

1 Accuracy, limitations and cost-efficiency of 2 eDNA-based community survey in tropical frogs

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4 Miklós Bálint^{1,2}, Carsten Nowak^{1,2}, Orsolya Márton^{1,3}, Steffen U. Pauls^{1,2}, Claudia
5 Wittwer¹, Jose Luis Aramayo B.⁴, Arne Schulze⁵, Thierry Chambert⁶, Berardino Cocchiararo^{1,2},
6 Martin Jansen¹

7

8 ¹Senckenberg Research Institute and Natural History Museum Frankfurt,
9 Senckenberganlage 25, 60325 Frankfurt, Germany

10 ²LOEWE Centre for Translational Biodiversity Genomics (LOEWE-TBG),
11 Senckenberganlage 25, 60325 Frankfurt, Germany

12 ³Institute for Soil Sciences and Agricultural Chemistry, Centre for Agricultural Research,
13 Hungarian Academy of Sciences, Herman Otto str. 15, H-1022 Budapest, Hungary

14 ⁴Museo Historia Natural Noel Kempff Mercado (Universidad Autónoma Gabriel Rene
15 Moreno), Av. Irala 565, Santa Cruz de la Sierra, Bolivia

16 ⁵Hessisches Landesmuseum Darmstadt (HLMD), Friedensplatz 1, 64283 Darmstadt,
17 Germany

18 ⁶Pennsylvania State University, Department of Ecosystem Science and Management,
19 University Park, PA 16802, USA

20 Abstract

21Rapid environmental change in highly biodiverse tropical regions demands efficient
22biomonitoring programs. While existing metrics of species diversity and community composition
23rely on encounter-based survey data, eDNA recently emerged as alternative approach. Costs
24and ecological value of eDNA-based methods have rarely been evaluated in tropical regions,
25where high species richness is accompanied by high functional diversity (e.g. the use of
26different microhabitats by different species and life-stages). We first tested whether estimation
27of tropical frogs' community structure derived from eDNA data is compatible with expert field
28assessments. Next we evaluated whether eDNA is a financially viable solution for biodiversity
29monitoring in tropical regions. We applied eDNA metabarcoding to investigate frog species
30occurrence in five ponds in the Chiquitano dry forest region in Bolivia and compared our data
31with a simultaneous visual and audio encounter survey (VAES). We found that taxon lists and
32community structure generated with eDNA and VAES correspond closely, and most deviations
33are attributable to different species' life histories. Cost efficiency of eDNA surveys was mostly
34influenced by the richness of local fauna and the number of surveyed sites: VAES may be less
35costly in low-diversity regions, but eDNA quickly becomes more cost-efficient in high-diversity
36regions with many sites sampled. The results highlight that eDNA is suitable for large-scale
37biodiversity surveys in high-diversity areas if life history is considered, and certain precautions in
38sampling, genetic analyses and data interpretation are taken. We anticipate that spatially
39extensive, standardized eDNA biodiversity surveys will quickly emerge in the future.

40 Keywords

41 Amphibians, metabarcoding, tropical biodiversity, conservation, community ecology, cost
42comparison

43 Introduction

44 Improvements on most biodiversity loss indicators lag behind the 20 “Aichi Biodiversity
45Targets” (UNEP 2016) that aim to reduce the decline of biodiversity by 2020 (Tittensor *et al.*
462014). An important component of the biodiversity crisis is the extinction of species. Based on
47current trends in mammals, birds, reptiles and amphibians, it has been projected that the
48biodiversity crisis may lead to the 6th Mass Extinction over the next three centuries if all
49threatened species go extinct (Barnosky *et al.* 2011). Current rates of extinction may even be
50much higher if one considers the extinction that likely occurred during the last few decades-
51centuries, but went unnoticed because the now-extinct species had small ranges, were never
52described or only described on the eve of their extinction (Pimm *et al.* 2014; Lees & Pimm
532015). It is however difficult to assess which species are endangered and to what extent. First,
54most taxa remain poorly described: in some highly diverse regions many species will likely go
55extinct before they are discovered (Costello *et al.* 2013; Lees & Pimm 2015). Second, cryptic
56genetic diversity is common within morphospecies (Pfenninger & Schwenk 2007; Pauls *et al.*
572013), and global change may impact cryptic diversity more severely than morphospecies
58(Bálint *et al.* 2011). Third, data on population-level trends is scarce, even for well-known species
59(Butchart *et al.* 2010). Better population-level biodiversity data is thus urgently needed to 1)
60understand biodiversity patterns and extinction threats, 2) improve forecasting abilities about
61future biodiversity, and 3) improve humanity’s responses to the challenges of biodiversity loss.
62This data is crucial in times when conservation action is increasingly demanded by society
63(Tittensor *et al.* 2014).

64 The importance of internationally coordinated, standardized biodiversity data collection is
65long recognized both in science and in conservation (Henry *et al.* 2008). This is particularly true
66for the most biodiverse areas. The tropics are generally underrepresented in ecological studies

67(Clarke *et al.* 2017; Stroud & Feeley 2017). In addition, encounter-based data collection is
68logistically challenging since it needs considerable funds to bring enough specialists for different
69organismic groups to remote areas, thus insufficient funds and expertise often limit
70comprehensive surveys. Indirect species records through environmental DNA are increasingly
71heralded as an alternative to encounter-based surveys (Thomsen & Willerslev 2015; Pedersen
72*et al.* 2015). eDNA also facilitates the standardization of biodiversity surveys at both regional
73and global scales since community composition data can be obtained by high-throughput-
74sequencing of standardized, taxonomically informative marker genes (metabarcoding) (Taberlet
75*et al.* 2012a; Cristescu 2014). Aquatic or semiaquatic vertebrates such as frogs and other
76amphibians or fish have been early targets of eDNA based studies, either focusing on the
77detection of single species (e.g. Ficetola *et al.* 2008; Goldberg *et al.* 2011; Jerde *et al.* 2011;
78Thomsen *et al.* 2012a) or entire communities (e.g. Thomsen *et al.* 2012b; Valentini *et al.* 2016;
79Shaw *et al.* 2016; Yamamoto *et al.* 2017). The use of next-generation sequencing approaches
80led to a boost in data acquisition (Taberlet *et al.* 2012b) and is considered to make important
81contributions to biodiversity research (Rees *et al.* 2014; Bohmann *et al.* 2014). eDNA-based
82metabarcoding may present one of several tools needed to globally coordinate initiatives for
83ecosystem monitoring and sustainable management (Bush *et al.* 2017; Schmeller *et al.* 2017).

