

1 **Approaches to integrating genetic data into ecological networks**

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3 Running Head: Molecular Food Webs

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5 Elizabeth L. Clare^{1,2}, Aron J. Fazekas³, Natalia V. Ivanova², Robin M. Floyd⁴, Paul D.N.

6 Hebert², Amanda M. Adams⁵, Juliet Nagel⁶, Rebecca Girton¹, Steven G. Newmaster³, M.

7 Brock Fenton⁷

8

9 ¹ School of Biological and Chemical Sciences, Queen Mary University of London.

10 London UK. E14NS

11 ²Centre for Biodiversity Genomics, University of Guelph, Guelph Ontario, Canada N1G

12 2W1

13 ³Biodiversity Institute of Ontario, University of Guelph, Guelph Ontario, Canada N1G

14 2W1

15 ⁴Welcome Trust Stem Cell Institute, University of Cambridge, Cambridge, UK

16 ⁵Department of Biology, Texas A&M University, 3258 TAMU, College Station 77843

17 USA

18 ⁶University of Maryland, Center for Environmental Science, Frostburg, MD, USA

19 ⁷Department of Biology, University of Western Ontario, London, ON N6A 5B7, Canada

20

21 Corresponding Author: Elizabeth Clare, School of Biological and Chemical Sciences,

22 Queen Mary University of London. London UK. E14NS, e.clare@qmul.ac.uk

23

24 **Abstract**

25 As molecular tools for assessing trophic interactions become common, research is
26 increasingly focused on the construction of interaction networks. Here we demonstrate
27 three key methods for incorporating DNA data into network ecology and discuss
28 analytical considerations using a model consisting of plants, insects, bats and their
29 parasites from the Costa Rican dry forest. The simplest method involves the use of
30 Sanger sequencing to acquire long sequences to validate or refine field identifications, for
31 example of bats and their parasites, where one specimen yields one sequence and one
32 identification. This method can be fully quantified and resolved and these data resemble
33 traditional ecological networks. For more complex taxonomic identifications, we target
34 multiple DNA loci e.g. from a seed or fruit pulp sample in faeces. These networks are
35 also well resolved but gene targets vary in resolution and quantification is difficult.
36 Finally for mixed templates such as faecal contents of insectivorous bats we use DNA
37 metabarcoding targeting two sequence lengths (157bp, 407bp) of one gene region and a
38 MOTU, BLAST and BIN association approach to resolve nodes. This network type is
39 complex to generate and analyse and we discuss the implications of this type of
40 resolution on network analysis. Using these data we construct the first molecular-based
41 network of networks containing 3304 interactions between 762 nodes of 8 trophic
42 functions and involving parasitic, mutualistic, and predatory interactions. We provide a
43 comparison of the relative strengths and weaknesses of these data types in network
44 ecology.

45 Key Words: food webs, interaction networks, DNA barcoding, metabarcoding, high-
46 throughput sequencing, bats

47 **Introduction:**

48 *Ecological Networks, DNA & Opportunities*

49 Ecosystem functioning is driven by a network of interactions among species
50 affected by diverse abiotic and biotic variables such as climate, habitat, and resource
51 distribution (McCann, 2007) with global economic (Costanza et al., 1997) and
52 conservation (Worm et al., 2006) impacts. The analysis of interaction networks is of
53 increasing interest across many disciplines, spurring the development of new
54 mathematical and statistical tools (Poisot, Stouffer, & Kéfi, 2016). In ecology, visual
55 representations provide a synoptic view of complex interactions and are primarily
56 displayed as bipartite networks, where trophic levels are depicted as layers (upper and
57 lower) composed of species as nodes connected by links representing interactions
58 (Dormann, Fründ, Blüthgen, & Gruber, 2009). When multiple networks are combined, it
59 is possible to conceptualise multiple trophic levels simultaneously (e.g., Pocock, Evans
60 and Memmott 2012) clarifying ecosystem assembly and structure (Milo et al., 2002),
61 functional roles, and mechanisms of stability (McCann, 2000; Thébault & Fontaine,
62 2010). Comparisons between networks can assess natural or anthropogenic impacts
63 (McCann, 2007), the evolution of networks (Guimarães Jr, Jordano, & Thompson, 2011;
64 Nuismer, Jordano, & Bascompte, 2013) and the role and response of specific nodes
65 (Martín González, Dalsgaard, & Olesen, 2010; McDonald-Madden et al., 2016; Strona &
66 Lafferty, 2016).

67 Many networks are incredibly complex with multiple trophic levels and high
68 taxonomic diversity (e.g. Pocock et al. 2012), and are therefore time consuming to
69 construct, often requiring years of ecological observations and considerable taxonomic

70 expertise (Evans, Kitson, Lunt, Straw, & Pocock, 2016). Consequently, they are not
71 readily scalable to rapid bio-monitoring or, if they can scale, they routinely suffer from
72 problems of network completeness and poor or uneven resolution of taxa (Hemprich-
73 Bennett, Oliveira, Le Comber, Rossiter, & Clare, 2018; Ings et al., 2009). Incorporating
74 dietary tracers such as fatty acids, isotopes, and genetic tools such as DNA sequencing is
75 a growing trend for measuring species interactions, though each has advantages and
76 disadvantages (reviewed in Nielsen, Clare, Hayden, Brett, & Kratina, 2018). Genetic
77 analyses are expanding at a remarkable rate and have evolved from enzyme-linked
78 immunosorbent assay (ELISA) and targeted species detection (Symondson, 2002) to the
79 use of high-throughput sequencing (HTS) for the analysis of target markers or
80 “metabarcoding” (reviewed in Pompanon et al., 2012). While these techniques are
81 quickly becoming common for the dietary analysis of single species with many proposed
82 applications (Clare, 2014), they have not been widely incorporated into network analysis
83 (but see González-Varo, Arroyo, & Jordano, 2014; Hemprich-Bennett et al. 2018; Wirta
84 et al., 2014) despite strong advocates (Evans et al., 2016; Roslin & Majaneva, 2016).

85 Many reviews, authors, and developers of these techniques have discussed the
86 challenges in DNA-based analyses of species interactions including the impacts of primer
87 choice on taxonomic coverage and resolution, the completeness of reference databases
88 (Pompanon et al., 2012), bioinformatics methods (Clare, Chain, Littlefair, & Cristescu,
89 2016) and the role of quantification (Deagle et al., 2018) but the specific implications for
90 constructing networks vary with data type. In traditional DNA barcoding a specimen’s ID
91 is delimited by generating one sequence per specimen and comparing it to a reference
92 dataset to confirm its identity. These data are not fundamentally different from traditional

93 observations for generating an interaction matrix. However, at the other end of the
94 complexity spectrum, metabarcoding represents a novel data type for network ecology.
95 HTS generates millions of sequences from each sample of mixed template. While the
96 process can uniquely deal with otherwise intractable sources such as trace material and
97 liquid feeders, it poses novel problems for ecological analysis and network ecology. First,
98 the data require complex bioinformatics handling to remove unwanted (often error prone)
99 data, but in many cases the impacts of these parameter choices on ecological analysis are
100 unknown (Clare et al., 2016). Second, the ability to quantify DNA within a sample is
101 highly controversial and while, in some cases, proportions of recovered sequence
102 correspond to biomass, in other cases the connection is not clear (Deagle et al., 2018;
103 Deagle, Thomas, Shaffer, Trites, & Jarman, 2013; Thomas, Deagle, Eveson, Harsch, &
104 Trites, 2016). Finally, in an ideal situation, the recovered sequences are matched to a
105 complete reference dataset to identify taxa, but in most cases the reference library is
106 incomplete. In these cases either an incomplete network is created biased towards the
107 contents of the reference collection (often larger, more charismatic or economically
108 important taxa) or the recovered pool of DNA is converted into molecular operational
109 taxonomic units - MOTUs (Floyd, Abebe, Papert, & Blaxter, 2002) - which are best
110 viewed as pools of equivalent genetic diversity that can be compared, whether we know
111 their identity or not (Clare et al., 2016; Floyd et al., 2002). In this case, each MOTU
112 becomes a node in the network and this level of the network is entirely resolved to a
113 common point of reference with both known and unknown items included, a distinct
114 advantage when mixed resolution presents an analytical problem (Hemprich-Bennett et
115 al., 2018; Ings et al., 2009). However, the level of this resolution is arbitrarily defined by

116 the bioinformatics assessment (see an analysis of parameters for MOTU definition in
117 ecological analysis (Clare et al., 2016; Flynn, Brown, Chain, MacIsaac, & Cristescu,
118 2015) which may collapse trophic levels and thus generate fundamentally different
119 structures. This is of importance when selecting what metrics can or should be measured.
120 Networks metrics can be divided into several broad classes. Network level metrics (e.g.
121 connectance, nestedness, generality) are measured across the entire network. Node level
122 metrics (e.g. centrality, species strength, partner diversity) are specific to the interactions
123 of a given node. Motif measurements are sub-network of a particular pattern which may
124 define specific ecological interaction types or functions (Milo et al., 2002). Each metric
125 type needs to be considered separately in light of the new data type. Many have
126 concluded that the molecular approach will transform the discipline of ecological
127 biomonitoring and ecological network analysis permitting rapid consistent assessments in
128 systems that are otherwise intractable (Gibson et al., 2015; Roslin & Majaneva, 2016;
129 Toju, 2015; Wirta et al., 2014) while others have advocated adoption but raised serious
130 analytical concerns (e.g. Evans et al., 2016). How then might we proceed?

