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Comparison of eDNA and visual surveys for rare and cryptic bromeliad-dwelling frogs

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Surveys of rare or cryptic species may miss individuals or populations that are actually present. Despite the increasing use of environmental DNA (eDNA) analysis to survey species in ponds, rivers, and lakes, very few studies have attempted to use eDNA for the detection of species using very small water bodies such as those accumulated within plants. Our aim was to investigate the feasibility of an eDNA sampling method for detecting Crossodactylodes itambe, an endemic bromeliad-dwelling frog from a remote location in Brazil. We collected water samples from 19 bromeliads for which we had observational data from direct visual surveys. We compared occupancy estimated from direct observations with the results from quantitative real-time PCR based eDNA assays. For observational surveys, we used a single season occupancy model. We applied a novel Bayesian occupancy model to estimate occupancy from eDNA samples, as well as false positives and false negatives at different stages of the workflow. eDNA from bromeliad tanks provided reliable estimates, with very low error levels and improved detection when compared to detectability from direct observation. Estimated occupancies using eDNA and visual survey methods were similar. The method is feasible for species restricted to small water bodies and exposed to direct UV radiation, and particularly useful to survey remote locations and confirm species presence. eDNA analysis provides a viable alternative to destructive sampling of bromeliads or direct observation methods that require logistically challenging repeated observations. Therefore, eDNA methods may be widely applicable to sampling programmes of other amphibians that live in plants.

Keywords: bromeliad, eDNA detection, false-positive, amphibian, occupancy, phytotelm

INTRODUCTION

Species surveys using direct observations suffer from the problem of individuals or populations being missed (MacKenzie et al., 2002). Such imperfect detection is caused by a wide variety of factors, including time of survey and temperature (Sewell et al., 2010), observer experience (Grant et al., 2005; Fitzpatrick et al., 2009) or simply individuals being obscured from view. When making observations, the presence of a surveyor can alter the behaviour of the target organism, reducing the likelihood of it being observed (Barata et al., 2017, 2018a). To account for imperfect detection, repeated visits are required to control for variation in detectability (MacKenzie et al., 2002). However, detection does not necessarily require direct observation. An increasing number of indirect survey methods are emerging such as environmental DNA (eDNA) analysis. Indeed, eDNA surveys can outperform direct observation surveys (Lopes et al., 2017; Burns et al., 2020), but this varies according to the ecological characteristics of the targeted organism (Takahara et al., 2019).

Surveys targeting eDNA involve the collection of environmental samples from a location which usually comprises water, soil or sediments (Turner et al., 2015; Buxton et al., 2018; Spitzen-van der Sluijs et al., 2020; Valentin et al., 2020). These samples are then processed following forensic protocols for the extraction of DNA that has been released by organisms into the environment. Where possible, the DNA that is extracted and amplified from these environmental samples is then identified to species level by comparing their sequences to a reference DNA library, thereby allowing for inferences concerning species presence in that habitat (Jane et al., 2015). Detection of species using eDNA methods has become commonplace in environments where detectability of a target species may be relatively low, such as ponds (Harper et al., 2018), lakes, rivers, and streams (Sales et al., 2019; Bedwell et al., 2020). However, very few studies have attempted to use eDNA methods for the detection of species using very small bodies of water such as aggregations of water collected within plants (also known as a phytotelm), e.g. within the tanks of bromeliads. Only two previous studies have used eDNA
methods in the survey of bromeliad tanks (Brozio et al., 2017; Torresdal et al., 2017), both of which targeted the Critically Endangered Trinidad golden treefrog (Phytotriades auratus), an elusive species that requires destructive sampling (i.e., bromeliad destruction).

Crossodactylodes itambe is a small frog species endemic to the summit of one mountain, the Itambe summit, Minas Gerais state, in south-eastern Brazil (18°23'54"S, 43°20'W; datum WGS84). The species exclusively lives within ground bromeliads (Vriesea medusa), which are found on high elevation rocky outcrops (Barata et al., 2013). Crossodactylodes itambe is restricted to bromeliads at 1800 m above sea level or higher with a total area of less than 0.5 km², although the plants can be found at lower altitudes (Santos et al., 2017). Species occupancy increases at higher elevation (Barata et al., 2017) and abundance of individuals is related to bromeliad structure, such as plant size and the volume of water retained by the central tank (Barata et al., 2018b). The restricted range and habitat requirements for the species make it highly vulnerable to extinction from climate change, wildfires or disease, and therefore a priority for conservation monitoring.