84 In this study we evaluate whether eDNA metabarcoding is suitable for inventories of
85frogs, a group with particular high species-diversity in tropical regions. Frogs and other
86amphibians are sentinel victims of the biodiversity crisis: more than one-third of the
87approximately 7500 described species are endangered (Stuart *et al.* 2004; Bishop *et al.* 2012;
88Whittaker *et al.* 2013). Frogs are also known for being a highly diverse, but incompletely
89described taxon, especially in the tropics (Ferrão *et al.* 2016; Caminer *et al.* 2017). Many
90“widespread” morphospecies harbor considerable cryptic genetic diversity and are better
91considered complexes of closely related species with much smaller ranges (Fouquet *et al.* 2007;

92Gehara *et al.* 2013, 2014; Ortega-Andrade *et al.* 2015). Efficient implementation of amphibian
93conservation measures (e.g. the prioritization of areas for conservation, or informing society and
94stakeholders about conservation needs) are only possible with geographically broad-scaled
95fine-grain, taxonomically well-resolved faunistic data, but our current understanding of present
96and future amphibian biodiversity is often based on rare, spatially and temporally scattered
97observations of phenotypically defined taxa.

98 First efforts have been made to test the suitability of eDNA for the survey of tropical frog
99biodiversity (Lopes *et al.* 2016), but important practical aspects remain unaddressed. First, it is
100not clear which fraction of the local species pool is represented by amphibian eDNA in tropical
101water bodies. Existing comparisons of encounter-based surveys and eDNA either do not include
102non-adult life stages, or they use already compiled fauna lists for the evaluation of eDNA
103performance without consideration of life history traits or behavioral aspects at the moment of
104sampling. However, temporal and spatial patterns of microhabitat use by frogs is largely
105species-specific and influenced by phenology and reproduction modes (e.g. Duellmann & Pyles
1061983, Haddad & Prado 2005, Wells 2010), which can induce strong biases in biodiversity data
107(Petitot *et al.* 2014). Second, the degree to which aquatic eDNA can provide accurate
108assessments of community structure remains largely unevaluated. Most studies to date have
109only investigated the correspondence between encounter-based and metabarcoded taxon lists
110(Miya *et al.* 2015; Valentini *et al.* 2016), although community structure assessments themselves
111may be of higher importance for many applications (Ji *et al.* 2013; Elbrecht *et al.* 2017). Third, it
112is not clear whether, and under what conditions eDNA is financially efficient since comparisons
113are lacking (Lopes *et al.* 2016), although these comparisons are essential for deciding on data
114collection strategies.

115 Here we address whether 1) detectability of tropical amphibians with eDNA is linked to
116species' life history, and 2) eDNA sampling provides accurate data for the characterization of

117community assembly. Finally, we present a framework for cost comparisons between
118encounter- and eDNA-based biodiversity survey that may be adapted to other systems beyond
119amphibians. We compare the results of long- and short-term encounter-based field surveys, and
120an eDNA survey of tropical amphibians in a well-characterized high-diversity area in Bolivia.

121

122

123 **Materials and methods**

124 The study area is located the Chiquitano region of Bolivia, a forest-savanna ecotone
125between Amazon, Cerrado and Gran Chaco in a transition zone among humid and dry forests
126that are special in regard to their taxonomic and functional diversity (Castro *et al.* 1999). The
127region contains the largest intact, old-growth block of seasonally dry tropical forests in South
128America (Miles *et al.* 2006; Power *et al.* 2016). Samples were collected from ponds in the
129vicinity of the Biological Station “Centro de Investigaciones Ecológicas Chiquitos” on the San
130Sebastián cattle ranch (S16.3622, W62.00225, 500 m a.s.l.), 24 km south of the town of
131Concepción, Province of Ñuflo de Chávez, Santa Cruz Department, Bolivia. A description of the
132area is given by Schulze *et al.* (2009). This area is well characterized with respect to
133amphibians, including both larvae and adults (Schulze *et al.* 2015). Intensive long-term
134assessments have resulted in the detection of 45 frog species in the area (e.g. Jansen 2009;
135Schulze *et al.* 2009, 2015; Jansen *et al.* 2011), as well as the discovery of cryptic intraspecific
136diversity (e.g. Jansen *et al.* 2011; Jansen *et al.* 2016).

137 We sampled five ponds (T1 – T5) for this study between 19 and 23 November 2014 (see
138also section “Site and ponds” in the Supplemental information). At these ponds, 35 species
139have been recorded in previous long-term surveys. Water samples were obtained from three
140sampling points at each pond (Fig. 1): These water samples consisted of four liters of water
141collected into two two-liter silicon bottles. One sample of 2 x 2 liters was taken at each sampling
142point of ponds T1 - T4. Three samples of 2 x 2 liters were taken on each sampling points of
143pond T5 to check for sampling variation, i.e. whether replication in the sampling records more
144variation in community composition compared to replication in PCR steps. We also sampled and
145filtered water from two field controls: Tap water from the station from a covered well, and water

146 from an aquarium with tadpoles of *Leptodactylus vastus* at the station. The water samples were
147 filtered at the station immediately after collection.

148 Samples were filtrated with a vacuum filtration unit (Duran Group, Wertheim, Germany),
149 which was connected to a vacuum membrane pump (Laboport, both Carl Roth GmbH + Co. KG,
150 Karlsruhe, Germany). Filtration was performed until complete clogging of the filters. For each
151 sample, the water from the two bottles was filtered separately on two glass-fiber filters (GFF,
152 pore size 2 μm). One of the filters was preserved in CTAB (A), the other filter was dried
153 immediately (B). Our intention here was to evaluate if filter preservation influences species
154 detection, since the transportation of dry filters may be considerably simpler compared to wet
155 filters. The flow-through from the two GFF filtrations was combined and filtered with a nylon filter
156 until clogging (NF, pore size 0.2 μm , Millipore, Merck, Darmstadt, Germany). This filter was
157 immediately dried after filtration (C). We intended to evaluate if the larger pore-size filters
158 sufficiently capture anuran eDNA, or finer pore filters allow for additional species detection. To
159 avoid cross-contamination we changed gloves with every new sample and disinfected hands
160 with ethanol (96%), as well we cleaned filtration unit with ethanol (96%) after each pond or site.

161 Combined visual and audio encounter surveys (VAES, Zimmerman 1994), and a tadpole
162 survey (TS, Schulze *et al.* 2015) were conducted by experienced observers (MJ and AS) to
163 generate a presence/absence matrix of species for each pond. Frogs surveys were performed
164 at night (between 21:00 and 00:30 hours) during 0.5 - 1 hrs transect walks along the ponds to
165 detect and identify specimens in vegetation, in water and on the ground around ponds.
166 Detections occurred from either visual (using flashlights and headlamps) or auditory species
167 identification. Tadpoles were sampled with dip nets once during daytime and once at night in
168 each pond as well as the riparian vegetation.

169 eDNA samples were processed in Germany in a laboratory dedicated to the pre-PCR
170 handling of environmental DNA samples. Several working routines have been implemented in
171 the laboratory to avoid contamination of samples and reactions, including spatial separation
172 from the other DNA facilities (separate room), strict decontamination protocols using UV light
173 and bleach, physical separation of extraction and PCR work spaces, automated extraction, and
174 restricted access to staff trained in the handling and analysis of forensic and environmental
175 samples.