131 Here we address this challenge by focussing on a single complex assemblage of
132 interacting species to demonstrate three approaches to the use of DNA data to resolve
133 interactions and measure several network and node level metrics. While these are not
134 without controversy, our objective is to present an example of methods of data integration
135 into a “network of networks” and we include the most commonly analysed interaction
136 types (antagonistic, mutualistic, parasitic) and the three key methods that have been
137 discussed for DNA and network integration. First, we use standard single-gene DNA
138 barcoding to resolve taxonomy in cryptic organisms and to validate field identifications.

139 Second, we use multi-gene DNA barcoding to resolve more complex taxa and single-
140 sourced trace material that cannot be identified by morphological methods. Third, we use
141 metabarcoding to resolve mixed material, and then discuss the advantages and challenges
142 of applying these approaches. While these have been used previously, our analysis
143 provides the first example of integrating these data types to form a multi-trophic level
144 assemblage resolved entirely with DNA and the first to contrast these data. We hope to
145 provoke discussion about the appropriate use of these data types.

146

147 **Materials and Methods:**

148 *A case study from Cost Rica: Plants, Bats, Insects, and Parasites*

149 From May to July 2009, a field team visited Sector Santa Rosa of the Area de
150 Conservación Guanacaste (ACG). The present analysis relies on material collected during
151 that period and a preliminary ecological analysis of this case is presented (Box 1). A total
152 of 801 bats were captured representing 26 species morphologically identified using
153 available field keys and checklists (Reid, 2009; Simmons, 2005). From these individuals
154 we analyzed 466 parasites that were sampled from 18 host species and 260 guano
155 samples from 21 species of which visual inspection led to 132 samples being classified as
156 containing plant materials (seeds or fruit pulp) and the rest insect material. Some species
157 are integrated into all trophic levels while others are only loosely associated, for example
158 the sanguivore *Desmodus rotundus* did not produce a faecal sample so was retained in the
159 dataset as a parasite host only.

160

161 *Method one: Sanger sequencing to resolve species ID of bats and parasites*

162 The simplest way to integrate DNA data involves the use of Sanger sequencing to
163 clarify species boundaries or to confirm and improve upon identifications made in the
164 field. For bats and parasites we targeted the 5' end of the mitochondrial cytochrome *c*
165 oxidase subunit 1 gene (COI) as described by Hebert, Ratnasingham, & DeWaard, (2003)
166 using full length (658 bp) DNA barcodes which provide taxonomic discrimination for
167 most animal groups (Hebert, Cywinska, Ball, & DeWaard, 2003). For bats we used small
168 tissue fragments from each individual captured and the “routine” method of DNA
169 barcoding described in Ivanova, Clare, & Borisenko, (2012) and the mammal primer
170 cocktail, PCR reagent mix and the thermocycler conditions “MamCOI” described in
171 Tables S1, S2 and S3 of that publication. We edited sequences in CodonCode Aligner
172 (CodonCode Corporation, Centerville, MA) and compared the resulting DNA barcodes to
173 existing reference databases (Clare, Lim, Fenton, & Hebert, 2011) using a Neighbor-
174 Joining (NJ) tree to confirm they clustered with other representatives of their species
175 assignment based on morphological inspection in the field (Figure S1). Sequences,
176 collection information, and primer names are available in the Barcode of Life Data
177 System (BOLD) (www.barcodinglife.org) (Ratnasingham & Hebert, 2007) project BCCR
178 for each recovered sequence.

179 For parasitic flies and mites, we extracted DNA from whole specimens using
180 voucher-retention procedures (Porco, Rougerie, Deharveng, & Hebert, 2010). Our
181 subsequent PCRs used a variety of primer combinations which are associated with
182 individual records available in the project BCPB available in the BOLD website with
183 corresponding primer sequences online at
184 http://www.boldsystems.org/index.php/Public_Primer_PrimerSearch. Our PCR protocols

185 followed (Hebert et al., 2013) with post sequence analysis employing CodonCode
186 Aligner (CodonCode Corporation, Centerville, MA). Unlike bats, field taxonomic
187 designations were minimal and a full reference database of voucher-linked barcodes
188 was not available. As a consequence, we employed the Barcode Identification
189 Number (BIN) (Ratnasingham & Hebert, 2013) method of delimiting MOTUs in
190 BOLD to identify species and compared this to terminal clusters in an NJ tree
191 generated in BOLD. Three clusters were unassigned to any BIN because their
192 sequence lengths were insufficient to provide a sequence match with high
193 confidence; therefore, we designate these as taxa based on reciprocal monophyly of
194 their sequences in an NJ tree (Figure S2).

195

196 *Method two: Sanger sequencing with multiple targets*

197 A more complex problem involves the analysis of material from one source when
198 that material is degraded, making DNA analysis a preferred option, but where the taxa
199 involved are difficult to resolve using this approach. In this case, the seeds defecated by
200 bats may be identifiable from morphology, but fragmented seeds and digested fruit pulp
201 are almost never identifiable morphologically. Consequently, plants whose seeds are too
202 large to be swallowed are often excluded from food webs and dietary analyses unless
203 direct observation confirms their consumption. Plants represent an additional hurdle as
204 species delimitation by DNA often requires multiple genetic markers (CBOL plant
205 working group, 2009).

206 For all guano samples containing seeds, we separated three to five intact
207 morphologically identical seeds from each sample. For samples containing only pulp or

208 pollen we separated approximately 10 mg of dried guano for DNA extraction. DNA
209 isolated employed the NucleoSpin® 96 Plant II DNA isolation kit (Macherey-Nagel)
210 following the manufacturer's protocol with an extension of lysis to one hour.

211 We amplified the *rbcL* and *trnH-psbA* regions using primers
212 *rbcLa_F/rbcLajf_634R* and *trnH/psbA* (Fazekas et al., 2008; Kress, Wurdack, Zimmer,
213 Weigt, & Janzen, 2005). We amplified *matK* using primers *1R_KIM/3F_KIM* (Fazekas
214 et al., 2008) and repeated the PCR for failed reactions using alternate primers:
215 *XF/3F_KIM* (Fazekas et al., 2008; Ford et al., 2009). PCRs were carried out in 10µL
216 volumes containing 2µL of 5X Phire® reaction buffer (Finnzymes), 0.05µL of 10mM
217 dNTPs, 0.1µL of each 10µM primer, and 0.125µL of Phire® Hot Start II polymerase
218 (Finnzymes) using the following protocol: initial denaturation at 98°C for 90s, 35 cycles
219 of 98°C for 5s, 55–66°C for 10s (depending on primer set), 72°C for 7–10s (depending
220 on region), followed by a final extension at 72°C for 60s and hold at 4°C (see primer
221 references for additional details).

222 We sequenced each amplicon bi-directionally with the same primers used for
223 amplification in 11µL reaction volumes containing 0.5 µL of BigDye terminator mix
224 (ABI), 2µL of 5X sequencing buffer, 1µL 10uM primer, and 0.5µL of undiluted PCR
225 product using the following protocol: initial denaturation at 96°C for 2min, 30 cycles of
226 96°C for 30s, 55°C for 15s, 60°C for 4min, followed by a hold at 4°C.

227 We assembled contigs and edited all sequences using Sequencher 4.8 (Gene
228 Codes Corp, Ann Arbor, MI). We then ascertained the percentage similarity of all
229 recovered sequences to available reference sequences in GenBank and BOLD, with the
230 exception of the *trnH-psbA* region which was not searchable within BOLD.

231 Identification to known taxa is more complex as the different regions provide
232 resolution at different taxonomic depths in different taxa. For example, *rbcL* typically
233 provides generic level resolution (CBOL plant working group, 2009) (occasionally to
234 species level), whereas the *matK* and *trnH-psbA* regions can provide resolution to species
235 in ~60-90% of cases (depending on the taxa and geographic scope) (Braukmann,
236 Kuzmina, Sills, Zakharov, & Hebert, 2017; Burgess et al., 2011; Lahaye et al., 2008).
237 Due to incompleteness of the reference sequence databases for the flora, many sequences
238 did not show 100% identity to any species in the reference database. We therefore
239 assigned sequences to family, genus, or species depending on the region and percent
240 identity using the following criteria. For *rbcL*, sequence matches with 99.75-100%
241 identity were assigned to a genus, while matches with 99-99.75% identity were only
242 placed to a family. For *matK*, matches with 100% identity were assigned to a species or a
243 species cluster when more than one species in the reference set matched with 100%
244 identity; matches with 99.0-99.9% identity were assigned to genus, while matches with
245 98-99% identity were only assigned to a family. For the *trnH-psbA* region, most matched
246 sequences ranged from 98-99% identity (no 100% matches were observed). The variable
247 length of the region, the presence of repeated sequence motifs, and the small number of
248 reference sequences complicated the interpretation of BLAST analysis with the GenBank
249 dataset so most assignments were only made to a genus. For two genera, however, the
250 *trnH-psbA* data corroborated the *matK* designation and enabled an increased level of
251 resolution. Unique sequences for these samples were therefore designated with a number
252 (in addition to genus) and treated as putative species. Species-level designation was also

253 accepted for sequences that matched a monotypic genus, and where sequences matched a
254 genus of three species, two of which occur outside the study area.

