

Biodiversity and vector-borne diseases: Host dilution and vector amplification occur simultaneously for Amazonian leishmaniases

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1 2	Biodiversity and vector-borne diseases: host dilution and vector amplification occur simultaneously for Amazonian leishmaniases
3	Running title: Biodiversity and Amazonian leishmaniases.
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24 Abstract

25 Changes in biodiversity may impact infectious disease transmission through multiple mechanisms. We explored the impact of biodiversity changes on the transmission of 26 27 Amazonian leishmaniases, a group of wild zoonoses transmitted by phlebotomine sand flies 28 (Psychodidae), which represent an important health burden in a region where biodiversity is 29 both rich and threatened. Using molecular analyses of sand fly pools and blood-fed dipterans. 30 we characterized the disease system in forest sites in French Guiana undergoing different levels 31 of human-induced disturbance. We show that the prevalence of *Leishmania* parasites in sand 32 flies correlates positively with the relative abundance of mammal species known as Leishmania reservoirs. In addition, Leishmania reservoirs tend to dominate in less diverse 33 34 mammal communities, in accordance with the dilution effect hypothesis. This results in a 35 negative relationship between *Leishmania* prevalence and mammal diversity. On the other 36 hand, higher mammal diversity is associated with higher sand fly density, possibly because 37 more diverse mammal communities harbor higher biomass and more abundant feeding 38 resources for sand flies, although more research is needed to identify the factors that shape 39 sand fly communities. As a consequence of these antagonistic effects, decreased mammal diversity comes with an increase of parasite prevalence in sand flies, but has no detectable 40 41 impact on the density of infected sand flies. These results represent additional evidence that 42 biodiversity changes may simultaneously dilute and amplify vector-borne disease transmission 43 through different mechanisms that need to be better understood before drawing generalities on 44 the biodiversity-disease relationship.

45 Keywords: zoonotic disease, dilution effect, metabarcoding, iDNA, phlebotomine sand fly,
46 Culicidae, amplification effect

47 Introduction

48 The current biodiversity crisis alters ecosystem functioning and human well-being through a 49 variety of processes that are still debated (Cardinale et al., 2012; Oliver et al., 2015). In this 50 context, the impact of biodiversity loss on infectious disease transmission and spread has 51 become an important research topic during the last two decades. Particular attention has been 52 given to the potential regulation of pathogens in ecosystems by the presence of species that are 53 inefficient for their transmission, a mechanism referred to as the "dilution effect" (Keesing et 54 al., 2010; Ostfeld & Keesing, 2012). Theoretical investigations as well as laboratory and field 55 experiments have allowed defining the conditions under which a dilution effect is expected to 56 occur (Dobson, 2004; Johnson, Hartson, Larson, & Sutherland, 2008; Johnson, Lund, Hartson, 57 & Yoshino, 2009; Mihaljevic, Joseph, Orlofske, & Paull, 2014; Norman, Bowers, Begon, & 58 Hudson, 1999; Ostfeld & LoGiudice, 2003; Roche, Dobson, Guégan, & Rohani, 2012; Rudolf 59 & Antonovics, 2005; Suzán et al., 2009; Van Buskirk & Ostfeld, 1995), and empirical studies 60 have suggested its existence in numerous disease systems (Clay, Lehmer, Jeor, & Dearing, 61 2009; Derne, Fearnley, Lau, Paynter, & Weinstein, 2011; Ezenwa, Godsey, King, & Guptill, 2006; Gilbert, Norman, Laurenson, Reid, & Hudson, 2001; Gottdenker, Chaves, Calzada, 62 Saldaña, & Carroll, 2012; LoGiudice, Ostfeld, Schmidt, & Keesing, 2003; Ostfeld & Keesing, 63 64 2000; Telfer et al., 2005; Weinstein, Titcomb, Agwanda, Riginos, & Young, 2017). 65 Considering a local community of hosts composed of species that differ in their competence 66 for a given pathogen (i.e., their ability to get infected and transmit), a dilution effect may occur 67 if the presence of the least competent hosts reduces contact rates between the most competent 68 hosts and the pathogen or decreases the density of competent hosts. Additionally, if the less 69 competent hosts are also those that tend to be extirpated from species-depleted communities,

70 biodiversity loss should result in enhanced transmission. The idea that the dilution effect may 71 produce a beneficial impact of biodiversity conservation on public health in most disease 72 systems, has triggered important interest as well as strong criticism (Civitello et al., 2015; 73 Halsey, 2019; Keesing et al., 2010; Lafferty & Wood, 2013; Levi et al., 2016; Randolph & 74 Dobson, 2012; Salkeld, Padgett, & Jones, 2013; Wood et al., 2014). Advances in disease 75 ecology have allowed a more detailed understanding of biodiversity-disease relationships, and 76 the field has progressed beyond initial debates about the generality of the dilution effect. 77 Studies have highlighted the importance of accounting for different factors such as the 78 geographical scale, the nature and extent of biodiversity changes, the transmission mode, and 79 the taxa involved in the disease system under consideration (Cohen et al., 2016; Faust et al., 80 2017; García-Peña et al., 2016; Gibb et al., 2018; Halliday & Rohr, 2019; Halliday, Rohr, & 81 Laine, 2020; Johnson, Calhoun Dana M., Riepe Tawni, McDevitt-Galles Travis, & 82 Koprivnikar Janet, 2019; Johnson, de Roode, & Fenton, 2015; Keesing & Ostfeld, 2021; 83 Morand, Jittapalapong, Suputtamongkol, Abdullah, & Huan, 2014; Rohr et al., 2020; 84 Weinstein et al., 2017; Wood & Lafferty, 2013; H. S. Young, Parker, Gilbert, Sofia Guerra, & 85 Nunn, 2017). It has been shown that changes in biodiversity can either amplify or dilute 86 pathogen transmission, through multiple mechanisms which can sometimes occur within the 87 same system (Clay, Lehmer, St. Jeor, & Dearing, 2009; Faust et al., 2017; Huang, van Langevelde, Prins, & de Boer, 2015; Luis, Kuenzi, & Mills, 2018; Miller & Huppert, 2013; 88 89 Ogden & Tsao, 2009; Roche & Guégan, 2011; Rohr et al., 2015; Swei, Ostfeld, Lane, & Briggs, 90 2011; Wood, Summerside, & Johnson, 2020). In the case of vector-borne diseases, it has been 91 stressed that the ecology and feeding habits of arthropod vectors must also be considered 92 (Carlson, Dyer, Omlin, & Beier, 2009; Hamer et al., 2011; Laporta, Prado, Kraenkel, Coutinho,

93	& Sallum, 2013; Loss et al., 2009; McGregor et al., 2018; Miller & Huppert, 2013; Ogden &
94	Tsao, 2009; Park, Cleveland, Dallas, & Corn, 2016; Randolph & Dobson, 2012; Roche &
95	Guégan, 2011; Roche, Rohani, Dobson, & Guégan, 2013; Swei et al., 2011; Titcomb et al.,
96	2017; Vinson & Park, 2019).