Due to the remote location, nocturnal activity of the species (Barata et al., 2018a) and a detection probability of 0.40-0.65 (Barata et al., 2017), visual surveys are labour intensive and costly. Power analysis conducted by Barata et al. (2017) suggests that when using visual encounters, an observer would be required to make at least three to four visits and 143 bromeliads would need to be surveyed to have an 80% chance of detecting a change of 30% in the population. Furthermore, when searching for new populations of rare and cryptic species, only a subset of the potentially highly suitable areas can be surveyed because of logistical and financial constraints (I.M. Barata, unpublished data). Consequently, it would be highly beneficial to develop a passive survey method with an equivalent or greater detectability from a single site visit than visual encounters. A passive method would provide substantial savings in terms of logistical and survey efforts, without compromising the ability to detect population changes within this highly vulnerable population or finding new populations at surrounding locations.

Here we develop species-specific PCR primers for C. itambe and test the practicalities of collecting eDNA samples from ground bromeliads in a remote location. Bromeliads in this location grow in an outcrop among shrub and herbs and are exposed to direct UV radiation, which can potentially increase DNA degradation rates (Strickler et al., 2015). We compare the known occupancy estimated from observational surveys (from multiple visits in 2014-17) with the results from quantitative real-time PCR (qPCR) based eDNA assays. Our main goal was to test the feasibility of a passive sampling method in detecting C. itambe in an exposed montane area with the challenges of high UV radiation, with a view to applying the method more widely to other amphibian species that are restricted to bromeliads.

**SITE SELECTION**

A tank bromeliad is a type of phytotelm that accumulates rainwater at the base of each leaf axil and in the central tank, thereby providing a microhabitat for other species (Lehtinen, 2004). We selected 19 bromeliads for eDNA sample collection, 11 of which were known to have been occupied by C. itambe based on previous studies and detection histories (Barata et al., 2017). The remaining eight bromeliads had no C. itambe recorded within them over the previous four years and were therefore considered to be unoccupied by frogs. All samples from bromeliads were from within the known range of the frog, on the Itambe summit, Minas Gerais.

Additionally, two samples were collected from flowing water in the vicinity of the study site as field negative samples. These were collected to ensure no contamination occurred in the field, as this is a potential risk when sampling in remote locations. Whilst we acknowledge that filtering sterile water on site as a field negative is a more standard approach, it was decided not to increase the volume and weight of materials needed to be carried due to the logistics of accessing the remote location. As flowing water is not used by the target species, it was highly unlikely to contain target DNA. If these results returned negative, we could be confident there was no in-field contamination.

**VISUAL ENCOUNTER SURVEYS**

Observational surveys were conducted across four years, from 2014 to 2017, during the rainy season (between October to March). Bromeliads were tagged with individual numbers that allowed repeated visits in multiple years. For each year, visual encounter surveys consisted of three to six consecutive visits to the same site to create a detection history of presence (1) and absence (0). Visual surveys were conducted at night by a team of two people, with only the most experienced observer taking notes on species presence/absence to avoid observational bias (Barata et al., 2017). Bromeliad selection for eDNA samples was based on previously known detection histories between 2014 and 2016 and an additional survey in 2017 during four consecutive nights. Surveys and detection histories from previous years provided estimates of species occupancy and detection (Barata et al., 2017). The surveys in 2017 confirmed the same species presence/absence pattern observed in previous studies and allowed the collection of water samples from selected bromeliads.

**ENVIRONMENTAL DNA SAMPLE COLLECTION**

Samples were collected using syringe filtration and 0.22 µm (PVDF membrane type, gamma irradiated) MilliporeTM SterivexTM filter capsules. To prevent contamination, we prepared individual sterile sample collection kits to be used at each bromeliad before conducting fieldwork. Each kit contained two pairs of gloves, a 0.22 µm filter capsule, a 60 mL luer-lock syringe, a 30 mL container filled with absolute ethanol, a 10 mL syringe with 1/2" needle, two luer-lock caps, a 50 mL centrifuge tube and a small zip-lock bag (Fig. 1A). In the
field, water was drawn from either the centre or lateral leaves of the bromeliad using the sterile 60 mL luer-lock syringe (Fig. 1B); the filter capsule was attached to this syringe which was used to push the water across the filter membrane (Fig. 1C). This procedure was repeated twice to filter a total of 120 mL of water. Once the whole water sample was filtered or the capsule had become blocked (which was the case for one sample), 10 mL of absolute ethanol was added to each filter as a preservative using the 10 mL syringe and ½” needle. Each capsule was sealed with luer-lock caps, re-sealed in an individual 50 mL centrifuge tube and stored in a sample bag to prevent contamination while in storage and transport (Fig. 1D). We numbered each sample (filters, containers and bags) with the reference number of the bromeliad from individual tags.