255

256 *Method three: Metabarcoding of mixed unknowns*

257 When the material to be analysed is a mixed sample of unknown taxa (in this case
258 arthropods), the entire assemblage must be targeted, followed by the use of
259 bioinformatics tools to process the sequences and ascertain the number of taxa in each
260 sample. In this case, we used DNA metabarcoding that targeted two segments of the COI
261 DNA barcode region and processed these data using a series of bioinformatics tools in a
262 well established analytical pipeline (e.g. Alberdi, Aizpurua, Gilbert, & Bohmann, 2018;
263 Clare, Symondson, & Fenton, 2014; Salinas-Ramos, Herrera Montalvo, León-Regagnon,
264 Arrizabalaga-Escudero, & Clare, 2015). In brief, DNA was extracted using Qiagen Stool
265 mini-kits (Qiagen CA) with modifications from (Clare et al., 2014; Zeale, Butlin,
266 Barker, Lees, & Jones, 2011) and eluted in 35µL of molecular grade water. We
267 targeted 157bp and 407bp amplicons of the DNA barcode region. PCRs were
268 conducted in 20µL reactions containing 10µL of Qiagen multiplex master mix
269 (Qiagen CA), 6µL of water, 1µL of each 10µM primer and 2µL of DNA. PCR reactions
270 were: 95°C – 15 min; 50 cycles of 95°C - 30 sec, 52°C – 30 sec, 72°C – 30 sec (1 min
271 for the 407bp region); 72°C – 10 min. Amplicons were visualized on 96 well 2%
272 agarose pre-cast E-gels (Invitrogen, Life Technologies). For the 157bp region we
273 used the Zeale primers (Zeale et al., 2011) which do not amplify bat DNA well,
274 modified with the two adaptor molecular identification tags (MIDs) system to
275 identify individual samples (Clare et al., 2014) without pooling. For the 407bp

276 region we used primers MLepF1 (GCTTTCCCACGAATAAATAATA) and
277 RonMWASPdeg (GGWTCWCCWGATATAKCWTTTCC) combined in equal quantities
278 with LepR1 (TAAACTTCTGGATGTCCAAAAAATCA) and HCO2198
279 (TAAACTTCAGGGTGACCAAAAAAATCA). Sequence recovery is predicted to be lower
280 with longer amplicons due to DNA degradation in digested material but longer reads
281 maximize identification. For this region we extracted and PCR-amplified all samples
282 independently but unlike the Zeal region we did not use (MIDs). This does not
283 impact MOTU estimates, but we cannot assign individual sequences to their source
284 but so they were analysed as a pool and we do not generate networks from these
285 data, just compare MOTU estimation from alternative regions.

286 PCR products were pooled without normalization and 70 μ L of the pooled product
287 was cleaned using the PCRClean DX kit (Aline Biosciences) for a double size selection
288 purification protocol (Table S1). Purified PCR products were eluted in 36 μ L of water.
289 The concentration was measured on the Qubit 2.0 spectrophotometer using a Qubit
290 dsDNA HS Assay Kit (Invitrogen, Life Technologies). All products were normalized to
291 1ng/ μ L prior to final library dilution. Sequencing was performed using the Ion PGM
292 Template OT2 400 kit for template preparation according to manufacturer's instructions,
293 except for a ~2-3x recommended dilution with water (Table S2) and a 316 chip. After the
294 chip check (prior to loading), the chip was flushed once with 100 μ L of 100% isopropanol
295 and three times with 100 μ L of annealing buffer.

296

297 Bioinformatics analysis

298 The sequences were processed using two analytical pipelines for comparison
299 First, we employed well established tools in the Galaxy platform (Afgan et al., 2016).
300 Reads from the 157bp Zeal region were separated by MID allowing 2 indels and 2
301 mismatches using the barcode splitter tool. For both the 157bp and 407bp datasets
302 primers, (MIDs for the 157bp region) and adaptors were removed using the clip tool
303 (both tools from the FASTX tool kit (Assaf Gordon (2010). FASTQ/A short-reads pre-
304 processing tools. http://hannonlab.cshl.edu/fastx_toolkit/). The resulting
305 amplicons were filtered for length (157bp or 407bp \pm 10bp) and dereplicated
306 (Figure S3) using the Collapose tool (FASTX tool kit). We used custom scripts to
307 remove singletons (Table S3).

308 For the 157bp dataset we clustered the remaining haplotypes into MOTUs at
309 90-97% similarity in QIIME using the pick_otu and uclust methods
310 (<http://qiime.sourceforge.net/>). See Clare et al., (2016) and Flynn et al., (2015) for a
311 discussion of MOTU thresholds. For each dataset we used a BLAST analysis
312 interpreted in MEGAN (Huson, Mitra, Ruscheweyh, Weber, & Schuster, 2011) to
313 filter out MOTUs that could not be reliably assigned to an arthropod order using a
314 reference database of >600,000 COI sequences extracted from GenBank. Parameters
315 in Megan were: Max Expected =0.01, Top Percent =10, Min Support Percent (Off),
316 Min Support =1, Min Complexity =0.2. Min Score =250. We tested a representative
317 sequence from each MOTU in UCHIME as implemented in MOTHUR (Schloss et al.,
318 2009) to filter out MOTUs that were likely to be chimeras.

319 For each MOTU dataset (90-97% clustering) we examined a BLAST
320 assignment for MOTU representatives in MEGAN. If two or more reads were

321 assigned to the same species we considered MOTUs oversplit, rejected that
322 threshold and tested the next most conservative option. We particularly considered
323 assignments in the Lepidoptera because this order was heavily represented in the
324 reference database. A QIIME threshold of 92 minimized MOTU oversplitting and this
325 data set was used for network construction. The same analysis was performed for
326 the 407bp dataset (MEGAN Min Score =500) but without network construction
327 (Figure S4).

328 We further queried the 157bp and 407bp datasets by comparing all sequences
329 to the same reference sequence library and to a reference library provided by D. Janzen
330 and W. Hallwachs generated from specimens (primarily Lepidoptera) from the study area
331 visualised in MEGAN (Figure S5 and S6) and with custom BLAST parsing scripts. This
332 analysis extracts species-level identifications, but is biased towards identification of
333 Lepidoptera, which dominate the reference database from the study location, and the
334 accuracy of database curation (e.g., databases generally provide better resolution of
335 larger, more charismatic, and economically important species).

336 For a second comparative analytical approach we used a non-MOTU based
337 method. Initial steps were similar with reads processed in Galaxy to split by MID and
338 remove primers using cutadapt (<https://cutadapt.readthedocs.io/en/stable/guide.html>).
339 FastQ files were then transferred to the mBRAVE platform (<http://www.mbrave.net/>) and
340 processed using the parameters trim front 0bp, trim end 0bp (primers and adaptors had
341 already been removed via cutadapt) trim length 500bp and filtering of MinQV 0qv, min
342 length 147bp, and max bases with low or ultra low QV of 100% (to avoid specific quality
343 filter parameters. We set a pre-clustering threshold of none, and ID distance threshold of

344 2% and left OTU thresholds as pre-set standards as we ignored MOTU analysis for this
345 comparison. Chimera screening and dereplication was performed automatically by the
346 mBRAVE platform. The resulting data is automatically compared to the BOLD system
347 library for insects using the BIN approach to attempt to associate the reads with known
348 BINs (this library contained 580, 824 reference sequences from 434, 878 known BINs,
349 last updated 21, Oct. 2018). The resulting dataset was then converted to a matrix of bat
350 species vs. associated prey BINs for further analysis.

351

352 *Network analysis:*

353 Using the data produced by all three approaches, we constructed a “network of
354 networks” in Cheddar (Hudson et al., 2013) and Bipartite (Dormann et al., 2009) in R (R
355 Development Core Team 2008) which represents all identified taxa or MOTUs. This
356 network has differing levels of resolution based on the trophic level or taxonomic group.
357 As the bats are, with a single exception, resolved to species level, they are fully
358 quantified. The mites and flies that feed on them are identified by a Barcode
359 Identification Number (BIN) (Ratnasingham & Hebert, 2013) as a proxy for species. This
360 trophic level is also well resolved, but individual taxa are only partially quantified from
361 each bat (finding all individual parasites is not practically possible). Similarly, the plant
362 and arthropod prey levels are frequency-based as it is not possible to assess ingested plant
363 biomass from seeds (plant ID) and metabarcoding data are poorly suited to quantify the
364 biomass or abundance of species represented in the data (Deagle et al., 2018). The
365 arthropods are represented by MOTUs (Floyd et al., 2002). In addition we produced a
366 separate network of bats and prey employing the non-OTU based BIN association matrix.

367

368 **Results**

369 *Molecular Analysis*

370 We recovered DNA barcodes from 698 bats representing 24 species belonging to
371 17 genera. The barcode results generally confirmed the field IDs, but could not
372 distinguish *Artibeus lituratus* and *Artibeus intermedius* (Clare, Lim, Fenton, & Hebert,
373 2011) leaving this node unresolved. However, other cases of taxonomic uncertainty were
374 resolved. An unknown species of *Carollia* was identified as *C. sowelli* and members of
375 two genera (*Glossophaga*, *Micronycteris*) gained species assignments. We suspect one
376 genetically divergent specimen of *Sturnira parvidens* may be a sister taxon, but since this
377 outcome could not be confirmed, it was retained as a single node (Figure S1).

378 We recovered DNA barcodes from 445 of the 466 mites and flies found on 18
379 host species. Parasite diversity varied from a single ectoparasite species per bat species (9
380 cases) to nine ectoparasite species on *A. jamaicensis*. Among the seven bat families, the
381 Emballonuridae, Mormoopidae and Phyllostomidae hosted the greatest diversity of
382 parasite species whereas individual of the Noctilionidae were only associated with two
383 ectoparasite species and individual Molossidae, Natalidae and Vespertilionidae were only
384 parasitized by one ectoparasite species at a time. Two thirds of the 34 ectoparasite
385 species, nine mites and 13 flies were only collected from one host species. The maximum
386 number of host species inhabited by a mite or fly species was four.