97 Overall, there is a need for more field studies on various systems to further disentangle the 98 mechanisms underlying biodiversity-disease relationships and to inform epidemiological 99 predictions. However, conducting such studies can be highly challenging, since it requires 100 generating data on wild vertebrate and arthropod fauna as well as on circulating pathogens in 101 numerous study sites and, often, in difficult environmental contexts (e.g., tropical regions). 102 Here, we used recently developed molecular tools to explore the effects of mammal diversity 103 on the transmission of Amazonian leishmaniases. Leishmaniases are a group of human vector-104 borne diseases endemic to different tropical and subtropical regions, caused by parasites of the 105 genus Leishmania (Kinetoplastida: Trypanosomatidae) and transmitted by hematophagous 106 phlebotomine sand flies (Psychodidae) (reviewed in Bañuls, Hide, & Prugnolle, 2007). In 107 Amazonian ecosystems, several zoonotic *Leishmania* species typically coexist, with distinct 108 sylvatic transmission cycles involving different sand fly vector species and wild mammal 109 reservoir hosts (Ralph Lainson & Shaw, 2010; Rotureau, 2006). Amazonian leishmaniases 110 represent a significant public health burden in a biodiversity-rich region threatened by human 111 activities (Chavy et al., 2019; Pan American Health Organization, 2019; Rangel, Costa, & 112 Carvalho, 2014). However, the mechanisms through which biodiversity may impact the 113 transmission of these wild zoonoses remain largely unexplored.

114

Material and methods

115 Sampling

116 Sampling was performed between 2015 and 2017 in 19 different forest sites in French Guiana 117 (Figure 1, Table 1, Data S1). Sites were separated by at least ~5-km distance and chosen to 118 represent sylvatic environments with variable levels of human-induced disturbance, ranging 119 from remote and protected areas to forest patches in the vicinity of urbanized zones. When 120 several sites were sampled within a same area (i.e., Saint-Georges, Belizon, Kaw or Counami), 121 those were specifically chosen to represent contrasting situations with respect to hunting 122 pressure (e.g., closer to or further from the nearest city or accessible road, inside or outside of 123 a protected area). Human-induced disturbance was measured using the average Human 124 FootPrint index [HFP; (de Thoisy et al., 2010); updated in 2012; Table 1, Data S1]. In each 125 site, sand flies were collected using US Centers for Disease Control and Prevention (CDC) 126 miniature light traps set up across a *ca*. 1 ha plot. Traps were separated by *ca*. 50-m distance 127 from each other and left for up to six consecutive nights. Each morning, the contents of each 128 trap was collected. Sand fly females were sorted using a stereo microscope and kept in pools (corresponding to each trap-night) in microcentrifuge tubes with 95% ethanol for later 129 130 molecular analyses. A maximum of 50 individuals was included in a pool, and several pools 131 were made when more than 50 specimens were caught in a given trap (with a maximum of 132 four pools per trap, *i.e.*, 200 individuals). The total number of sand flies caught in each trap was systematically recorded, unless the contents of the trap was importantly damaged or 133 134 partially lost (mainly due to rainy conditions or manipulation during collection). Visibly blood-135 fed dipterans, including sand flies, mosquitoes (Culicidae), and biting midges 136 (Ceratopogonidae), were kept individually in microcentrifuge tubes with 95% ethanol for molecular analyses. Additional blood-fed dipterans resting during the day along tree trunks
were collected using a Prokopack aspirator (John W. Hock co., Gainesville, FL, USA) and
conserved in the same way.

140 *Laboratory*

141 We analyzed sand fly pools to identify their species composition using a previously developed 142 DNA metabarcoding protocol (Kocher, Gantier, et al., 2017). Leishmania DNA detection and 143 identification was performed on the same pools using high-throughput sequencing of kDNA 144 minicircle amplicons (Kocher, Valière, Bañuls, & Murienne, 2018). For each blood-fed 145 specimen, the dipteran species and blood meal source were identified individually as 146 previously described (Kocher, de Thoisy, Catzeflis, Valière, et al., 2017). Sand fly pools were 147 homogenized using a *Qiagen TissueLyser 2* (Qiagen, Valencia, CA, USA), and DNA was 148 extracted with the *Oiagen DNeasy Blood and Tissue kit*. For individual blood-fed specimens, 149 a modified Chelex (Bio-Rad, Hercules, CA, USA) protocol was used for DNA extraction 150 (Casquet, Thebaud, & Gillespie, 2012). The Ins16S 1 [F: TRRGACGAGAAGACCCTATA; 151 R: TCTTAATCCAACATCGAGGTC; (Clarke, Soubrier, Weyrich, & Cooper, 2014)], 12S-152 V5 [F: TAGAACAGGCTCCTCTAG; R: TTAGATACCCCACTATGC; (Riaz et al., 2011)] and leishmini [F: 5'-GGKAGGGGCGTTCTGC-3'; R: 5'-STATWTTACACCAACCCC-3'; 153 154 Kocher, Valière, Bañuls, & Murienne, 2018] PCR primers were used to amplify short 155 fragments of dipteran, vertebrate and *Leishmania* DNA, respectively. Tags of eight base pairs 156 with at least five differences between them were added at the 5' end of each primer to enable 157 multiplexing of PCR products for subsequent sequencing (Binladen et al., 2007). A Latin 158 square design was used for PCR multiplexing to allow for the detection and filtering of mistagged sequencing read (Esling, Lejzerowicz, & Pawlowski, 2015). For sand fly 159

metabarcoding, two PCR replicates were performed. PCR products were pooled according to
the multiplexing design and used for sequencing library preparation and high-throughput
sequencing on Illumina Hiseq or Miseq platforms at the GeT-PlaGe core facilities of Genotoul
(Toulouse, France).