176 DNA was extracted from GFF samples (A, B, see above) with a CTAB chloroform
177 extraction method according to Strand *et al.* (2014) and Wittwer *et al.* (2017). Dried nylon filters
178 (C) were extracted with DNeasy Blood & Tissue Kit (Qiagen, Hilden, Germany) following
179 Thomsen *et al.* (2012b). Negative controls were included in all extraction sessions (see also
180 DNA extraction in the Supplemental information). The barcode amplification targeted
181 mitochondrial 16S ribosomal DNA. We designed primers (see Primer design in Supplemental
182 information for details on design and primer tests) for a 150 bp fragment with the reference
183 database of local species (sequences from 159 specimens, Jansen *et al.* 2011; Schulze *et al.*
184 2015). The primers were tested on a subset of 12 species. PCRs were run on 96-well plates in
185 15- μ l reaction volumes with a touchdown protocol, with four technical PCR replicates per
186 sample (see PCR setup in the Supplemental information for PCR protocol and cycling
187 conditions). We included four negative PCR controls (ultra-sterile water), four extraction blanks,
188 and two positive controls on each plate. Positive controls consisted of 1) equimolar positives
189 where DNA of the twelve frog species (primer development) was pooled in equimolar
190 concentration (5 ng/ μ l), and 2) non-equimolar positives where a concentration series was used
191 by diluting the 5 ng/ μ l templates 1x to 512x dilution. PCR replicates were individually labeled
192 with multiplexing indices designed by Kozarewa and Turner (2011). PCR products were purified
193 with Agencourt AMPure XP SPRI magnetic beads (Beckman Coulter, Brea, CA), pooled in

194equal volumes, and paired-end-sequenced (2 x 150 bp) on an Illumina NextSeq 500 sequencer
195at StarSEQ GmbH (Mainz, Germany).

196 The bioinformatic pipeline for sequence processing mostly relied on OBITools v1.2.0
197(Boyer *et al.* 2016, see Sequence processing in the Supplemental information for details). We
198performed two reference-based taxonomic assignments, first with a custom database of 16S
199sequences of 159 specimens from the regional amphibian fauna (16S_custom), and second
200with all 16S sequences found in the EMBL (release 125, 16S_EMBL). Further filtering was done
201in R (R Core Team 2017) with a script supplied on GitHub:
202<https://github.com/MikiBalint/amphibian-eDNA>. This includes an ordination of all PCR replicates,
203negative and positive controls. This ordination was used to identify outlier replicates of samples
204from different ponds (Supplemental Fig. S1).

205 We followed recommendations for the statistical analyses of marker gene community
206data (Bálint *et al.* 2016). eDNA faunistic differences among the five ponds were visualized with a
207latent variable model-based ordination in R ('boral', Hui *et al.* 2015), and tested with
208multispecies generalized linear models (GLMs in 'mvabund', Wang *et al.* 2012). We used read
209abundances as input data for the boral ordination and the multispecies GLMs since presence-
210absences inferred from read abundances may overestimate the importance of rare sequence
211variants on community structure (Bálint *et al.* 2016). Different filtrates were included as distinct
212samples into the analyses since we wanted to see the variation among closely related samples
213(e.g. the three filtrates taken at the same point of a pond) *versus* the variation among distinct
214samples (e.g. originating from different ponds). We assumed a negative binomial distribution for
215both boral and mvabund analyses since read abundances are overdispersed counts, with a
216strong uniform prior on the dispersion parameter in boral (0-3). The surveyed ponds were
217markedly different in size, vegetation, water depth, etc, thus we considered pond identity as a
218good predictor of community composition. The pond effect was evaluated with analysis of

219variance (likelihood-ratio test, PIT-trap resampling, 1 000 bootstrap replicates). We performed a
220model-based ordination also for the VAES presence/absence data (since no abundance data
221was collected during the VAES), and then used a Procrustean superimposition (Peres-Neto &
222Jackson 2001) to evaluate how the VAES-based ordination of ponds matches the ordination of
223centroids of eDNA samples. We compared the efficiency of filter preservation (CTAB or dried)
224on the successful detection of species with a site occupancy model (MacKenzie *et al.* 2002;
225Bailey *et al.* 2014), implemented in the R package 'unmarked' (Fiske *et al.* 2011). We used the
226single season false-positive occupancy model developed by Miller *et al.* (2011, for details see
227Site occupancy models in Supplemental information).

228 For cost comparisons, we considered a typical eDNA survey scenario: samples are
229collected in the field and later processed in a dedicated laboratory. For the encounter-based
230survey we considered a scenario with a similar separation of the fieldwork and species
231identification: species records (audio or visual) are collected by a field biologist, and later
232identified or verified by an expert in the office/laboratory. The parameters in our cost models are
233an approximation of the variables involved in the present study and involve a learning effect in
234the efficiency of the taxonomic expert (see Cost comparison in the Supplemental information for
235details). The sampling and identification costs of VAES are dependent both on the number of
236sampling sites, and the number of species since each species needs to be recorded. During the
237eDNA survey samples are collected by a field biologist, analyzed in a lab and sequenced by an
238external provider. The costs of eDNA sampling and identification depend on the number of sites,
239but not on the number of species. We kept some cost factors constant: the costs for training the
240frog taxonomic expert and the VAES observers, the costs for building up the eDNA
241metabarcoding facilities (clean rooms and equipment to perform DNA manipulations, except
242sequencing), and the databases necessary for the sequence assignment. We assume that
243travel costs are the same for the two survey types, and that the time necessary to walk between

244frog observations and eDNA sampling points is the same. All model parameters and
245calculations are accessible on FigShare (Bálint *et al.* 2017).

246 Results

247 The sequencing resulted in 12 742 273 read pairs (deposited in ENA as PRJEB22113),
248 of which 9 479 299 were identified as complete 16S amplicons. De-replication of the reads (see
249 Supplemental Information: Sequence processing for more information on the steps) resulted in
250 631 003 unique sequences variants. Only 22 706 of these variants were represented by at least
251 110 reads and retained for further processing. The sequence cleanup resulted in 14 442 high
252 quality sequence variants, 13 497 coming from the present experiment. Of these, 4 805
253 sequence variants were taxonomically assigned by the *ecotag* command with the 16S_custom
254 database, and 8 692 with the 16S_EMBL database. The *ecotag* outputs are accessible on
255 FigShare (<https://doi.org/10.6084/m9.figshare.5099842.v5>), and can be combined and simplified
256 with the R script provided on GitHub (<https://github.com/MikiBalint/amphibian-eDNA>). The
257 assigned sequence variants represented 8 011 631 sequences. After bioinformatic processing
258 with OBITools 561 of these sequence variants were identified as ‘head’ sequences, i.e.
259 sequences that have no variants more abundant than a predefined percentage of their own
260 count, here 5% (see also Sequence processing in the Supplemental information) in at least one
261 sample (6 965 866 reads). The original read numbers were re-assigned to these head sequence
262 variants, and the read numbers were used in downstream analyses. Several of these 561 head
263 sequence variants were found also in the negative controls. After removing the reads assigned
264 to these, the final sample - sequence variant abundance matrix contained 5 815 014
265 sequences. These belonged to several groups: frogs (2 158 534), fish (1 692 613), insects (304
266 059), mammals (14 006), birds (967), and bacteria and other groups (1 063 156).