387 We recovered plant DNA from 112 guano samples from 12 species of bat. Guano
388 from seven bats contained two seed morphotypes analysed separately, producing 119
389 sequenced seed samples. We recovered *rbcL* from 102, *matK* from 81 and *trnH-psbA*

390 from 106 samples. Through comparison to GenBank and BOLD, 103 samples had
391 sequences assigned to eight genera based on at least two of the three loci. Of these, 97
392 seed samples had sequences assigned to a putative species and 16 samples had sequences
393 placed to a genus based on a single gene region (Table S4). Comparison of *rbcL*
394 sequences to GenBank often returned multiple BLAST hits with equivalent best scores.
395 For example, top BLAST matches to *Ficus* or *Solanum* matched (100% or 99% identity
396 respectively) multiple species within these genera. Although some sequences did not
397 have an identical match on GenBank, all *rbcL* sequences matched with 100% identity to a
398 sequence on BOLD, presumably reflecting the greater diversity of taxa present in the
399 latter database. Similarly, all *matK* sequences matched with 100% identity to sequences
400 on BOLD versus lower values on GenBank (94-100%). In some cases this allowed a
401 more precise taxonomic assignment on BOLD, either to a species (e.g., *Guazuma*
402 *ulmifolia*), or species cluster (e.g., *Cecropia obtusifolia* / *peltata* / *insignis*) versus
403 assignment to a higher taxonomic rank (e.g., Urticaceae or *Cecropia* sp.; *Ficus* sp.).

404 The GenBank BLAST of *trnH-psbA* sequences corroborated results obtained with
405 *rbcL* and *matK*. In all cases, samples that yielded unique sequences for *matK* also had
406 unique sequences for *trnH-psbA*. Although the limited taxonomic coverage for the latter
407 gene region on GenBank often prevented an assignment to a known species, these
408 sequence variants were treated as putative species. We also detected a probable case of
409 taxonomic error in GenBank. Two *trnH-psbA* sequences from our samples showed high
410 similarity (98% identity) to *Cecropia obtusifolia*, an unexpected result as several other
411 sequences of almost twice the length showed nearly 100% similarity to several other

412 species of *Cecropia* on GenBank. Further investigation revealed that these sequences
413 likely belong to a species of *Vismia* (Hypericaceae).

414 We used HTS to recover two regions of the mitochondrial COI gene (157bp, and
415 407bp, Tables S2-S3). The 157bp region has been used extensively (Alberdi et al., 2018;
416 Zeale et al., 2011), and generated high recovery rates in the present study; it is fully
417 analysed and generated 686 MOTU at the given parameters. Surprisingly, given the
418 degradation induced by digestion, the 407bp region also showed high sequencing success.
419 These two regions (Tables S5 and S6) identified a similar number of species (118 versus
420 109 taxa for the 157bp and 407bp regions respectively) from all the same classes and
421 orders of arthropods (excepting one mantid). Many of the same species, for example, 32
422 species of Lepidoptera, were common in the two lists. However, there were also different
423 species identified and in a number of cases identifications were improved using the
424 longer target region. For example, sequences assigned to the genus *Culex* by the short
425 region could be identified as *Culex nigripalpus* by the 407bp region. This outcome
426 suggests these two regions may be complementary, adding confidence to the general
427 diversity recovered and the specific taxa identified. However, the 407bp region pushes
428 the current limits of amplicon size recovery on most HTS platforms, creating constraints
429 on quality and recovery rate. Analysis with the BIN association method in mBRAVE
430 identified 212 potential prey in the 157bp dataset.

431

432 *The impact of OTUs on network metrics*

433 The most novel data type generated is the metabarcoded data that underlie the bat-
434 prey network because the prey nodes do not represent a particular taxonomic level or

435 taxon but a measure of prey genetic diversity. As a result, we investigated the impact of
436 the key bioinformatics step – that of generating MOTUs – on the measurement of
437 common network variables. Our data suggests that MOTU thresholds have a significant
438 impact on standard network metrics as taxa are lumped or split to a greater or lesser
439 extent. For most network metrics (Figure 4), an increase in the MOTU threshold (e.g.,
440 from 90% to 99%) split taxa so the resource level in our network increased in richness
441 relative to the consumers with expected outcomes for each metric. In the case of links
442 between species, connectance, nestedness, and vulnerability this variation can result in
443 different relative rankings of these metrics between network types. For a complete
444 analysis see Hemprich-Bennett et al., (2018). The effect is consistent but less predictable
445 in measures of robustness (Figure 4), but in all cases we would have drawn the same
446 conclusion. The BIN association network (Figure 5) contained substantially fewer prey
447 nodes, which is to be expected, as the reference database for the area is minimal. Of
448 these, 75% were Lepidoptera reflecting the substantial effort to create a Lepidoptera
449 reference library for the site (see below). Interestingly, the actual measurements of
450 network properties did not differ substantially (Table 1) which reflects the tremendous
451 prey diversity represented by any method.

452

453 **Discussion**

454 We have demonstrated that three types of molecular data can be incorporated into
455 network analysis. DNA can be used to confirm field identities (e.g. bats) or differentiate
456 cryptic taxa (e.g. parasites) and to identify morphologically compromised material (e.g.
457 plant pulp). DNA can also be used to generate complex and fundamentally novel data via

458 metabarcoding of mixed material (e.g. faeces) that can be analysed using MOTU or
459 association with taxa in reference collections (e.g. the BIN method). While these data
460 types can effectively generate rapid, scalable analyses of entire communities, there are
461 challenges in both generating data and in the interpretation of network metrics to ensure
462 biologically meaningful results (Table 2, Figure 4).

463 The incorporation of DNA analysis into networks presents both straightforward
464 use cases and challenges. Confirming field IDs is a common molecular procedure
465 (Borisenko, Lim, Ivanova, Hanner, & Hebert, 2008) and differentiating cryptic or
466 taxonomically complex species is now routine (Smith, Woodley, Janzen, Hallwachs, &
467 Hebert, 2006). These approaches have successfully been incorporated into network
468 analysis (e.g., Wirta et al., 2014). However, the inclusion of metabarcoding results is
469 more challenging and requires special consideration to integrate with network analysis.
470 Metabarcoding is best applied to mixed faecal samples, gut contents (particularly liquid
471 feeders e.g. Piñol, San Andrés, Clare, Mir, & Symondson, 2014) or pollen carried by
472 generalists. However, it is challenging to generate reliable metabarcoded data (Alberdi et
473 al., 2018; Arrizabalaga-Escudero et al., 2018; Atwell et al., 2010; King, Read, Traugott,
474 & Symondson, 2008). The methods of interpreting individual dietary analyses using these
475 data have been studied in several contexts (Clare et al., 2016; Flynn et al., 2015;
476 Pompanon et al., 2012; Symondson, 2002). However, certain challenges are unique to the
477 interpretation of food webs. Debate about the quantification of metabarcoding data
478 centres largely around whether sequence recovery is linked to original biomass (Deagle et
479 al., 2013; Nielsen et al., 2018; Pompanon et al., 2012). While this is possible in restricted
480 scenarios (Bowles, Schulte, Tollit, Deagle, & Trites, 2011; Thomas et al., 2016), in many

481 cases frequency-based measures of interactions are more appropriate (Nielsen et al.,
482 2018). Frequency-based approaches are common in network ecology, for example,
483 visitation frequency to specific flowers is a standard metric of the strength of mutualistic
484 interactions (Memmott, Waser, & Price, 2004; Vázquez, Morris, & Jordano, 2005).
485 However, incomplete quantification needs to be considered when weighted metrics are
486 used (Kaiser-Bunbury, Muff, Memmott, Müller, & Caflisch, 2010), as rare and common
487 interactions may be equally weighted (Clare, 2014)

488 We suggest two alternative ways to incorporate metabarcoding data: using
489 MOTUs and screening for taxonomic identities (e.g. BINs, similarity searches). The
490 advantage of MOTUs is that all data are incorporated, both known and unknown taxa.
491 However, by incorporating unknowns, one may inadvertently include non-target taxa
492 (e.g. intestinal parasites or bacteria that are not screened out bioinformatically) that may
493 generate nodes in networks unrelated to the behaviour under study or even false nodes
494 from sequencing error (Clare et al., 2016; Flynn et al., 2015). In many systems, MOTUs
495 collapse all prey levels into one “resource” level rather than revealing the complexity
496 among trophic levels. For example, in our case some insects were primary consumers
497 while others were predators, but all were treated as MOTU “prey” of an undifferentiated
498 consumer level. Our data further suggest that the protocols used to differentiate MOTUs
499 will themselves impact network metrics (Figure 4). The effect of node resolution has
500 been discussed for decades with analyses showing that the impact of resolution on node,
501 chain length and trophic levels significantly alters the observations of network properties
502 (Brose, Ostling, Harrison, & Martinez, 2004). The situation is similar but not identical to
503 the node resolution issue of employing MOTU. The impacts of the informatic steps used

504 to generate MOTU are only starting to be considered in ecological (Clare et al., 2016) or
505 network analysis (Hemprich-Bennett et al., 2018). Any network that incorporates taxa
506 with different levels of resolution (e.g. mixing genus and species designations) faces the
507 same challenge (Hemprich-Bennett et al., 2018; Martinez, 1991). However MOTUs make
508 it possible to easily re-analyse any dataset to empirically estimate that impact (Figure 4)
509 and one potential advantage is that MOTUs generate a uniform level of resolution in a
510 network. By their nature MOTUs represent equal and repeatable measures of biodiversity
511 (Floyd et al., 2002), even if that level does not equate to a standard level of taxonomy.
512 This may represent a powerful advantage in comparing network structure across systems,
513 but presents a challenge in interpretation. For example unnamed MOTUs of unknown
514 life-cycle and unknown affinity to each other provide limited information on the nature of
515 the ecological interaction being measured beyond the general structure of the community.
516 Similarly, if the numbers of nodes and their connections vary with analysis parameters
517 (e.g. MOTU threshold) a network on its own holds little biological meaning. However, if
518 the same methods are replicated a biological picture can emerge. For example, if the prey
519 level undergoes a population crash, the genetic diversity and the MOTU number would
520 similarly decline relative to the consumer level and fluctuations in parameters such as
521 generality or nestedness would be measureable. The key then is to compare only analyses
522 that employ the same methods from sequencing platform and field and lab protocol to
523 informatics choices, just as sampling protocols and node resolution should be maintained
524 in traditional networks being compared. This would be required to avoid context
525 specificity. It is also necessary to pick specific metrics; for example network level metrics

526 may be more reliable than motif measurements (though see Hemprich-Bennett et al.,
527 2018).