**eDNA extraction**

DNA extraction was undertaken in a dedicated lab, within a UV hood. All equipment and work stations were sterilised using a combination of 10 % bleach solution and/or UV light in advance of use. Standard laboratory protective equipment was worn at all times. DNA extraction followed a modified Qiagen® DNeasy® blood and tissue kit protocol, adapted from Spens et al. (2016). The ethanol preservative was removed from the MilliporeTM SterivexTM filter capsule by attaching a sterile syringe and passing air across the capsule, collecting the liquid in a 2 mL micro-centrifuge tube. 50 µL of 3 M sodium acetate solution per milliliter of ethanol recovered was added, these samples were then incubated at -80 °C for 10 minutes to aid in DNA precipitation before being centrifuged at 14000 RPM for 15 minutes to collect any precipitate as a pellet on the side of the tube with supernatant discarded. 180 µL of ATL buffer and 20 µL of proteinase K from the extraction kit was added to each micro-centrifuge tube, which was then vortexed for 15 seconds to suspend the pellet and mix. Samples were incubated on a rotating block at 56 °C overnight. AL buffer and ice-cold absolute ethanol was then added to each micro-centrifuge tube in a 1:1:1 ratio with the incubated contents of the tube.

720 µL ATL buffer and 80 µL of proteinase K from the extraction kit were added directly into each of the filter capsules, the caps replaced, and filter units sealed with Parafilm®, and incubated at 56 °C overnight on a rotating block. The liquid was removed from the filter using a sterile syringe and by passing air through the capsule, collecting the buffer in a fresh micro-centrifuge tubes. AL buffer from the extraction kit and ice-cold ethanol were then added in a 1:1:1 ratio (samples were split across two tubes per capsule to accommodate the volume). Product extracted from the ethanol preservation buffer and the filter capsule for each sample were then combined, passing both across the same mini-spin column from the extraction kit. Extraction continued as per extraction kit manufacturer’s protocol eluting into 200 µL of warm AE buffer.

**Primer development**

Primers suitable for use with eDNA were developed to amplify a short region of the Cytochrome Oxidase 1 gene (COI) of *C. itambe*, based on the sequence identified by Santos et al. (2017); NCBI accession number KY362551. Primers were designed using the program Primer 3 (Koressaar & Remm, 2007; Untergasser et al., 2012; Koressaar et al., 2018). Conditions were set to identify primers between 18 and 23 base pairs in length to amplify a region between 75 and 100 base pairs long. We specified that no runs of greater than three base pairs should be included, with a GC content of between 40 and 50 %, and an optimum melt temperature of 60 °C. A set of primers was identified to amplify an 83 base pair sequence (Table 1). The primer sequences were tested in silico with a NCBI blast search to check for cross amplification with other species. No species were found to have a 85 % or greater match to either the forward or reverse primer. Additionally, whilst other

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**Figure 1.** Environmental DNA sample collection demonstration: A) sample collection kit; B) water collection from lateral leaves of the bromeliad using a sterile 60 mL syringe; C) pushing the water across the 0.22 µm filter capsule; and D) individual capsule sealed in a tube and stored in a sample bag to prevent contamination.
frogs (*Bokermannohyla nanuzae*) were occasionally seen using the bromeliads in the study area over this 4-year period (Barata I.M., personal observation), no other amphibian species using the bromeliads were recorded during the observational surveys (i.e., the night before eDNA samples were taken). Primers could not be tested in vitro due to the difficulties in obtaining and exporting tissue samples of the relevant species.