164 *Bioinformatics*

165 Bioinformatic analyses were performed using the *OBITools 1.2.9* package (Boyer et al., 2016) 166 and R 4.0.3 (R Core team, 2020). Paired-end reads were merged with *illuminapairedend* and 167 demultiplexed based on PCR primer tags using *ngsfilter*. Reads were dereplicated using 168 *obiuniq* and sequences supported by less than ten reads in a given sample were discarded using 169 *obigrep.* Taxonomic assignments were performed using *ecotag*, with customized reference 170 DNA sequence datasets for each studied taxonomic group. For dipteran identifications, we used previously published reference datasets for neotropical sand flies (Kocher, Gantier, et al., 171 172 2017) and mosquitoes (Talaga et al., 2017), to which we added mosquito reference sequences 173 corresponding to the targeted 16S region, which we extracted from NCBI GenBank using 174 *ecoPCR*. For vertebrate identifications, we used a previously published reference dataset for 175 Amazonian mammals (Kocher, de Thoisy, Catzeflis, Huguin, et al., 2017), completed with 176 vertebrate reference sequences corresponding to the targeted 12S region extracted from 177 GenBank. For Leishmania identifications, we used a previously published dataset of kDNA 178 minicircle reference sequences (Kocher et al., 2018). ecotag employs a Lower Common 179 Ancestor algorithm that allows to perform taxonomic assignments based on the percentage of 180 identity with multiple matches in a reference dataset. In other words, a sequence matching 181 similarly to several members of a given taxon will be assigned to the corresponding taxon. For 182 dipteran and vertebrate identifications, we considered taxonomic assignments at the genus

183 level at best if the percentage of identity with the closest match was lower than 97%, in order 184 to avoid biases due to reference dataset incompleteness (*i.e.* artifactual species-level 185 identifications in cases where only one species was represented in the dataset for a given 186 genus). We then performed *de novo* sequence clustering using sumaclust 1.0.31 (Mercier, 187 Boyer, Bonin, & Coissac, 2013) with a 97% threshold. We defined molecular taxonomic units 188 (MOTU) based on *ecotag* results in case of species-level identifications, and based on *de novo* 189 clustering otherwise, in order to identify putative species within upper-level taxa. Vertebrate 190 identifications were adjusted when only a subset of the matched species was known to be 191 present in French Guiana. Because no reference sequence was available for local biting midge 192 species, we defined MOTUs within the Ceratopogonidae family based on *de novo* clustering 193 only.

194 For each sequencing library, we used the number of sequencing reads found with non-used 195 primer tag combinations to perform MOTU-based filtering of mistagged reads, as suggested 196 previously (Esling et al., 2015). Additionally, MOTUs supported by less than 100 reads in a 197 given sample were filtered out. For sand fly metabarcoding, we further filtered MOTUs that 198 (i) were not identified as Phlebotominae, (ii) were not recovered in two PCR replicates, and 199 (iii) were supported by least 2% of the sequencing reads in a given sample (a maximum of 50 200 sand flies were included in each analyzed pool). For individual blood-fed specimens, the most 201 supported dipteran and vertebrate MOTUs were retained (i.e. we did not consider the 202 possibility of multiple blood meal sources, as a conservative measure). In a few cases, the 203 resulting dipteran identification did not match the expected dipteran group (sand fly, mosquito, 204 or biting midge) and was therefore discarded. For blood meal identifications, results were 205 discarded if the majority sequence was identified as human or other potential laboratory

contaminants which were not expected in our study sites, as well as sequences assigned to
 above-order taxonomic levels which likely represented molecular or sequencing artifacts. For
 Leishmania detection, only species-level identifications were considered, and the majority
 Leishmania species was retained in each positive sample.

210 *Estimating mammal diversity from individual dipteran blood meals*

211 iDNA has recently emerged as a promising tool to perform vertebrate inventories at lower cost 212 and effort (Calvignac-Spencer et al., 2013; Kocher, de Thoisy, Catzeflis, Valière, et al., 2017; 213 Schnell et al., 2015). Here, we used a probabilistic approach to generate estimates of host 214 community composition and diversity from our iDNA data (i.e. blood meal identifications of individual dipteran specimens) while accounting for invertebrate host preferences. The model 215 is represented by a Bayesian network in Figure S1. We assume that the probability p_{sih} that a 216 blood-fed invertebrate of species *i* has fed on a host of species *h* in site *s* depends on r_{sh} , the 217 relative abundance of h in s, and on α_{sh} , the relative preference of i for h. We further assume 218 219 that the relative probability of insect *i* feeding on host h_i rather than on host h_k in site *s* is given 220 by:

221
$$\frac{p_{sih_j}}{p_{sih_k}} = \frac{r_{sh_j}\alpha_{ih_j}}{r_{sh_k}\alpha_{ih_k}} \qquad (1)$$

From (1) it follows that the probability p'_{sih} that an invertebrate of species *i* has fed on host *h* in site *s*, given that it has fed on one of the *H* host species identified in our dataset, is:

224
$$p'_{sih} = \frac{r_{sh}\alpha_{ih}}{\sum_{k=1}^{H} r_{sk} \alpha_{ik}}$$
(2)

225 Denote $\{N\}_{si} = \{n_{si1}, \dots, n_{siH}\}$ the data vector of the number of occurrences of invertebrate *i* 226 having fed on each of the *H* host species in site *s*. The probability of this data is given by the 227 multinomial distribution with parameters $\{p'\}_{si} = \{p'_{si1}, \dots, p'_{siH}\}$ and $Y_{si} = \sum_{k=1}^{H} n_{sik}$:

228
$$P(\{N\}_{si}|\{p'\}_{si}, Y_{si}) = Multinomial(\{N\}_{si}|\{p'\}_{si}, Y_{si})$$
(3)