267 Read numbers assigned to frog species varied considerably among the spatial replicates
268 of the eDNA samples (see *species_abundance_matrix.csv* – zipped - on FigShare:
269 https://figshare.com/articles/_/5099842). Numbers of frog species detected by both methods,

270eDNA and VAES, varied between 3 and 18 per pond (Table1). Several of the species present
271at the ponds had no aquatic life phase during the time of survey (i.e. no larvae or adults in the
272water), and this likely resulted in lack of detection. To better evaluate the performance of the
273eDNA approach we thus assembled “reduced” VAES data sets containing only those species
274that are known to have an aquatic life phase during the survey period. For example, we
275excluded *Leptodactylus fuscus*, a species that was observed calling from the shore of the pond
276during the survey, but for which no tadpoles were recorded at that time.

277

278 In summary, in the five ponds 31 frog species were detected in total with both VAES and
279eDNA. Each of these methods detected 25 species, while the TS detected 4 species; 19
280species were detected by both eDNA survey and VAES (Fig. 2A; Table 1). Six species were
281detected only by eDNA, and six species were detected only by VAES. If we consider only
282species with aquatic life phase during the sampling (“reduced” VAES data set; N=20 for all
283ponds), of these eDNA detected 19 (Fig. 2B, Table 1, Supplemental Table S1).

284 We detected 11 species (of the twelve) from the equimolar DNA concentration positive
285controls, and 6 in the non-equimolar DNA concentration controls (Supplemental Fig. S2). Read
286numbers in the equimolar PCR controls were highly variable, but strongly correlated among the
287controls ($R > 0.7$ for each pair of positive controls, Supplemental Fig. S2A). The read numbers
288in the non-equimolar positive control were strongly linked to the DNA template concentrations of
289the PCRs (Pearson correlation coefficient $R = 0.97$, Supplemental Fig. S2B). Only
290*Leptodactylus vastus* was recovered in the field positive control (water from an aquarium with *L.*
291*vastus* tadpoles).

292 We found no difference in detection probability between the filter preservation methods
293(CTAB buffer - A: detection probability $p_1=0.278$ [0.242; 0.317]; dry - B: $p_1=0.243$ [0.209; 0.281]),

294Supplemental Fig. S3). Interestingly, subsequent filtering of water filtered through glass fiber
295filters with a 0.2 μm nylon filter - C, appeared to show an increased detection probability
296($p_1=0.327$, 95% C.I. = [0.285; 0.372]. The three approaches were equivalent with respect to
297false positive probabilities (Supplemental Fig. S3).

298 Regarding the community structure assessment based on the eDNA data, replicate
299samples of each pond grouped relatively tightly on the latent variable model ordination (Fig. 3A).
300The pond identity was a statistically significant predictor of frog communities in the five ponds
301(ANOVA, $df = 6$, $dev = 534.99$, $p < 0.01$). This is reflected in the 95% confidence intervals of the
302group centroids on the ordination which clearly separates all ponds except T1 and T3 (Fig. 3A).
303The ordination of the eDNA pond centroids closely corresponds with the ordination of
304observations from the five ponds with VAES (Procrustes permutation test, $R = 0.8$, $p = 0.03$, Fig.
3053B).

306 The cost model of VAES and eDNA showed that the starting costs (i.e. with few
307sampling sites) for VAES are relatively low, but these costs rapidly increase until the taxonomic
308expert becomes familiar with the regional frog fauna (Fig. 4). The VAES price is dependent on
309the species richness: first, the VAES observer needs to record each species on the field, and
310then the taxonomic expert needs to listen to each recording (Fig. 4). eDNA metabarcoding has a
311relatively high entry price since consumables and sequencing are costly, regardless of the
312number of sites. eDNA survey prices are then a linear function of the number of sampling sites,
313and an increase in the site numbers simply adds to sampling and consumable costs, but does
314not influence neither the time spent in the laboratory, nor the sequencing costs (Fig. 4).

315

316 Discussion

317 The one-time eDNA survey detected 25 species in the studied ponds and confirmed the
318 presence of (i) almost all species that were in contact with water during the survey time, (ii) all
319 species that had tadpoles in the ponds, (iii) more than half of all 45 species ever recorded from
320 the area, and (iv) about 65% of the 35 species ever recorded in the five surveyed ponds. The
321 eDNA dataset recorded clear differences among the frog communities in the surveyed ponds, a
322 result also in accordance with the VAES data. The comparison of VAES and eDNA cost models
323 show that eDNA biodiversity surveys may be a cost-efficient alternative to VAES in species-rich
324 areas, but not necessarily in areas with low species numbers. Specific recommendations and
325 technical remarks are presented in the Supplemental information.

326 eDNA approaches for routine use in tropical regions will benefit from the implementation
327 of straightforward and robust sampling techniques. The comparison of the filtration-preservation
328 approaches shows that filters can be dried in the field before being sent to a lab. Currently only
329 few comparative studies exist regarding the preservation of eDNA filtrates on filters (e.g., Hinlo
330 *et al.*, 2017; Spens *et al.*, 2017). Hinlo *et al.* (2017) showed that the simple refrigeration of filters
331 may be preferred to frozen storage. Here we show that filters can be dried in the field: this
332 simplifies transportation and storage since no special precautions are needed, unlike for liquid
333 handling. Interestingly, small pore-size nylon filters seemed to recover a high proportion of
334 target DNA that had previously passed through a 2 μm mesh sized glass fibre filter. While this
335 result may be caused by the different extraction methods used, we consider it likely that a large
336 proportion of DNA occurred in extracellular form. This result suggests that alternative filtering
337 methods based on fine-scale mesh sized filtration with limited filtering volumes may further
338 increase success rates in eDNA studies on amphibian communities. By limited volumes we do
339 not suggest sampling only a few hundred ml of water, but rather to sample enough from a

340waterbody to ensure representativeness, and then filter from this well-mixed sample until the
341filter clogs.