528 In contrast, similarity based searchers and BIN association type data provide better
529 ability to determine exactly what is being included as a node (e.g. Figure 5) and yield
530 greater ecological information about the type of interaction being measured, but will be
531 biased by the contents of the reference library being used. In this case, the network
532 metrics were similar enough that comparative conclusions about bat-prey/BIN, bat-
533 parasite and bat-plant networks would remain the same but some specific indicators
534 change. For example, generality of the bat-BIN network was much lower reflecting the
535 substantial reduction in prey nodes when relying exclusively on reference collections for
536 the inclusion of a prey node. As reference collections improve this effect will diminish
537 but it is a very important factor in relatively unexplored faunas.

538

539 *Three distinct data types*

540 This paper has considered three distinct types of molecular data. The bat and plant
541 identifications provided by DNA deliver nearly perfect resolution of the network. Such
542 analysis generates data similar to that employed in traditional network ecology, the only
543 major difference being the need for multi-locus data to obtain species-level resolution for
544 plants. The parasite identifications were generated in a similar fashion to the bat data (one
545 sequence per specimen), but with the crucial difference that current reference databases
546 are very incomplete. As a consequence, we employed an alternative taxonomic system,
547 the Barcode Index Number (BIN). The performance of the BIN system has been
548 extensively tested (Ratnasingham & Hebert, 2013) and these studies have shown that it

549 delivers taxonomic resolution that is very close to traditional taxonomy. This data type
550 (Table 2) has the advantage of making it possible to incorporate taxa which are
551 apparently different species but where the current taxonomic system is incomplete. BIN
552 analysis avoids unresolved nodes in network construction, but imposes a constraint that
553 the identification is based on a measure of sequence differentiation observed in related
554 taxa. Unlike other MOTU-generating methods, the BIN system is not based on strict *a*
555 *prior* threshold delimitation but has been trained specifically using the large Sanger data
556 sets for the DNA barcode region. In a test of 1400 species spanning birds, bees, fishes
557 and Lepidoptera, the correspondence between species counts based on traditional
558 taxonomy and BINs was very high ($r^2=0.99$) and the actual mapping of species to BIN
559 was approximately 90% (varying from 79%-97% between taxonomic group)
560 (Ratnasingham & Hebert, 2013). Thus, when viewed from the context of DNA
561 barcoding, BINs are a strong proxy for species. Because the definition of new BINs
562 requires at least 500bp of sequence information from the COI barcode region, the short
563 reads generated by most current HTS platforms cannot be used to delineate new BINs
564 although they can be matched to existing BINs. Reflecting this constraint, there is a need
565 for other methods of MOTU generation. For this third data type, data are most often
566 analysed using MOTU without much (if any) taxonomic identification (e.g. Figure 1).
567 This has the advantage of making it possible to analyzed mixed sources (e.g. stomach
568 contents) but imposes unique problems for network ecology as it compresses trophic
569 levels and dispenses with traditional taxonomy. While such analysis can generate data for
570 a comparable interaction network model, it may not represent a trophic food web. In the
571 study location, most arthropods remain undescribed despite decades of intense taxonomic

572 work. For example, some 400,000 arthropod species are estimated to be present in the
573 ACG, but just 43,000 of these species have been barcoded over 14 years, revealing the
574 scope of the taxonomic challenge (D.H. Janzen Pers. Comm.). In such locations, a
575 complete food web or interaction network is impossible and restricting analysis to those
576 species which can gain a full taxonomic designation (either by morphology or DNA)
577 would introduce a substantial bias (Table S6). As a consequence our bat-BIN network
578 contained substantially fewer nodes than our bat-prey network based on MOTU and
579 would be less comparable to a network generated in an area with a different/reduced
580 reference database. In such cases, a BIN or MOTU approach to generating a reference
581 collection and then some sort of association or matching system is the only means of
582 developing an ecosystem network model. The use of reference databases can to provide a
583 familiar binomial designation on some nodes by similarity searchers or BIN association
584 but imposes a significant bias on the data, which is then composed of “things found in
585 databases” while novel BINs, and MOTUs do not impose this bias. On the other hand,
586 novel BINs and MOTUs may include non-target taxa such as parasites, parasitoids, or
587 taxa acquired via secondary predation.

588

589 ***Comparison of 157 and 407bp datasets:***

590 Current consensus suggests that short reads are required to maximize MOTU
591 and taxonomic ID recovery in digested material because of DNA degradation.
592 Contrary to this expectation, the 407bp region had higher MOTU estimates and
593 broader taxonomic coverage when evaluated using BLAST, suggesting it has less
594 amplification bias and hence a complementary region for arthropod diversity

595 analysis. However, this conclusions need to be considered with caution. Longer
596 reads should generate better taxon identification scores (more information) but will
597 also generate high rates of low quality BLAST scores (local alignments). We
598 modified the MEGAN Min Support value to partially compensate for this and to
599 maximize assignment with most scores >98% similar to references. Taxonomic
600 assignments of MOTUs (e.g. Table S5 and S6) should be considered for interest's
601 sake only in this dataset, particularly when a reference databases contains errors or
602 skewed coverage. For this reason we considered only MOTUs for network analysis.
603 Despite the promise of the 407bp region we used the 157bp region MOTUs for
604 network analysis for two reasons. First, unlike the 157bp region, the 407bp region is
605 new to NGS analysis and has not been evaluated for this purpose before. We
606 consider it an interesting and potentially important tool but are hesitant to rely on it
607 until further testing has been completed. Second, the 407bp region is long compared
608 to the capacity of most high throughput sequencing platforms which limits its use
609 and prevented us from employing MIDx to separate samples. Platform read length
610 has generally fallen since the first highly popular Roche454 platform capable of
611 1000bp reads to the now standard 250bp paired end reads of the MiSeq, thus while
612 promising, the 407bp read will be analytically challenging. Newer platforms such as
613 the SMRT sequencing platform (PACBIO, Pacific Biosciences) can overcome this
614 problem allowing longer reads and thus higher taxonomic resolution assuming that
615 digestion has not substantially compromised the DNA.

616

617 *DNA integration into network ecology.*

618 Despite challenges, incorporating DNA into networks has significant advantages.
619 First, the technique does not rely on the need to observe interactions or the time
620 consuming rearing practices used to establish many cases of parasitism (Wirta et al.,
621 2014). It can be applied to broken seeds, fruit pulp (e.g. Lim et al., 2017), single grains of
622 pollen or morphologically destroyed material (e.g., digested remains) as well as entire
623 specimens. Even traces of DNA (eDNA) with no observable material are amenable
624 (Bohmann et al., 2014; Drinkwater, Clare, & Rossiter, In Review); for example, seeds
625 dropped on the forest floor will have DNA of the plant, but also of the animal that
626 dropped them from either saliva or cells from a digestive tract. González-Varo et al.,
627 (2014) have spectacularly demonstrated this method to capture bird DNA on the surface
628 of dispersed seeds. Similarly, the detection of cryptic species and relationships represents
629 a huge shift in the resolution of interaction networks. This was demonstrated by Wirta et
630 al., (2014) who observed that DNA dramatically increased the number of identified
631 interaction types and altered the perceived host specificity of host-parasitoid networks.

632 A rapid DNA-based network biomonitoring tool will require us to understand: first,
633 which data can be quantified (Deagle et al., 2018) and second, which metrics are reliable,
634 in relative or absolute terms, to ensure we produce biologically meaningful outputs
635 (Hemprich-Bennett et al., 2018, Clare et al. 2016, Ings et al. 2009,). However, these
636 datasets are already being demonstrated as powerful tools to resolve complex interaction
637 networks quickly and in exquisite detail. Here we have generated a detailed network of
638 networks in a complex tropical ecosystem incorporating different molecular data types as
639 a case study. Ecologically, our data suggest a hitherto unrecognised keystone species and
640 behavioural flexibility that may be critical to the success of insectivores (Box 1).

641 Methodologically, our findings support the approach, but also highlight the need for
642 rigorous testing of methods. The rapidly advancing technology of this field means that
643 such analyses will soon become a common and relatively inexpensive tool for
644 understanding biostructure (McCann, 2007). While a fully resolved and taxonomically
645 identified network will always be the goal, our analysis demonstrates the utility of these
646 tools for network ecology and produces the first full network of networks resolved
647 entirely by DNA.