Given S sites and I invertebrate species across all sites, we denote $\{N\} = \{\{N\}_{si}\}$ the whole dataset, $\{p'\} = \{\{p'\}_{si}\}$ the set of probability vectors and $\{Y\} = \{Y_{si}\}$ the set of sample sizes, with $i = 1, 2 \dots I$ and $s = 1, 2, \dots, S$. The likelihood of the full dataset is then given by:

232
$$P(\{N\}|\{p'\},\{Y\}) = \prod_{s=1}^{S} \prod_{i=1}^{I} Multinomial(\{N\}_{si}|\{p'\}_{si},Y_{si}) \quad (4)$$

233 The model was implemented in a Bayesian framework in *Stan* (Carpenter et al., 2017) through 234 its R interface *rstan*, in order to sample the posterior probability density of the parameters using 235 Monte Carlo Markov chain (MCMC; see Supporting Information and Figure S2 for details about the choice of priors and simulation results, as well as the Stan code). Joint samples of 236 mammal's relative abundances in each site were used to derive the Shannon index of diversity 237 $(-\sum_{h} r_{sh} \log(r_{sh}))$ and the overall proportion of species known as *Leishmania* reservoirs. 238 239 Closely related dipteran species appeared to feed on similar ranges of hosts (e.g. sand flies of the Nyssomyia genus feeding mostly on arboreal mammals, sand flies of the Psychodopygus 240 241 genus feeding mostly on armadillos, mosquitoes of the *Culex* genus feeding on a wide range 242 of vertebrates or observed ceratopogonids feeding mostly on amphibians; Table 2; Figure S3). 243 Therefore, we estimated feeding preferences at the genus level (or at the family level in the 244 case of ceratopogonids), in order to increase statistical power. Additionally, we assumed that 245 dipterans had identical preferences for vertebrates of the same order which exhibit similar

246 morphological and ecological features (with the exception of rodents which were separated 247 into terrestrial and arboreal rodents; Figure S3). Sites in which less than five dipteran blood 248 meals could be identified were not retained for the analysis. The posterior distribution of the 249 parameters was sampled using MCMC with 3 chains of 40,000 iterations, including 4,000 250 iterations for warmup. Convergence and mixing were assessed using trace plots and ESS 251 values, which were >200 for all parameters (Figures S4 and S5). The prior distribution of the 252 parameters was obtained by running the same MCMC sampling scheme as for the posterior, 253 with the likelihood fixed to a constant. The comparison of posterior and prior densities allowed 254 us to assess to what extent the posterior was driven by the data (Figures S4 and S5). The mean 255 posterior estimates of parameters were computed and used as variables for the generalized 256 linear models (GLM) described in the next section (we initially attempted to jointly estimate 257 mammal relative abundances and GLM's parameter but this led to MCMC mixing issues).

258 Effect of mammal diversity on Leishmania transmission

259 In addition to the measures of vertebrate communities obtained from individual blood meal 260 identifications, sand fly counts in traps and molecular analyses of sand fly pools were used to 261 estimate the abundance and diversity of sand flies as well as the *Leishmania* prevalence rate in 262 sand flies. All of these variables were used in a series of Bayesian GLMs to explore the impact 263 of mammal diversity changes on the disease system (Figure 1, Supporting Information, Data 264 S2). We assessed the relationship between human-induced disturbance and mammal diversity 265 using a regression of the Shannon index of mammals on the HFP. We assessed whether 266 changes of mammal diversity led to predictable changes of the mammal community 267 competence for Leishmania parasites using a regression of the relative abundance of 268 Leishmania reservoirs on the Shannon index of mammals. Furthermore, we assessed the effect

269 of changes of mammal diversity on sand fly density and diversity by regressing the number of 270 sand flies collected in each trap and the Shannon index of sand flies (estimated from 271 metabarcoding results with the R package *iNext*; (Hsieh, Ma, & Chao, 2016)) on the Shannon 272 index of mammals. We then assessed the effect of mammal community competence, sand fly 273 density and sand fly diversity on *Leishmania* transmission using a regression of *Leishmania* 274 prevalence rates in sand flies on the relative abundance of *Leishmania* reservoirs, the expected 275 number of sand flies collected per trap and the Shannon index of sand flies (used as predictor 276 variables in the same regression). We checked for spatial autocorrelation of the different 277 variables with the Moran's I autocorrelation index as implemented in the R package ape 278 (Paradis, Claude, & Strimmer, 2004), using the inverse of pairwise geographical distances 279 between sites as the weight matrix. For all regressions, predictor variables were standardized. 280 and weakly informative normal priors were used for regression coefficients. Prior distributions 281 of slope coefficients were centered around 0, while that of regression intercepts were centered 282 around the mid-range value of the corresponding dependent variable. The posterior 283 distributions of slope coefficients were used to assess the significance of inferred relationships. 284 We evaluated the cumulative effect of mammal diversity changes on *Leishmania* transmission 285 by generating posterior predictions of the Leishmania prevalence rate and of the expected 286 number of infected sand flies per trap for different values of the Shannon index of mammals. 287 We visualized sampled mean prediction curves against the Shannon index of mammals (across 288 the range of values observed in our study), and used the distribution of their average slope to 289 measure the predicted effects.

290 **Results**

291 Sampling and molecular analyses

292 In total, we collected 18,508 sand fly females, which were gathered in 666 pools used for sand 293 fly metabarcoding and *Leishmania* detection (Table 1, Data S1). After DNA extraction and 294 amplification, high-throughput sequencing and bioinformatic filtering, 600 (90.1%) sand fly 295 pools could be characterized with metabarcoding, and Leishmania DNA was detected in 175 296 (26.3%) of them. We further collected 855 blood-fed dipterans, including 715 sand flies, 123 297 mosquitoes and 17 biting midges that were analyzed individually (Table S1). Dipteran 298 identification was successful in 91.7% of the individuals (although not necessarily at the 299 species level), and their blood meal content was identified in 75.9% of the cases. Both dipteran 300 and blood meal identifications were successful for 602 (70.4%) individuals. Three sites in 301 which less than five blood-fed dipterans could be identified were not retained for statistical 302 analyses (Figure 1, Table 1).