342 During the one-time surveys, six species were only detected with eDNA, and not with
343VAES (Fig. 2A). Four of these species (*Dermatonotus muelleri*, *Leptodactylus elenae*,
344*Osteocephalus taurinus*, *Rhinella schneideri*) are quite common in the area. The detection of
345these four species by eDNA but not by VAES may result from the low abundance of adults or
346tadpoles in the ponds, or a lack in acoustic activity. In some cases eDNA detections are even
347likely based on single tadpoles or single adults only. For example, one adult *L. syphax* was
348heard calling from a crevice of a rocky outcrop, its usual habitat (de Sá *et al.* 2014),
349approximately 30 m away from T2, but detection by eDNA most probably was due to the
350presence of tadpoles in pond T2. All other species that had tadpoles in the ponds were also
351detected by eDNA (Table 1). Furthermore, our results suggest that detection of rare or solitary
352species by eDNA is possible, since the presence of *L. vastus* in the eDNA most possibly can be
353traced back to single, territorial adult males that were present during VAES of T4 and T5. These
354examples show the great potential of eDNA for detection of elusive life stages like tadpoles and
355less abundant species. The remaining two species detected only with eDNA are not known to
356occur in the area, thus they are candidates for false assignments. The low read numbers
357assigned to these species also supports this (Supplemental Table S1). Both of these were
358assigned with sequence data from EMBL (*L. latinasus*: KM091595, *L. laevis*: AY843696).
359*Lysapsus laevis* would be the first record of this genus in the study area, but the identification of
360*Lysapsus* populations only using short 16S eDNA sequences is questionable, especially when
361considering the unclear taxonomy of the group in Bolivia. Nevertheless, based on their
362distribution and ecology all of the four known species of *Lysapsus* (*L. boliviana*, *L. caraya*, *L.*
363*laevis*, *L. limellum*) may occur at the site (De la Riva *et al.* 2000; Lavilla *et al.* 2004; Reichle
3642004a; b; Angulo 2008; Jansen *et al.* 2011; Frost 2016). They may have remained undetected

365 thus far simply because of their small size and rather inconspicuous advertisement call.

366 *Leptodactylus latinasus* is another species possibly occurring in the area that had so far never
367 been recorded (de Sá *et al.* 2014) and we cannot exclude the possibility that the corresponding
368 sequence variant is actually an erroneous variant of one of the six local *Leptodactylus* species.

369 Species' ecology might explain why some species were detected only with VAES, but
370 not by eDNA (Fig. 2A, Supplemental Table S1). None of these species are strictly bound to
371 ponds in the life stages occurring during our sampling: *Dendropsophus arndti* and *D. salli*,
372 usually call from plants on the pond shores (Schulze *et al.* 2009) and have only sporadic contact
373 with water. *Leptodactylus fuscus* and *Pseudopaludicola* sp. also do not enter the water but
374 usually call from nearby muddy grounds or grasslands. We did not detect tadpoles of these
375 species in the ponds: the tadpoles of these species develop during the rainy season which
376 triggers reproduction, but our sampling slightly preceded the rainy season. Some of the other
377 undetected species have terrestrial or quasi-terrestrial life histories: *L. fuscus* deposits eggs
378 within foam nests in underground burrows at some distance from ponds and these are washed
379 into nearby water bodies by floods that follow heavy rains (Heyer 1978; Lucas *et al.* 2008).
380 *Boana geographica* was likely missed as a result of PCR bias: we could never recover it from
381 positive controls when DNA from other species was also present (Supplemental Fig. S2),
382 although single-template PCR reactions worked. *H. geographicus* was also not detected by
383 VAES, so it is possible that the species was not present during the survey. If we consider the
384 specific life histories, the one-time eDNA survey only missed a single species
385 (*Sphaenorhynchus lacteus*) in the area which was found calling from the water during the VAES
386 (Fig. 2B, Supplemental Table S1). Regarding the whole local species pool (45 species in 10
387 years), some of the species (10) may not be detectable by eDNA since they reproduce outside
388 of water (e.g. *Leptodactylus mystacinus*), or are forest dwellers (e.g. *Leptodactylus* cf. *didymus*,
389 Fig. 2D, Table S2). Overall, the results suggests that eDNA performed similarly well in detecting

390species as VAES, but it was not successful to identify all species occurring at the specific
391ponds. Differences among individual ponds may result from the differential observational biases
392of eDNA and VAES approaches, none of which is providing a “true” list of species (hence the
393long-standing need to model species occupancy). eDNA methods seem to be a powerful tool for
394the detection of elusive life stages and less abundant species in tropical communities: with a
395single sampling eDNA detected more than half of the 45 species known to be present in the
396area, and about 65% of the detectable species from the area (23 out of 35).

397 The sampling was done at the beginning of the rainy season when only few species
398reproduced in the ponds. Repeating the sampling during the rainy season could have increased
399species detections by eDNA. The results show the importance of life history in the
400metabarcoding-based survey of tropical frogs and emphasize that sampling at multiple time
401points may be essential for more complete species pools also with eDNA (see
402Recommendations and technical remarks in the Supplemental information). Comprehensive
403and taxonomically sound sequence databases are essential for eDNA metabarcoding studies:
404we had a database that contained all 45 species that were ever recorded in the area. This
405database was essential for both initial primer development, and taxonomic assignments: indeed,
406both eDNA-recorded taxa that were not present on the complete local species list of the present
407study (Fig. 2, Supplemental Table S2) were identified in the EMBL-based assignment that
408followed the assignment with the custom local 16S database. The importance of sequence
409assignment databases is long recognized, with considerable efforts underway to establish them
410(Ratnasingham & Hebert 2007; Coissac *et al.* 2016). There is an urgent need of further DNA
411sampling to create regional reference data banks, but only a few studies aim at completing DNA
412sampling of local anurofaunas in South America (e.g. Jansen *et al.* 2011; Guarnizo *et al.* 2015;
413Moraes *et al.* 2017). However, a preferably complete geographical and taxonomic coverage is
414necessary for continent-wide eDNA-based frog inventories. Increased DNA sampling will also

415increase knowledge about actual distributions and taxonomy of Neotropical frogs (including
416cryptic lineages). This will improve IUCN evaluations, which currently clearly lack distributional
417information (Supplemental Table S1).

418 There was considerable variation in the species recorded with the spatial replicates of
419the eDNA samples, and this underlines that eDNA sampling should be replicated for a good
420representation of community composition. These samples may be pooled before DNA extraction
421to optimize costs if site-level variation is not of interest. Interestingly, community structures
422across ponds inferred from both eDNA and VAES datasets were highly similar (Fig. 3). The
423eDNA data did not distinguish the communities from ponds T1 and T3, and these communities
424also grouped closely in the VAES results. Although the comparison of frog community structures
425among the assessed ponds may be confounded by the different sample sizes (with the eDNA
426ordination based on many spatial replicates and the VAES ordination on a single observation
427event per pond), and the eDNA ordinations are based on read-abundances in contrast to the
428VAES presence-absence ordinations, the correspondence of the results is still striking. Similar
429results were found in insect (Ji *et al.* 2013; Elbrecht *et al.* 2017) and plant communities
430(Niemeyer *et al.* 2017) where eDNA-based and morphology-based identifications result in
431similar conclusions about community structure, despite markedly different species lists. Our
432results provide further evidence that eDNA-based biodiversity surveys are highly sensitive to
433differences among ecological communities. These inferences are comparable to those derived
434with encounter-based observations and are informative about processes that underlay
435community assembly.