648

649 **Acknowledgments:** We are indebted to collaborators who have supported this research
650 including staff at the Canadian Centre for DNA Barcoding and the Biodiversity Institute
651 of Ontario and the staff of Sector Santa Rosa in the Area de Conservación Guanacaste
652 (ACG) Costa Rica particularly Daniel Janzen, Winnie Hallwachs, Roger Blanco and
653 Maria Marta Chavarria.

654

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656

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910 Box 1: A preliminary analysis of a tropical bat community.

911

912 Field Methods: All materials were acquired from past research at the field location
913 and/or held in personal collections. All bats were caught over a six-week period
914 from late May – early July of 2009 using mistnets or harptraps in Sector Santa Rosa
915 of the Area de Conservación Guanacaste. Net locations were alternated nightly
916 between the Bosque Humeda, La Casona and the Picnic area with an extra netting
917 night at the Playa Naranjo targeting *Noctilio*. Each bat was identified and placed in a
918 cloth bag for approximately one hour. Any guano produced was collected for
919 taxonomic identification of prey items and the bats were released at the point of
920 capture. Ectoparasites and wing biopsies were preserved in isopropyl alcohol;
921 guano was frozen. Morphological identification of the ectoparasites to fly or mite
922 was made in order to separate functional groups. Guano samples were screened for
923 seed fragments and insect remains and classified as containing plant material or
924 insect material. Two genera, *Glossophaga* and *Micronycteris*, were left with
925 provisional species level ID. *Artibeus lituratus* and *A. intermedius*, could not be
926 distinguished in the field and are referred to as *A. sp.*

927

928 Network Analysis: We visualised the interaction networks using Bipartite (Dormann
929 et al., 2009) and Cheddar (Hudson et al., 2013) as implemented in R (R Development
930 Core Team, 2015). We compared the structural metrics (links per species,
931 asymmetry, connectance, nestedness, generality, and vulnerability) of each
932 traditional bipartite sub-network (bat-parasite, bat-plant, bat-insect). We evaluated
933 the robustness of each network and modelled the effects of species loss and
934 restoration within the networks. We employed three extinction models: species
935 removed randomly (null model), species removed from most to least connected (Rd-
936 worst case scenario) (Kaiser-Bunbury, Muff, Memmott, Müller, & Caflisch, 2010) and
937 species removed from least to most frequently detected (Ra-best case scenario).
938 Species lose connections within the network when their hosts, prey, predators, or
939 mutualists are eliminated. From each of our three component networks (parasitism,
940 mutualism, predation) we measured network robustness (Kaiser-Bunbury et al.,
941 2010; Memmott, Waser, & Price, 2004). We then modelled a restoration scenario
942 where bat species are re-introduced from greatest to least connected (best-case)
943 and assessed the proportion of links restored to the structure. To pinpoint possible
944 keystone species, we examined the role of each bat species within the entire
945 network of network using betweenness and closeness centrality scores (Martín
946 González, Dalsgaard, & Olesen, 2010) in igraph (Csardi and Nepusz 2006). For
947 simplicity, when individual networks are depicted, we present bats on the top rather
948 than arranging these by trophic level (bats occupy multiple trophic levels making
949 any other display exceedingly complex). Finally, we analysed the impact of OTU
950 clustering thresholds of insects in the bat-prey network on the measurement of
951 these metrics considering clustering thresholds from 90-99%.

952

953 A network of networks: Using these data, we present the first “network of networks”
954 where all underlying data have been generated using a molecular approach (Figure
955 1a). We evaluated the structural metrics (Table 1) and robustness (Figure 2) of

956 traditional sub-networks (Figure 1bcd) and modelled the effects of bat species loss
957 on parasite persistence (Figure 2a), plant mutualism (Figure 2b), predation (Figure
958 2c), and secondary extinction of bats from prey loss (Figure 2d). Under all models,
959 parasite networks were less robust ($R_a=0.69/R_d=0.36$) and mutualistic
960 relationships were only slightly more robust ($R_a=0.74/R_d=0.4$). However,
961 arthropods responded differently to models of extinction: a high proportion of prey
962 face predation, even when the highest-ranking bat species by abundance are
963 eliminated ($R_a=0.86$), while, conversely, arthropods experience a tremendous
964 release from predation when bat species go extinct by connectance ($R_d=0.28$).
965 Insectivorous bats appear robust to the loss of prey species ($R_a=0.998/R_d=0.85$).
966 Even under the worst-case scenario, the first bat species is not lost until 32% of
967 arthropods are extinct, and even when >90% of arthropods are lost, >70% of bat
968 species remain in the network if prey biomass was sufficient (Figure 2d). Only *G.*
969 *soricina* showed significant trophic flexibility operating in both a mutualistic and
970 strong predatory role (high centrality scores, Supplemental Information Table S8).
971 This is also evident in our restoration ecology model (Figure 3) where the third bat
972 returned based on connectance is *G. soricina* introducing parasites, insects, and
973 plants at the same time.

974
975 A snap shot of a bat community: Even considering the variability of metrics across
976 multiple MOTU resolutions (Figure 4), the generality of bat-prey networks is
977 extreme compared to the bat-parasite and bat-plant networks. This significantly
978 impacts on our understanding of robustness in this system and may provide
979 evidence in the diversity vs. stability debate (McCann, 2000). The data suggest
980 extraordinary behavioural flexibility of insectivorous bats and their lack of reliance
981 on specific prey. While there is evidence for resource specialisation (e.g. the
982 preference for beetles in *Eptesicus* (Clare, Symondson, & Fenton, 2014) or moths in
983 sibling rhinolophids (Arrizabalaga-Escudero et al., 2018)) most studies that have
984 employed molecular techniques have observed very generalist flexible behaviour in
985 foraging (Salinas-Ramos, Herrera Montalvo, León-Regagnon, Arrizabalaga-Escudero,
986 & Clare, 2015; Sedlock, Krüger, & Clare, 2014) though none have examined a
987 community on this scale. Second, perhaps the most interesting observation is the
988 position of the bat *Glossophaga soricina* in the network. Clare, Goerlitz, et al., (2013)
989 used a molecular dietary analysis to identify a novel hunting strategy that permits
990 this supposed “nectar bat” to sneak up on insects. The bats’ echolocation calls are
991 low enough in intensity that prey with ears do not detect the approaching threat in
992 time to evade it (Clare, Goerlitz, et al., 2013). Our network analysis suggests that
993 insectivory in *Glossophaga* is not a rare behaviour but rather, during the period of
994 this study, *G. soricina* was the third best-connected insectivore in the community in
995 addition to its role in pollination and, seed dispersal and as a parasite host. Its
996 diverse functional roles make it a probable keystone species with very high
997 betweenness and closeness centrality (Supplemental Information Table S8). This
998 distinguishes it as the only bat occupying all these functional roles in the network
999 and thus a species of special conservation interest. In contrast some species are only
1000 very tangentially associated with this network. For example *Desmodus rotundus*, the

1001 vampire bat, is connected to only one parasite and thus forms its own module of
1002 interactions unconnected to the rest of the community.
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1005 **Data Accessibility:** All molecular data can be found in Dryad
1006 <https://doi.org/10.5061/dryad.0k90c0v> and BOLD projects (BCCR Bats of Costa Rica
1007 ACG & BCPB Parasites of tropical bats) also contain sequences and collection metadata
1008 and associated GenBank accessions.

1009

1010

1011 **Author Contributions.** ELC, AMA, and JN performed fieldwork. ELC, AJF, NVI and
1012 RMF performed laboratory work. ELC and RG performed analysis. PDNH, MBF and
1013 SGN helped design the study. All authors contributed to the writing of the manuscript.

1014

1015 Tables:

1016

1017 Table 1: Structure of the sub-networks

	Links per species	Asymmetry (A)	Connectance (C)	Nestedness (N)	Generality (G)Ψ	Vulnerability (V) Ψ
Bat-Parasite	1.02	-0.31*	0.09	12.60	1.95*	1.32*
Bat-Plant	1.37	-0.33	0.21	26.25	2.74	1.80
Bat-Prey	1.70	-0.97	0.16	14.11	76.72	1.52
Bat-BIN	1.17	-0.91	0.12	14.79	41.25	1.37

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1019 *see Supplemental Information for an interpretation of positive vs. negative values and structural arrangement

1020 Ψ unweight following (11) but see Supplemental Information for the appropriateness of unweight measures