303

3 Sand fly, vertebrate and Leishmania identifications

304 In total, we identified 63 sand fly MOTUs in sand fly pools (Table S1, Data S1), which is fairly 305 consistent with the known sand fly species richness of French Guiana (about 80 species 306 recorded so far). 34 of these were identified at the species level, including seven known or 307 suspected Leishmania vector species in the region: Bichromomyia flaviscutellata, Nyssomyia 308 umbratilis, Psychodopygus ayrozai, Ps. panamensis, Ps. squamiventris maripaensis, 309 Trichophoromyia ubiquitalis, Viannamyia furcata (Rotureau, 2006). The most abundant 310 species were Ps. squamiventris maripaensis, Th. ininii, Trichopygomyia trichopyga and Th. 311 ubiquitalis (25.2%, 18.3%, 18.0% and 10.9% of the estimated number of individuals,

312 respectively). Five Leishmania species were detected in sand fly pools: L. lainsoni, L. 313 amazonensis, L. naiffi, L. braziliensis, L. guvanensis; the most frequent being L. lainsoni and 314 L. naiffi (48.6% and 43.4%% of the positive samples respectively). Blood-fed specimens 315 belonged to 51 dipteran MOTUs, and blood meal analyses revealed a total of 52 vertebrate 316 MOTUs (Table 2, Figure S3, Data S1), including 28 mammals, among which 11 were 317 recognized Leishmania reservoir hosts in the region: Didelphis marsupialis, Philander 318 opossum, Metachirus nudicaudatus, Choloepus didactylus, Dasypus novemcinctus, Tamandua 319 tetradactyla, Coendou melanurus, C. prehensilis, Dasyprocta leporina, Proechymis cuvieri 320 and *P. guyannensis* (Rotureau, 2006). Our results revealed contrasting host preferences across 321 sand fly species, and were consistent with existing knowledge [e.g. Nyssomyia spp. feeding 322 mostly on sloth and other arboreal mammals (Christensen, Arias, de Vasquez, & de Freitas, 323 1982), B. flaviscutellata feeding mostly on terrestrial rodents (R. Lainson & Shaw, 1968), and 324 *Psychodopygus* spp. feeding mostly on armadillos (Le Pont, 1990); Table 2; Figure S3].

325 Statistical analyses

326 We estimated mammal diversity (Shannon index) and the relative abundance of *Leishmania* 327 reservoirs in each site based on dipteran blood meal identifications using a probabilistic approach (Figure S1, S2). We then used these estimates in a series of Bayesian regressions to 328 329 investigate the effect of mammal diversity on the transmission of *Leishmania* parasites. 330 MCMC trace plots and ESS values indicated convergence and correct sampling of the posterior 331 distribution for all parameters (Figures S4, S5, S6). Mean estimates of the Shannon index of 332 mammals across our study sites ranged from 0.46 to 2.7 (Table 1), and these correlated 333 negatively with the human footprint index (HFP; (de Thoisy et al., 2010); mean effect: -0.34, 334 90%CI [-0.61,-0.07]; Figure 1B). Spatial autocorrelation was not detected for the HFP

335	(expected/observed Moran's I: -0.067/0.056; p-value: 0.3), nor for the Shannon index of
336	mammals (expected/observed Moran's I: -0.067/-0.13; p-value: 0.6). The Shannon index of
337	mammals correlated negatively with the relative abundance of Leishmania reservoirs (-0.68,
338	90%CI [-0.99,-0.38]; Figure 2A), positively with sand fly density (0.4, 90%CI [0.13,0.67];
339	Figure 2B), and negatively (but weakly) with the Shannon index of sand flies (-0.19, 90%CI [-
340	0.38,0]; Figure 4SA). Furthermore, we estimated that the prevalence rate of <i>Leishmania</i> in sand
341	flies correlated positively with the relative abundance of Leishmania reservoirs (0.43, 90%CI
342	[0.27,0.59]; Figure 2C), but did not correlate with sand fly density or diversity (0.03, 90%CI
343	[-0.12,0.17] and 0.08, 90%CI [-0.07,0.23], respectively; Figure S7B,C). Finally, posterior
344	predictions indicated that the Shannon index of mammals correlated negatively with the
345	Leishmania prevalence rate in sand flies but did not correlate with the density of infected sand
346	flies (Figure 2D,E).

347 **Discussion**

348 With this work, we show that wild vector-borne disease systems can be efficiently studied 349 using DNA metabarcoding of arthropod vectors, which allows measuring arthropod, vertebrate 350 and parasite communities altogether. Dipteran blood meal analyses allowed to identify a large 351 variety of vertebrates across our study sites, confirming the potential of invertebrate-derived 352 DNA (iDNA) approaches for biodiversity monitoring (Calvignac-Spencer et al., 2013; Kocher, 353 de Thoisy, Catzeflis, Valière, et al., 2017; Schnell et al., 2015). Based on blood meal 354 identifications, we estimated the Shannon index of mammals, which correlated negatively with 355 the human footprint index (Figure 1B), as expected (de Thoisy et al., 2010). This suggests that 356 the variation of mammal diversity across our study sites was, at least partly, linked to humaninduced disturbance. We then explored the effect of mammal diversity on Leishmania disease 357

358 systems and transmission. In particular, we assessed the occurrence of a dilution effect 359 potentially leading to reduced transmission with higher mammal diversity. In the case of 360 vector-borne diseases, a dilution effect can be expected only if arthropod vectors feed on 361 various host species, including some that are poorly competent for pathogen transmission. This 362 is the case for Amazonian leishmaniases, since sand fly vector species from which blood meals 363 could be analyzed were observed to have fed on different hosts, including some that are not 364 known as Leishmania reservoirs (Table 2, Figure S3). For example, while most Ny. umbratilis 365 individuals had fed on two-toed sloths (main vector and reservoir of L. guyanensis, 366 respectively), and most *Psychodopygus* spp. had fed on nine-banded armadillos (main vector 367 and reservoirs of *L. naiffi*), a significant proportion of them had fed, respectively, on different 368 primates and large terrestrial mammals which are not known to act as *Leishmania* reservoirs. 369 Therefore, Leishmania parasites may indeed end up in "diluting" hosts which could contribute 370 to reduced transmission of these disease agents. This was further supported by the observation 371 of a positive correlation between the relative abundance of *Leishmania* reservoir hosts and 372 *Leishmania* prevalence rate in sand flies (Figure 2C).