436 Although central to deciding on a method for biodiversity surveys, cost comparisons are
437not straightforward since they need to be based on expert knowledge both in VAES and eDNA.
438Cost comparisons were performed for single species eDNA detection (Huver *et al.* 2015; Davy
439*et al.* 2015; Smart *et al.* 2016), but we are not aware of frameworks suitable for eDNA

440metabarcoding. Here the VAES cost estimation is informed by over a decade of field and
441integrative taxonomic work with tropical frogs (Jansen *et al.* 2007, 2016; Schulze *et al.* 2009;
442Brusquetti *et al.* 2014), while the eDNA part is only informed by a few years of eDNA
443biodiversity surveys (Bálint *et al.* 2016; Vörös *et al.* 2017). Such comparisons are urgently
444needed due to stakeholder demands in eDNA (e.g. governmental agencies, conservation
445NGOs, fisheries, etc.).

446 The two species richness scenarios we defined for cost comparisons (Fig. 4) shows that
447VAES costs become high in regions with high frog richness since they are a function of both the
448number of sites and the number of species. Biodiversity surveys with eDNA are not necessarily
449cheaper in low-richness regions since entry costs for the eDNA work are high: lab consumables
450and sequencing are costly. However, eDNA survey costs are not dependent on the local
451biodiversity since metabarcoding can consider thousands of species simultaneously in a sample
452(Taberlet *et al.* 2012b). Consequently, eDNA costs are function of sample numbers, which
453influence the collection time spent on the field and consumables. eDNA metabarcoding
454operations are easily scaled up in a sense that hundreds of samples can be simultaneously
455processed (Ficetola *et al.* 2015).

456 Several aspects of our cost models are contentious. One issue is whether the relatively
457untrained VAES observers, or taxonomic experts perform the fieldwork, since taxonomic experts
458may identify many of the targeted frog species immediately on the field. Currently, most surveys
459of high-diversity areas are directly done by experts interested in the local fauna, but we argue
460that this will not work for continental - global biodiversity surveys simply because there are not
461enough taxonomic experts (Buyck 1999; Haas & Häuser 2003). We also did not consider a
462scenario when VAES surveys are performed with automated recording devices (ARDs), and
463sounds are automatically identified by algorithms (see Recommendations and technical remarks
464in the Supplemental Information). The sound complexity in tropical environments currently

465prohibits the use of automated sound identifications (Campos-Cerqueira & Aide 2016). It is also
466difficult to compare fundamental infrastructure and training costs (i.e. the establishment of an
467eDNA laboratory versus the training of taxonomic experts). Discussions about cost models are
468timely since they will play major roles in devising much needed continental and global
469biodiversity surveys (Tittensor *et al.* 2014).

470 Conclusions

471 eDNA seems to be suitable to standardized biodiversity surveys of frogs even in
472species-rich areas, but it may be overly costly for smaller studies in low-richness regions.
473Differences between eDNA and traditional surveys seem to result largely from different
474observational biases. Consideration of life histories promises to improve comprehensiveness of
475both types of surveys and thus also their correspondence. The eDNA data is certainly suitable
476to characterize not only community composition, but also the factors that shape communities:
477this gives an unprecedented opportunity to incorporate eDNA as a standard toolkit for
478community ecology and macroecology. Assessing community structure with eDNA-based
479community data foresees global biodiversity surveys and monitoring that will support both
480biodiversity research, and informed decisions on sustainable use of biological diversity.

481

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739 **Data accessibility**

740 DNA sequences: EMBL accession PRJEB22113

741 Cost model parameters and calculations: FigShare,

742 <https://doi.org/10.6084/m9.figshare.5099842.v5>

743 Input data for statistical analyses: FigShare,

744 <https://doi.org/10.6084/m9.figshare.5099842.v5>.

745 R-script for bioinformatic filtering and statistical analyses:

746 <https://doi.org/10.5281/zenodo.1294092>

747

748 **Author contributions**

749 M.B., M.J., C.N., C.W. designed research; J.L.A., M.B., M.J., O.M., A.S., C.W., B.C.

750performed research; B.C., T.C., C.N., S.U.P. contributed reagents or analytic tools; M.B., M.J.

751analysed data; M.B., T.C., M.J., O.M., C.N., S.U.P., wrote the manuscript.

752

753 Tables

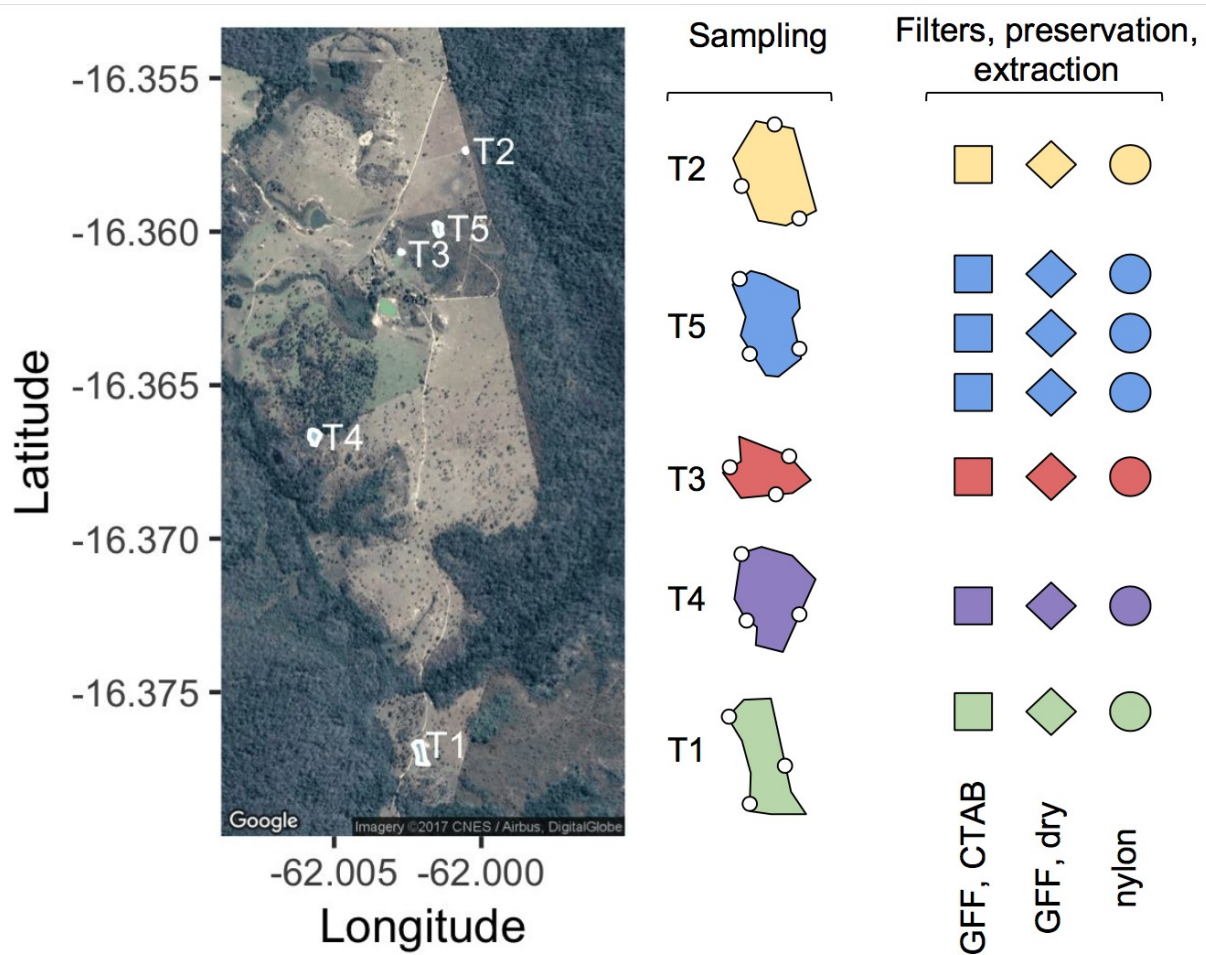
754 Table 1.

