3	
รมาวignol suidohว่าว่าวาดจ																		15.9	15.6	15.3	14.1	
Paratrichobius dunni																	8.9	15.2	16.0	15.2	13.7	
Nycterophilia parnelli																18.7	18.6	17.2	17.5	16.8	18.1	
Nycterophilia fairchildi															7.4	18.9	19.7	16.5	14.5	16.7	18.6	
2 nənana aboqotsigəM														16.8	17.7	14.4	13.2	14.5	20.2	15.1	14.0	
1 nənna nboqotsigəM													3.0	17.9	17.5	15.9	14.4	15.5	21.6	16.0	15.0	
sutonorsphae suidodoiT												18.2	15.8	14.6	15.8	13.6	13.9	13.2	11.9	13.4	14.7	
suibsmrstni suidodoinT											15.4	15.5	15.3	17.0	16.5	15.6	15.7	15.6	15.8	14.1	9.3	
iisəgub zuidodəirT										9.2	15.1	14.4	13.5	16.6	16.3	13.5	13.9	13.3	17.0	12.4	7.5	
2 suidorhiT									5.7	11.0	13.0	15.1	14.3	16.5	17.8	12.4	14.3	14.4	13.3	13.2	8.7	
1 suidohin1								13.5	13.1	14.8	6.2	19.3	17.7	15.3	15.8	13.5	13.5	13.3	10.8	13.4	13.9	
1 bildər)2							12.1	15.5	14.2	15.9	10.8	14.8	14.3	14.8	14.9	14.5	14.3	13.3	13.1	13.7	15.3	
Paratrichobius 1						11.9	13.6	12.8	12.5	14.8	13.3	13.4	12.0	19.6	17.1	8.4	4.5	14.5	15.3	14.3	13.9	-531219
υγοτεγορημια ςοχατα					18.9	15.5	16.3	18.1	17.3	17.3	17.8	19.0	18.8	7.1	3.1	20.2	21.0	18.2	13.7	17.6	19.6	umber <mark>E</mark> l
ds ppodojsiĝaM				18.4	13.3	15.2	17.3	15.0	13.7	16.0	16.9	3.0	4.0	17.5	17.6 8	15.4	14.1	15.0	50.0	15.8	[4.9	ession n
			4	.6	ω.	1.2		1.2	9.0	9.6	6.9	с С	~	7.9	č.	4.	1.	9.1	0.3	4.0	1.2	ank acc
asnana aranea Bela		Ŀ	.8	0 18	.2 11	.1	.6 17	17	3 13	.9 15	.9 15	.0	.0	.1 16	.6 17	.5 14	.4	.0	.4 20	.2 15	14	: GenB
sitamotsollynq arstqobiqsA	s.	15	14	17.	13	15	13	9.6	8.0	10	13	16	14	17.	17.	13	13	15	13	14	7.7	ranea 1
	. phyllostomati	1. aranea	1. sp	. coxata	aratrichobius 1	treblid 1	richobius 1	richobius 2	dugesii	intermedius	sphaeronotus	1. aranea 1 ^a	1. aranea 2	I. fairchildi	I. parnelli	dunni	longicrus	trebla sp.	major	richobius sp	urodermae	legistopoda a
	4	2	2	2	4	3				-	-	~	<	~	~	4	4	5	Ľ	L		2

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Estimates of evolutionary divergence (COI) in percentage over sequence-pairs between groups (diagonal left matrix), and within groups (rightmost column) of bat ticks (family Argasidae), showing the lineages obtained in this study (row label shaded) versus closely related reference sequences from GenBank. n/c - not calculated as only a single sequence available.

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squorg nirthiW	0.99	0.10	0.13	0.56	4.97	0.33	0.07	11.83	11.12	n/c	18.70	14.74	14.67	0.99
Grnithodoros Africa														
Ornithodoros Americ														16.4
sizA zovobodiinvO													17.7	18.1
.qs siqspottoN												17.7	15.9	16.3
.qs soino											15.0	19.1	15.2	15.9
.qs nitricola sp.										15.1	14.2	17.8	15.0	16.6
Тіс қ 6									14.3	15.2	16.8	19.3	13.9	16.5
ΤίςΚ 5								15.4	14.2	14.1	14.6	17.9	15.7	16.3
Tick 4							11.4	14.3	14.8	14.7	14.6	17.2	14.7	16.4
Тіск 3						12.1	12.6	15.0	15.0	14.4	15.8	17.9	15.5	16.3
Tick 2					12.7	11.5	11.3	14.2	14.4	13.9	13.7	17.1	14.2	15.2
Tick 1				11.4	12.9	11.9	12.5	13.4	15.2	14.3	14.8	18.5	15.1	16.4
Carios kelleyi			11.8	6.1	12.5	11.9	12.7	14.5	14.6	14.0	13.5	18.3	14.6	16.2
L plozintnA		13.9	13.1	13.9	13.3	13.7	12.7	13.7	13.7	14.0	14.4	17.6	15.3	16.2
	Antricola 1	Carios kelleyi	Tick 1	Tick 2	Tick 3	Tick 4	Tick 5	Tick 6	Antricola sp.	Carios	Nothoaspis	Ornithodoros Asia	Ornithodoros America	Ornithodoros Africa