373 Another important assumption of the dilution effect hypothesis is that species contributing the 374 most to pathogen transmission dominate in disturbed and less diverse communities. Ecological 375 and evolutionary hypotheses have suggested the existence of such a positive relationship 376 between host competence for pathogens and resilience to disturbance, leading to a general 377 increase of the overall community competence with biodiversity loss (Johnson, Ostfeld, & 378 Keesing, 2015; Keesing et al., 2010; Ostfeld & Keesing, 2012). Species that are resilient to 379 changing environments are frequently characterized by fast life history strategies, including 380 low investment in adaptive immunity and high reproductive rate, yielding an important influx 381 of susceptible individuals in the population. In addition, pathogens may adapt predominantly 382 to resilient host species, because these hosts are generally widespread, mobile and abundant, 383 therefore constituting the most frequently encountered resource. Empirical evidence has 384 supported these ideas (García-Peña et al., 2016; Han, Schmidt, Bowden, & Drake, 2015; 385 Johnson et al., 2019; Johnson, Preston, Hoverman, & Richgels, 2013; Johnson et al., 2012), 386 although it seems that the situation may vary depending on the taxa under consideration (Gibb 387 et al., 2018; H. Young, Griffin, Wood, & Nunn, 2013). Here, we show that mammal species 388 known as *Leishmania* reservoirs indeed dominate less diverse mammal communities, with a 389 233.6% (90%CI: [293.0%,190.8%]) increase of their relative abundance along the range of 390 estimated mammal's Shannon index (Figure 2A). In sum, these results suggest a predictable 391 effect of local mammal diversity changes on the overall host community competence for 392 Leishmania parasites, contributing to an increase of vectorial transmission with decreasing 393 mammal diversity, in accordance with the dilution effect hypothesis.

394 However, arthropod vector ecology should also be accounted for when investigating the impact 395 of biodiversity changes on a vector-borne disease. In particular, higher vector density should 396 be associated with more frequent host-pathogen contacts and increase transmission (Smith et 397 al., 2012). Additionally, when several arthropod species can act as vectors for a given 398 pathogen, higher arthropod diversity can result in higher pathogen transmission through an 399 overall increase of vector abundance, or due to functional complementarity between vector 400 species (Park et al., 2016; Roche et al., 2013). Given that vertebrates constitute trophic 401 resources for blood-feeding arthropods, it can be expected that arthropod communities are 402 partially driven by the abundance and composition of local vertebrate fauna. This might, 403 however, depend on the ecology of the considered arthropod species. For instance, highly

404 mobile mosquitoes might be little affected by changes in host density, which is a classical 405 assumption in epidemiological models (Dobson, 2004), while ticks may be more sensitive to 406 the presence of suitable hosts in their immediate environment (Ogden & Tsao, 2009; Randolph 407 & Dobson, 2012; Swei et al., 2011; Titcomb et al., 2017). Little is known in this regard for 408 sand flies, which, despite being flying insects, have relatively small flight ranges (Casanova, 409 Costa, & Natal, 2005; Morrison, Ferro, Morales, Tesh, & Wilson, 1993). Here, we observed a 410 positive relationship between mammal diversity and sand fly density (Figure 2B), which 411 suggests that sand flies may indeed be affected by changes in local mammal communities. 412 Such a relationship could be mediated by a correlation between mammal diversity and overall 413 mammal biomass, resulting in greater availability of blood meal resources for sand flies in 414 more diverse ecosystems. However, our data does not provide information regarding the 415 absolute abundance of mammals, and it is difficult draw conclusions about the causality of 416 such a relationship since some environmental factors could affect both mammal and arthropod 417 species communities. On the other hand, our results point to a negative correlation between 418 mammal and sand fly diversity (although weakly significant; effect 90%CI: [-0.38%,0.00%]; 419 Figure S7A), which suggest that different factors might shape mammal and sand fly 420 communities. This further highlights the need for more research to understand the factors 421 shaping sand fly assemblages in sylvatic systems and their potential consequences for 422 *Leishmania* transmission.

423 Our results did not reveal an effect of sand fly density or diversity on the prevalence rate of 424 *Leishmania* parasites in sand flies (Figure S7B, C). Thus, the proportion of *Leishmania* 425 reservoir hosts in mammal communities appeared as the main driver of *Leishmania* 426 transmission, resulting in a negative effect of mammal diversity on *Leishmania* prevalence

427 rate, through host dilution (Figure 2D). However, the prevalence of a pathogen in vectors is 428 not necessarily a relevant measure of disease transmission, and the density of infected vectors 429 should rather be considered for this matter. For a given prevalence rate, higher vector density 430 should be associated with a higher density of infected vectors. Therefore, the positive 431 relationship observed between mammal diversity and sand fly density across our study sites 432 might act on leishmaniasis transmission in opposition to the dilution effect, which it may 433 attenuate, cancel, or even reverse. This shows that the alteration of mammal diversity is 434 associated with changes in the ecosystem that independently impact *Leishmania* transmission 435 in contrasting ways. Posterior predictions of the density of infected sand flies indicate a weak 436 overall impact of mammal diversity changes on Leishmania transmission (Figure 2E). 437 Therefore, it seems that the observed dilution and amplification effects compensate for each 438 other in the system studied here. However, one or the other could predominate in other contexts 439 depending on characteristics of the considered system and environmental conditions that need 440 to be further determined. This constitutes additional evidence that biodiversity changes may 441 impact vector-borne pathogen transmission through concurrent mechanisms, and further 442 stresses the importance of better accounting for arthropod vector ecology in biodiversity-443 disease research.

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716 Data Accessibility and Benefit-Sharing

717 Sample metadata: Data and code used for statistical analyses are available as supplementary

718 material.

- 719 Genetic data: Sequencing data has been deposited at the Dryad database (doi:
 720 10.5061/dryad.44j0zpcfp).
- 721 Benefits generated: Benefits from this research accrue from the sharing of our data and results
- 722 on public databases as described above.