755 Numbers of detected species by method per pond (VAES = Visual-audio encounter
756 surveys. Reduced VAES = only considering species that were in contact with water during
757 survey).

Pond	eDNA	VAES	Reduced VAES	Tadpole survey
T1	10	8	6	4
T2	7	4	4	3
T3	3	7	3	0
T4	9	15	13	1
T5	13	18	14	1

758

759 Figures



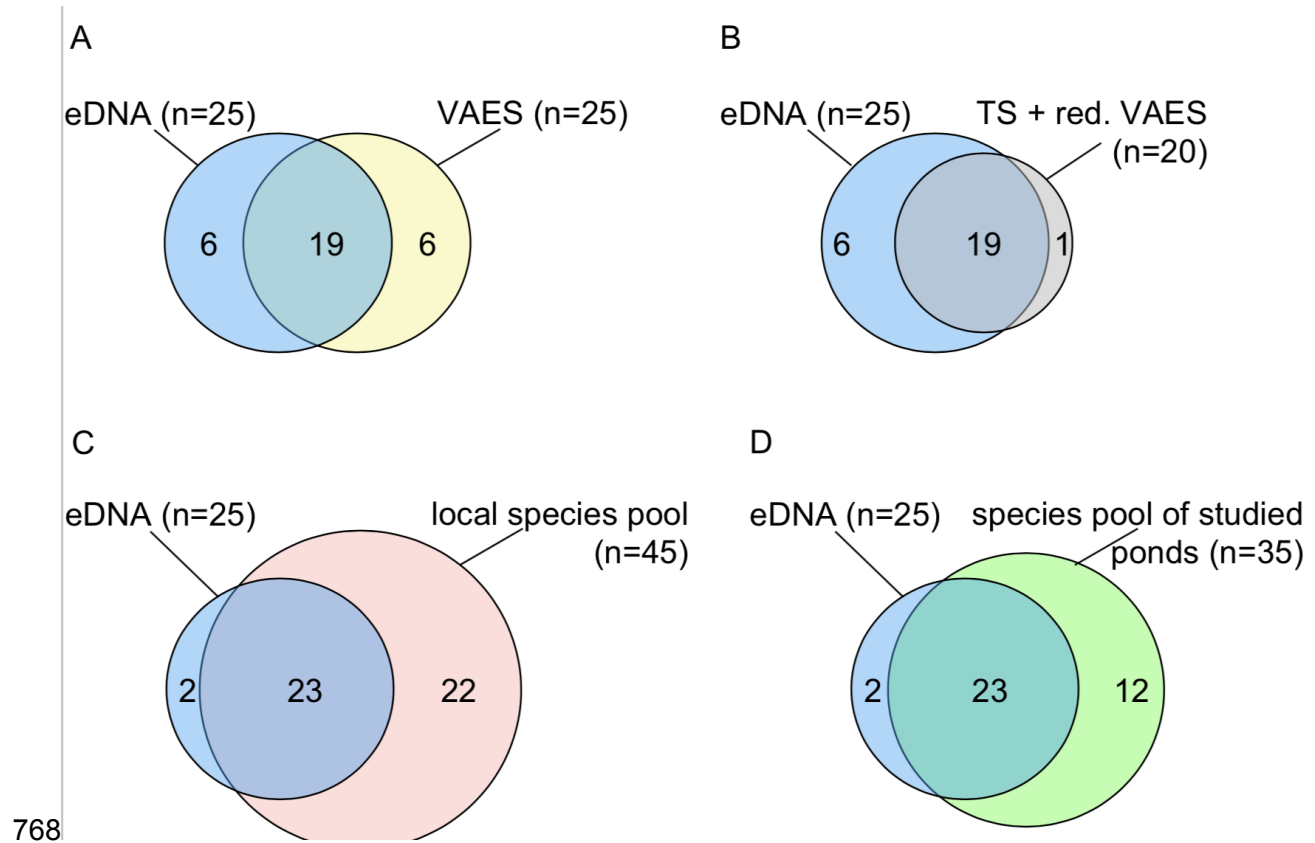
760

761 Fig. 1.

762 Sampled ponds, sampling replication, and filtration - filter preservation strategy. GFF -
763 glass-fiber filter, 2 μm ; nylon - nylon filter, 0.2 μm . Each sample (three per pond, marked with
764 small circles) were processed with each three filtration strategies. Multiple samples of pond T5
765 were three times processed with each filtration strategy.

766

767



768

769 **Fig. 2.**

770 Comparison of species lists generated by eDNA (blue) and A) visual and audio

771 encounter surveys (VAES, yellow), B) tadpole survey (TS) plus reduced VAES - only species

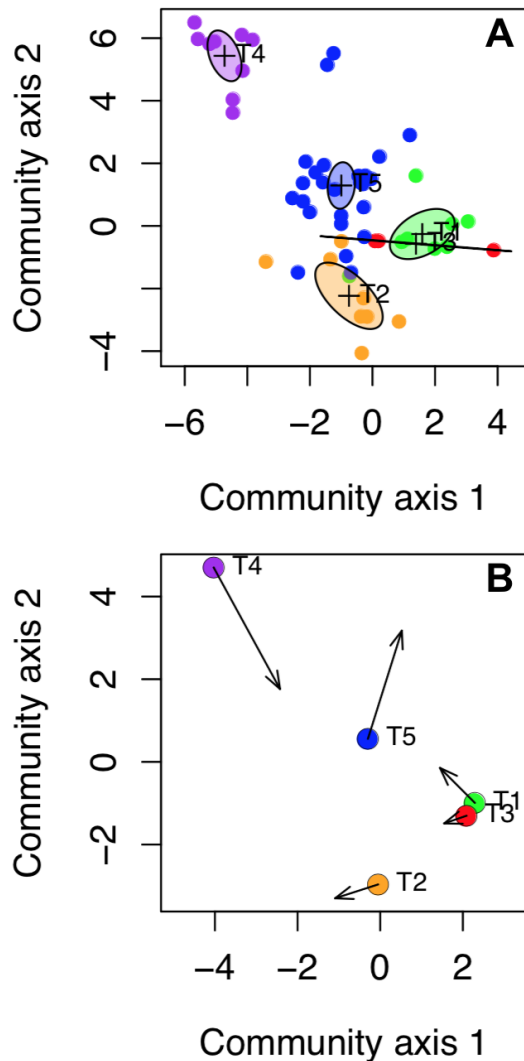
772 with high eDNA detectability in ponds (see text for details; grey), C) complete local species pool

773 (red), and D) species pool of studied ponds based on long-term monitoring (green).