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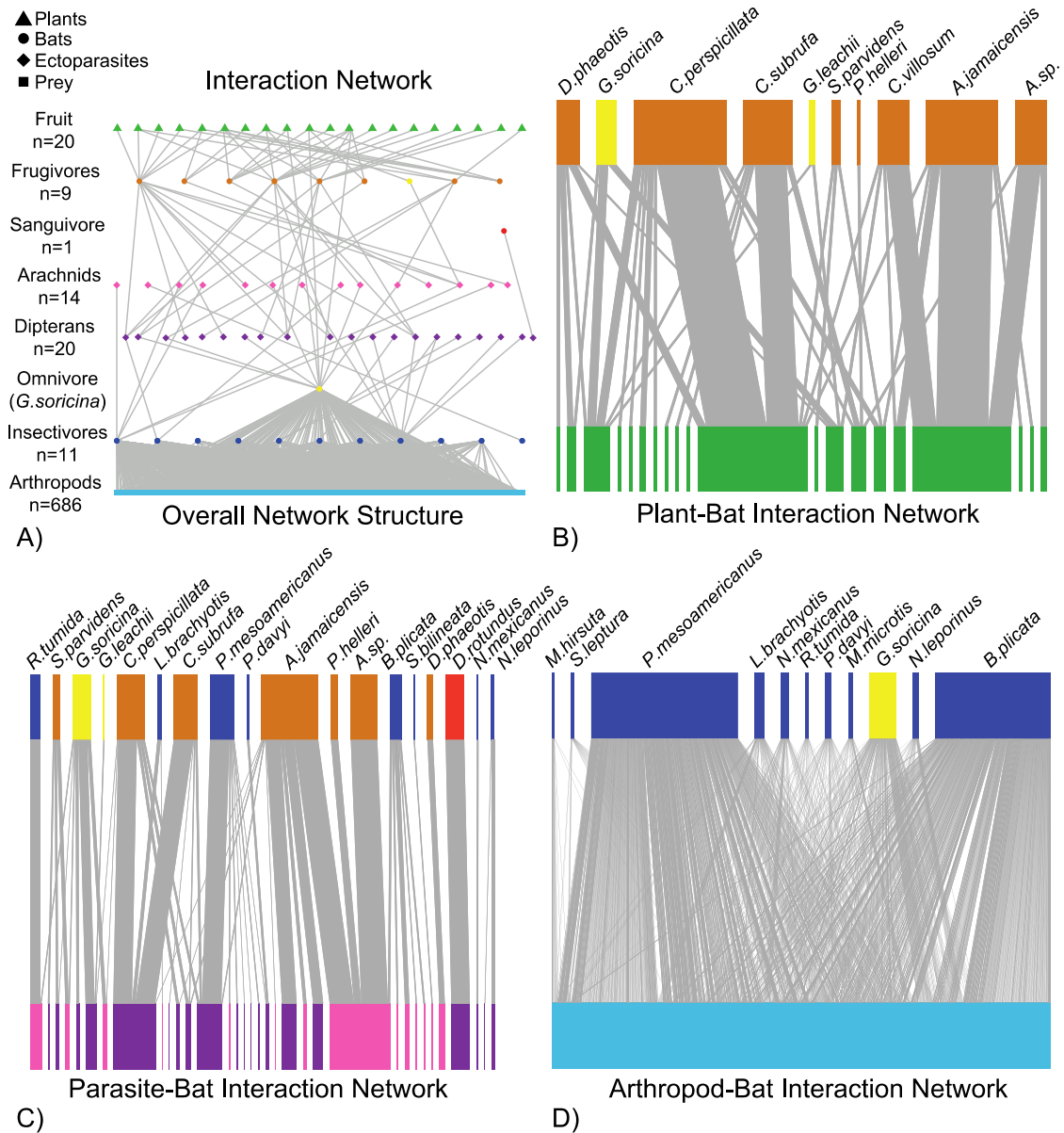
1024 Table 2: A comparison of three data types for network analysis

	Data Type	Taxonomic Resolution	Application	Required References	Advantages	Disadvantages	Network Implications
Resolving ID with DNA -The plant network	A few sequences per individual	Species	Identification of fragments	Complete database	-can deal with forensic trace material -produces traditional networks -minimal taxonomic expertise	- requires molecular expertise - requires a well resolved database	-Produces a network equivalent to a traditional food web
Resolving species when taxonomy is not known - The parasite network	A few sequences per individual	Species but without names	Identification of taxa where taxonomy may be incomplete or cryptic	Incomplete database	-can deal with forensic trace material -can include species with incomplete taxonomic investigation - resolves cryptic taxa	requires molecular expertise -may not encourage taxonomic work	-Similar to a traditional food web, but may collapse trophic levels where ID is not clear
Using MOTUs without a taxonomic unit - The arthropod prey network	Millions of sequences per sample	Arbitrary but comparable units	Rapid surveys where identification is not possible	No database required	-can deal with forensic trace material -may include both known and unknown data (MOTUs) -can be used in any context regardless of taxonomic knowledge -rapid and MOTUs are mathematically and genetically identical yielding perfect resolution -quantification is controversial*	- requires molecular expertise -does not represent real taxa -may include error- prone data -actual MOTU numbers are meaningless -may be biased by primers or other protocol choices	-Fundamentally different MOTUs are not species and likely collapse trophic levels but allow rapid structural comparisons -Node numbers are meaningless

1025 * See a review by Deagle et al. 2018.

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Figures:



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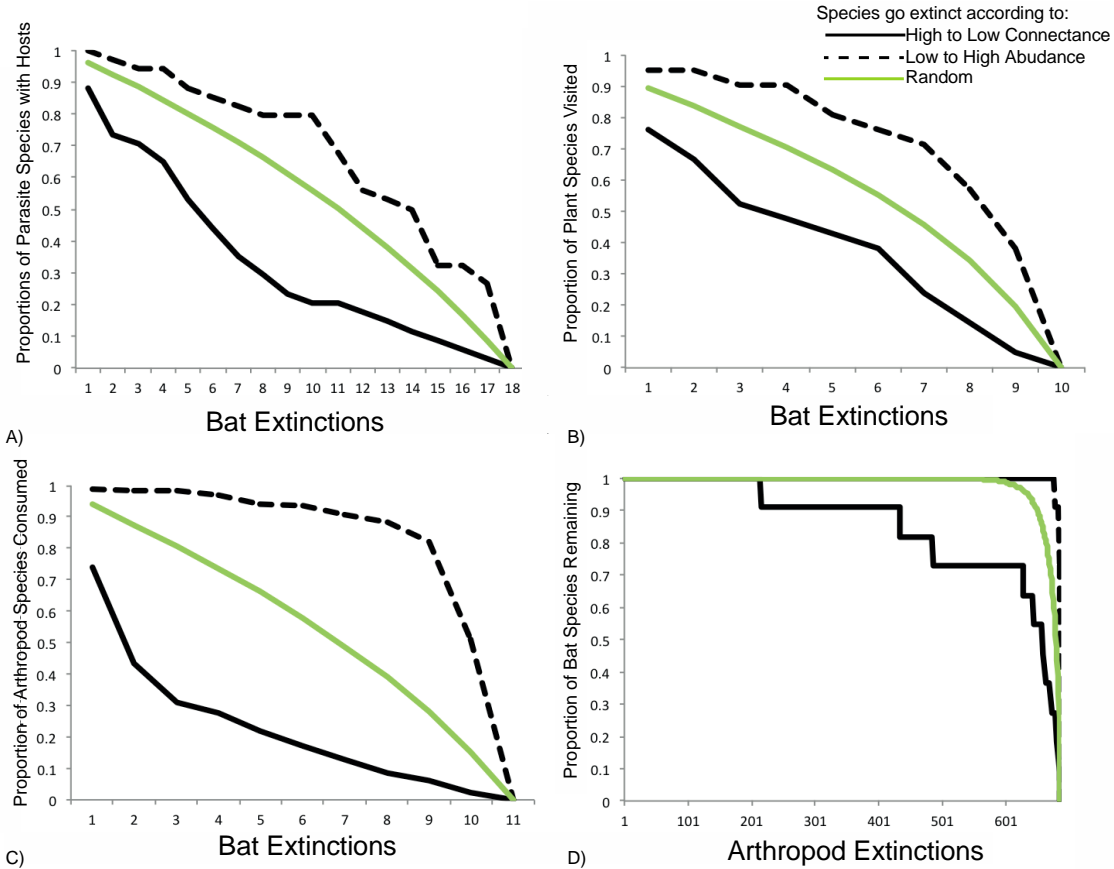
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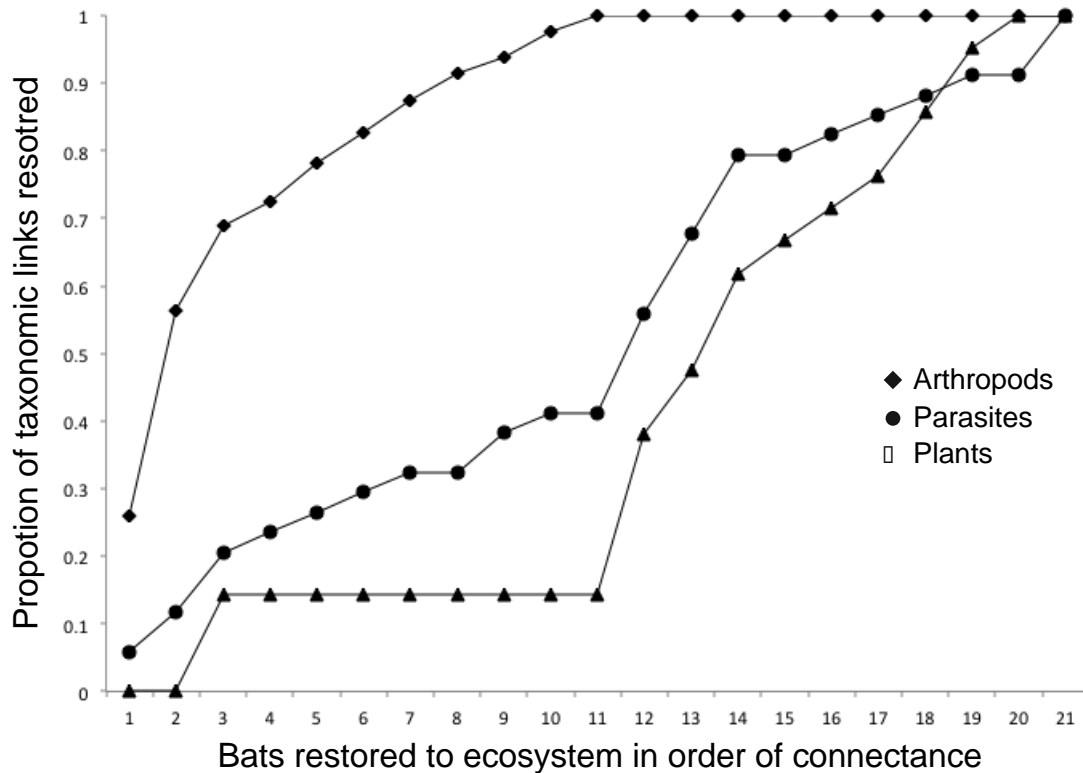
Figure 1: Species' interaction networks. The network of networks (A) displays interaction structure organised by behavioural ecology (rather than traditional trophic structure). The visualization of this network is not presented as standard trophic levels for two reasons. First, the arthropod prey represent multiple trophic levels themselves which cannot be differentiated. Second, the density of connections make links to plants impossible to distinguish if the plants are presented as the lowest trophic level. In this case the network has been structured to depict function rather than trophic levels. For example, arachnid mites of bats are parasites that spend their entire life cycle on their host (Christe, Arlettaz, & Vogel, 2000) which restricts their dispersal so horizontal transmission primarily occurs via host-to-host contact. Therefore, mites and their hosts are usually regarded as the product of long co-evolution. In comparison, parasitic Diptera (flies) can be highly