723 Author Contributions

- JM, AK, BdT, ALB and JFG designed the study. AK, JM, JCG, AC, BdT, ALB, MG, GP, RG,
- 725 ID and PMF conducted the field work. AK and SM performed the laboratory work. AK and
- JC analyzed the data. AK wrote the initial version of the manuscript, which was edited by JM,
- 727 JC, BdT and JFG, and all authors contributed to its improvement.

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730 Figures



Figure 1: (A) Location of study sites (French Guiana). (B) Regression of the Shannon index of mammals on human-induced disturbance. Shannon indices of mammals were estimated through dipteran blood meals in each site. The level of human-induced disturbance was measured using the human footprint index (de Thoisy et al., 2010). The mean prediction curve is depicted with 90% CI. In the inlet figure, the posterior density of the regression's slope coefficient is represented (dotted lines are positioned at 5% and 95% quantiles; the red line is positioned at x=0).

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740Figure 2: Effect of mammal diversity changes on Leishmania transmission, assessed by a series of Bayesian741regressions and posterior simulations. The most significant relationships are summarized in the top-panel742diagram (but see Figure S7 for the complete model). Direct and indirect relationships are represented with743solid and dashed arrows, respectively. Positive, negative, and non-significant effects are represented by744red, blue and black arrows, respectively. Light grey arrows represent deterministic relationships. For each

745 studied relationship, regression plots are presented and referenced with the corresponding letter in the 746 lower panels. (A,B,C) Regression curves are depicted with 90% CI, and the posterior density of regression's 747 slopes are represented in inlet plots (dotted lines are positioned at 5% and 95% quantiles; the red line is 748 positioned at x=0). (D,E) The indirect (cumulative) effect of mammal diversity on Leishmania transmission 749 was assessed by using posterior samples of model parameters to predict the Leishmania prevalence rate 750 and the expected number of infected sand flies per trap. Sampled mean prediction curves are plotted (a 751 subset of 200 curves, in order to facilitate visualization). Curves indicating an overall increase or decrease 752 across the range of predictor values are depicted in red and blue, respectively. Inner plots represent the 753 posterior density of the average slope of mean prediction curves.

Tables 755

756 757

Table 1: Details about the study sites, including sampling size and estimates of the different variables used to describe the ecological system.

Site	Nb. sand flies	Nb. blood-fed dipt. (success. ident.)	HFP	Mammal diversity (Shannon index)†	Prop. <i>Leish</i> . reservoirs†	Sand fly density (indiv. per trap)†	Sand fly diversity (Shannon index)†	<i>Leishmania</i> prev. rate (%)†
Belizon: A	1,028	17 (13)	36.0	2.22 [1.53,2.83]	0.42 [0.19,0.67]	44.35 [42.27,46.45]	2.18 [2.16,2.16]	1.05 [0.8,1.33]
Belizon: B	1,176	67 (38)	30.1	1.68 [1.16,2.18]	0.66 [0.38,0.89]	55.75 [53.31,58.25]	2.07 [2.04,2.04]	1.64 [1.37,1.92]
Belizon: C	1,298	22 (10)	17.0	1.95 [1.23,2.66]	0.71 [0.43,0.95]	58.69 [56.17,61.26]	1.72 [1.71,1.71]	1.71 [1.46,1.98]
Camp du tigre: Layon BdT	450	43 (30)	40.9	0.46 [0.13,0.95]	0.98 [0.91,1]	11.87 [10.97,12.79]	2.38 [2.35,2.35]	3.13 [2.18,4.22]
Counami: Carbet	349	60 (40)	21.0	2.27 [1.69,2.79]	0.65 [0.43,0.87]	15.18 [14.11,16.3]	1.45 [1.4,1.4]	1.4 [1.09,1.77]
Counami: Crossing	719	35 (32)	17.0	1.1 [0.49,1.85]	0.87 [0.58,1]	38.17 [36.06,40.33]	1.48 [1.46,1.46]	2.2 [1.7,2.78]
Counami: Patagai	572	13 (6)	17.0	1.57 [0.69,2.45]	0.49 [0.13,0.87]	19.43 [18.16,20.7]	1.69 [1.67,1.67]	1.08 [0.83,1.35]
Counami: T1	376	13 (8)	9.7	1.3 [0.36,2.2]	0.3 [0.04,0.68]	12.6 [11.58,13.65]	2.52 [2.47,2.47]	0.89 [0.54,1.31]
Counami: T2	817	97 (74)	4.0	2.09 [1.52,2.61]	0.69 [0.49,0.87]	30.7 [29.06,32.39]	1.9 [1.88,1.88]	1.64 [1.38,1.93]
Kaw: NR Tresor	2,964	50 (31)	8.0	2.32 [1.75,2.83]	0.61 [0.41,0.82]	109.27 [106.04,112.56]	1.68 [1.68,1.68]	1.46 [1.15,1.79]
NR Nouragues: Museum plot	2,676	196 (146)	4.0	2.41 [2.08,2.7]	0.18 [0.09,0.29]	142.16 [138.52,145.78]	1.93 [1.93,1.93]	0.68 [0.47,0.93]
NR Trinite: Layon D	795	14 (9)	0.0	2.27 [1.59,2.87]	0.48 [0.27,0.72]	26.56 [25.05,28.13]	1.56 [1.55,1.55]	1.04 [0.81,1.3]
Saint-Georges: Bridge	1,647	100 (70)	32.4	1.75 [1.4,2.11]	0.53 [0.31,0.76]	54.83 [52.06,57.66]	0.94 [0.93,0.93]	1.06 [0.76,1.4]
Saint-Georges: SG30B	182	38 (36)	22.7	1.03 [0.37,1.75]	0.89 [0.66,1]	15.25 [13.49,17.05]	2.39 [2.32,2.32]	2.67 [1.93,3.52]
Saint-Georges: SG6A	1,606	38 (27)	26.5	1.98 [1.42,2.52]	0.67 [0.48,0.84]	89.1 [85.42,92.83]	1.2 [1.19,1.19]	1.49 [1.13,1.9]
Saint-Georges: Track ONF	749	35 (21)	14.7	2.66 [2.19,2.99]	0.35 [0.19,0.52]	42.64 [40.22,45.05]	1.4 [1.39,1.39]	0.8 [0.59,1.03]

[†]Mean estimate (90% HPD)

Table 2: List of vertebrates identified in blood meals of each dipteran species or MOTU