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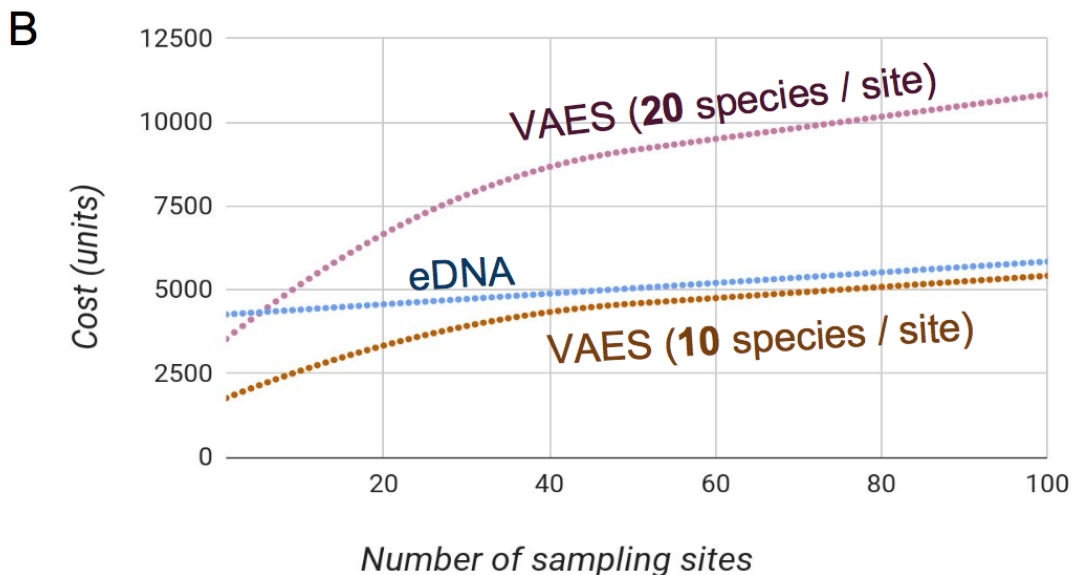
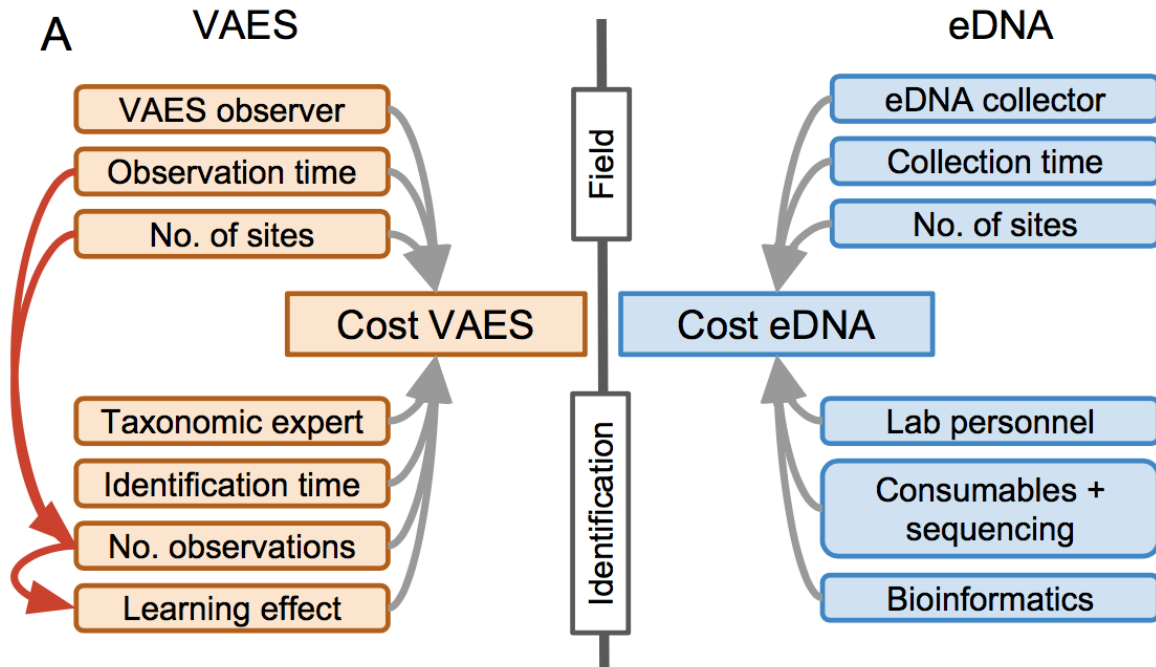


777

778 Fig. 3.

779 Comparison of ecological signal between visual and audio encounter surveys (VAES) and
780 eDNA metabarcoding surveys. A - Latent variable model ordination of the pond samples
781 according to the read numbers of species. Water samples from each pond were taken at three
782 locations. For ponds T1-T4, three samples at each location were filtered through two GFF (A,B),
783 and one nylon filter (C), resulting in nine samples per pond. For pond T5, each filtration was
784 done three times. Not all samples contained reads after the bioinformatic quality filtering and
785 these samples are not shown on the ordination (see e.g. T3 - red). The ellipses represent 95%
786 confidence intervals for the standard errors of the pond centroids (marked with +). B - Non-

787metric multidimensional scaling plot of the five ponds (Jaccard distance), according to VAES
788species presence-absences. The arrows represent the Procrustes rotation of the VAES pond
789ordination and they target the group centroids of the latent variable model ordination.



790

791 **Fig. 4.**

792 Cost comparisons between visual and audio encounter surveys (VAES) and eDNA
793 metabarcoding surveys. A - Schematic overview of the visual and audio encounter survey
794 (VAES) and eDNA cost models. Model parameters and calculations are accessible through
795 FigShare (Balint_et_al_survey_cost_calculations.xlsx,

796<https://doi.org/10.6084/m9.figshare.5099842.v5>). B - Cost comparisons of frog diversity surveys
797with eDNA, and two VAES scenarios (a low local species richness scenario - 10 species per
798site, and a medium-high species richness scenario - 20 species per site).