1040 mobile, and often spend part of their life cycle apart from their host (Fritz, 1983).
1041 Because of such distinct life history differences alternative hypotheses of function can be
1042 advanced. We depict them as separate functional groups (A) and in their traditional
1043 parasite role (C). N-values represent the number of taxa detected. Semi-quantified
1044 individual trophic networks (B-D) display traditional trophic organisations (though for
1045 simplicity of comparison bats are always presented on top). Detection frequency data for
1046 each species is given by the width of the block proportional to species' frequency in the
1047 network. Colours indicate behavioural role from A. See Supplemental Information for a
1048 discussion of visualization orientation and Supplemental tables S4 and S8 for matrices of
1049 parasite, plant and bat taxonomic identifications.



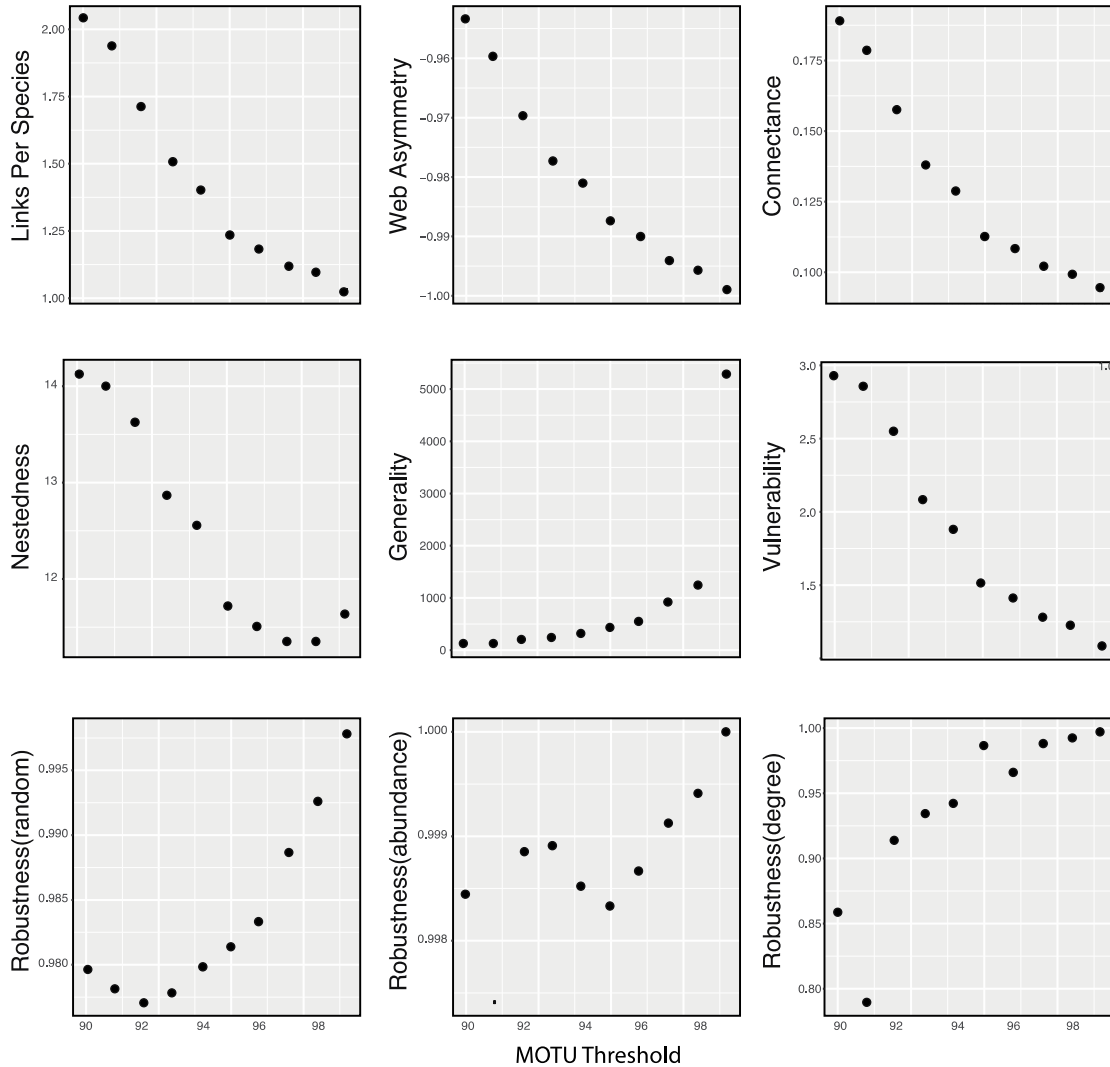
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Figure 2: The robustness of interaction network structure to the sequential removal of species under three extinction models. The number of bat species removed has an extreme impact on the loss of parasites (A), while plants are slightly more resilient (B). The proportion of arthropods released from predation (C) is strongly dependent on the model of extinction, while insectivorous bats are extremely resilient to the loss of prey under any model (D).



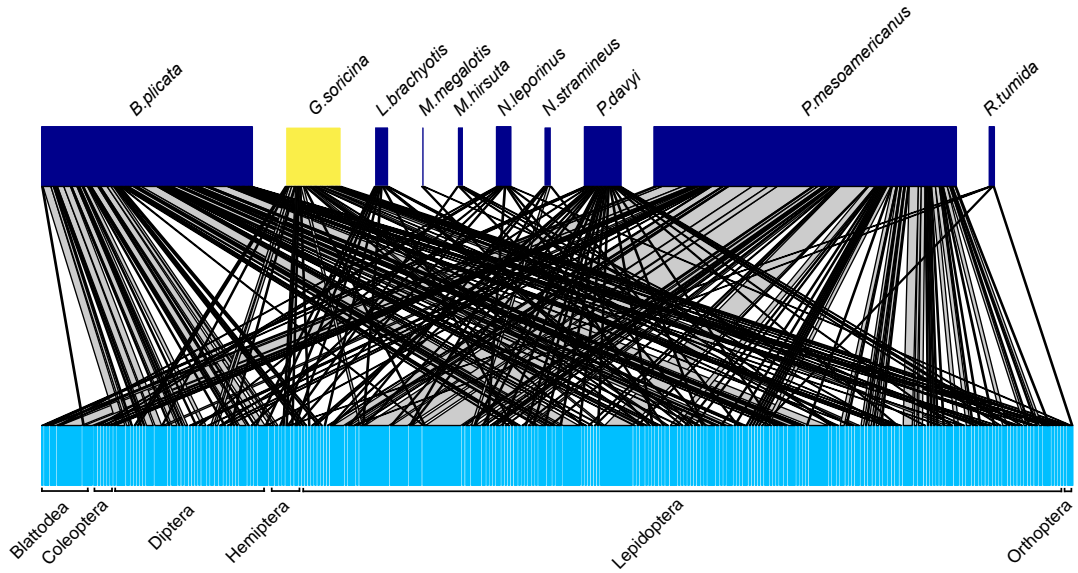
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Figure 3: A restoration ecology model showing the proportion of links restored if bats are introduced to the ecosystem in order of connectance (best case scenario). With the restoration of only the three most strongly connected species (*Pteronotus mesoamericanus*, *Balantiopteryx plicata* and *Glossophaga soricina*), 72% of arthropod species are under predation, 24% of parasite species have a host, and 14% of plant species are visited. See Clare et al., (2014) for a discussion of trophic roles of *Glossophaga*.



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Figure 4: Metabarcoding data is a fundamentally new type of data for network ecology. Nodes in metabarcoding normally do not represent a specific taxon or taxonomic level, but are molecular operational taxonomic units (MOTUs) best described as taxa that are defined by being genetically congruent pools of diversity. They are defined by a series of bioinformatics steps with the ultimate decision dependent on the threshold employed for splitting vs. lumping sequences into a MOTU (nodes in our networks). As the MOTU threshold changes, taxa are lumped or split to a greater or lesser extent. For most network metrics (top two rows), this has a predictable effect as the resource level in our networks increases in richness relative to the consumers. The same pattern is evident but less predictable in measures of robustness (bottom row). For a complete analysis see Hemprich-Bennett et al. (2018).



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Figure 5: Species' interaction networks for bats and prey identified using the BIN association method employed on the mBRAVE platform. See table S9 for a matrix of bats and BIN based nodes with full taxonomic identifications.