Species/MOTU	Nb.	Vertebrates identified			
Ceratopogonidae MOTU 1	3	Rhinella cf. margaritifera (67%); Hyloidea MOTU 1 (33%)			
Ceratopogonidae MOTU 2	2	Osteocephalus MOTU 1 (50%); R. cf. margaritifera (50%)			
Ceratopogonidae MOTU 3	1	Choloepus didactylus (100%)			
Ceratopogonidae MOTU 4	1	Osteocephalus MOTU 1 (100%)			
Ceratopogonidae MOTU 5	2	Osteocephalus MOTU 1 (100%)			
Aedini MOTU 1	1	Adenomera andreae (100%)			
Culex imitator	1	Squamata MOTU 1 (100%)			
Culex MOTU 1	58	Prochimus cuvier (36%). Thampohilus pigrocinereus (28%). Cupiculus paca (7%): Dasvorocta lenorina (3%). Didelphis			
	00	marsupialis (3%); Mazama nemorivaga (3%); Proechimys guyannensis (3%); Thamophilus MOTU 1 (3%); Bradypus tridactylus (2%); Chelonoidis denticulatus (2%); Dasppus novemcinctus (2%); Echimys chrysurus (2%); Metachirus nudicaudatus (2%); Oecomys rutilus (2%); Tapirus terrestris (2%)			
Culex MOTU 2	14	Squamata MOTU 1 (14%); <i>T. nigrocinereus</i> (14%); Accipitrinae MOTU 1 (7%); <i>Anolis fuscoauratus</i> (7%); <i>Bothrops atrox</i> (7%); <i>D. leporina</i> (7%); <i>D. novemcinctus</i> (7%); <i>Gonatodes humeralis</i> (7%); <i>Hypsiboas</i> MOTU 1 (7%); <i>Philander opossum</i> (7%); <i>Polychrus marmoratus</i> (7%); <i>Thamnophilus</i> MOTU 1 (7%)			
Culex MOTU 3	2	D. marsupialis (50%); T. nigrocinereus (50%)			
Culex sp.stJ	1	Thamnophilus MOTU 2 (100%)			
Culex sp.stK	1	D. leporina (100%)			
Culex sp.stL	1	Osteocephalus MOTU 1 (100%)			
Culicinae MOTU 1	3	P. cuvieri (67%): Thampophilus MOTU 1 (33%)			
Limatus flavisetosus	1	Tinamus maior (100%)			
Ochlerotatus serratus	5	D leporina (40%): Thampophilus MOTU 1 (40%): M nemoriyada (20%)			
Psorophora ferox	1	Avaparta acouchy (100%)			
Sabethini MOTU 1	4	D lenorina (25%). D novemcinctus (25%): G humeralis (25%): P onossum (25%)			
Bichromomyia flaviscutellata	5	D. leporina (40%); P. guyannensis (40%); P. cuvieri (20%)			
Evandromvia brachvphalla	1	P. cuvieri (100%)			
Ev infraspinosa	4	D powercipicitis (50%): C. didactylus (25%): M acouchy (25%)			
Ev monstruosa	1	D lengina (100%)			
Ev. sericea	5	Lachesis mita (60%): M. acouchy (40%)			
Ev walkeri	1	Isothix sinamariensis (100%)			
Pintomvia serrana	5	Aloualta seniculus macconnelli (60%): C. didactulus (20%): Pithecia pithecia (20%)			
Micropygomvia rorotaensis	1	Squamata MOTU 1 (100%)			
Nyssomvia sylvicola	2	D leporina (100%)			
Ny. umbratilis	13 8	C. didactylus (49%); Coendou melanurus (15%); A. macconnelli (14%); Ateles paniscus (9%); D. novemcinctus (4%); Coendou prehensilis (3%); Tamandua tetradactyla (2%); Cebus olivaceus (1%); Nasua nasua (1%); P. pithecia (1%); Psophia crepitans (1%);			
Ny. yuilli pajoti	18	A. macconnelli (17%); C. didactylus (17%); A. paniscus (11%); C. melanurus (11%); D. novemcinctus (11%); P. crepitans (11%); T. tetradactyla (11%); C. prehensilis (6%); T. terrestris (6%)			
Pi, pacae	1	M. nemorivaga (100%)			
Pressatia choti	7	D. Jeporina (86%): G. humeralis (14%)			
Psychodopygus amazonensis	32	D. novemcinctus (44%); D. leporina (16%); T. terrestris (16%); Dasypus kappleri (12%); D. marsupialis (6%); Mazama americana (3%); Tayassu pecari (3%)			
Ps. ayrozai	79	D. novemcinctus (84%); D. kappleri (15%); Thamnophilus MOTU 1 (1%)			
Ps. claustrei	11	C. paca (36%); M. acouchy (36%); D. leporina (18%); M. nemorivaga (9%)			
Ps. hirsutus	34	D. novemcinctus (59%); M. americana (15%); T. terrestris (15%); D. kappleri (12%)			
Ps. MOTU 1	8	D. novemcinctus (50%); D. leporina (12%); D. kappleri (12%); M. acouchy (12%); T. terrestris (12%)			
Ps. MOTU 10	1	D. novemcinctus (100%)			
Ps. MOTU 2	5	D. novemcinctus (40%); T. terrestris (40%); Pecari tajacu (20%)			
Ps. MOTU 3	5	D. novemcinctus (60%); D. kappleri (40%)			
Ps. MOTU 5	3	D. novemcinctus (100%)			
Ps. panamensis	3	D. leporina (100%)			
Ps. s. maripaensis	12	D. novemcinctus (67%): M. americana (11%): D. kappleri (10%): T. terrestris (5%): M. nemorivaga (3%): C. paca (2%): A.			
	0	paniscus (1%); P. guyannensis (1%); Thamnophilus MOTU 1 (1%)			
Sciopemyia sordellii	2	Chiasmocleis shudikarensis (50%); Hyloidea MOTU 2 (50%)			
Trichophoromyia ininii	1	C. paca (100%)			
Trichopygomyia trichopyga	6	D. novemcinctus (50%); C. paca (33%); D. kappleri (17%)			
Viannamyia tuberculata	1	C. melanurus (100%)			