**Table 1.** Primer sequences generated using Primer 3 for detection of *C. itambe* from eDNA.

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Length</th>
<th>Melting temperature (GC%)</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>CICO1-F</td>
<td>20</td>
<td>59.78 / 55</td>
<td>taccagctggcagtta</td>
</tr>
<tr>
<td>CICO1-R</td>
<td>20</td>
<td>59.59 / 55</td>
<td>ggcatggcctaagt-taccag</td>
</tr>
</tbody>
</table>

**qPCR**

qPCR was conducted in a separate room to DNA extraction. All equipment and work surfaces were sterilised using either a 10% bleach solution and/or UV light in advance of use and appropriate personal protective equipment was worn. Plate set-up was conducted in a UV hood dedicated to low concentration DNA work. qPCR was performed using a SYBR Green assay, with eight replicates per sample. The qPCR amplification procedure consisted of 10 µL of Applied Biosystems™ PowerUp™ SYBR™ Green Master Mix, 2 µL of each primer at a concentration of 10 µM/µL, and 4 µL of template DNA, in a final reaction volume of 20 µL. qPCR conditions consisted of two activation steps at 50 °C for 2 minutes followed by 95 °C for 2 minutes, then 40 cycles of 95 °C for 15 seconds, 59 °C for 15 seconds and 72 °C for 1 minute. A melt curve was then performed ramping up from 55 °C to 95 °C in 0.5 °C increments. Three negative control samples were included in each qPCR run, to check for contamination. A replicate was classed as positive when an exponential growth phase in relative fluorescence was identified and the melt curve indicated a temperature of between 82 °C and 82.5 °C, indicating the fluorescence was not caused by primer dimer.

Following qPCR analysis, each sample was checked for inhibition by adding a known quantity of non-target DNA to each sample. The assay consisted of 10 µL of Applied Biosystems™ PowerUp™ SYBR™ Green Master Mix, 4 µL of eDNA sample, 2 µL of a tissue extract from great crested newts (*Triturus cristatus*), 2 µL of forward and reverse primers for our targeted species (*C. itambe*), as well as the forward and reverse primers for great crested newts (TCCBL and TCCBR; Thomsen et al., 2012) at a concentration of 10 µM /µL, in a final reaction volume of 20 µL. The qPCR and melt curve conditions replicated those stated above, with two negative control samples included in each qPCR run. Samples failing to amplify our target species’ DNA or amplifying more than 1 cycle later than control samples were considered to contain PCR inhibitors.

**Data analysis for species detection**

We used a single-season occupancy model (MacKenzie et al., 2002) to estimate detection and occupancy rates from the observational data collected in the field. We used previously published detection histories (Barata et al., 2017) and included only bromeliads for which we had eDNA samples. Although occupancy models can accommodate covariates that explain both parameters (MacKenzie et al., 2002), we opted to run a null model (i.e., with no covariates). As our aim was to obtain overall estimates for comparisons without exploring covariates; given the small size of our dataset, we wanted to avoid overparameterisation of the model. We controlled for variation in detection in our data by using observations from a single experienced observer (Barata et al., 2017). We used a free online tool to analyse qPCR based eDNA data to generate occupancy and detectability information (https://blogs.kent.ac.uk/edna). This tool is based on Griffin et al. (2020) and uses a Bayesian framework to identify: the probability of species occupancy; stage 1 (the sample collection phase) true and false positive rates; and stage 2 (the laboratory phase) true and false positive rates. Stage 1 true positive rate (θ11) is the probability that a water sample collected from an occupied site includes DNA of the target species, with stage 1 false negative rate being the complement of this. Stage 1 false positive rate (θ10) is the probability that a water sample collected from an unoccupied site includes DNA of the target species. Stage 2 true positive rate (p11) is that an individual PCR replicate of a sample containing target DNA is positive, with stage 2 false negative being the complement of this. Stage 2 false positive rate (p10) is the probability that an individual PCR replicate of a sample that does not include DNA of the target species returns amplification. These differ from standard observational occupancy models as eDNA sampling is a two-phase process with potential for error to be introduced at both sample collection and sample analysis phases. Conversely, direct observation has a single phase where the species is either observed or not observed.

**RESULTS**

No target DNA was amplified from field or laboratory negative control samples by 40 qPCR cycles, and no sample demonstrated characteristics that indicated PCR inhibitors were present. All 11 samples collected from bromeliads with known species presence amplified target DNA, of which 10 showed amplification in all eight qPCR replicates, with the remaining sample showing amplification in seven of the eight qPCR replicates (Table 2). Additionally, one sample from a bromeliad with no known occupancy showed amplification in a single qPCR replicate (bromeliad 8112; Table 2). One sample from a previously unoccupied bromeliad had amplification in five of the eight replicates (bromeliad 7007; Table 2). However, this filter was damaged and leaked during transportation, possibly leading to contamination and a false positive. Therefore, we excluded this result from any further data analysis. These two eDNA positive but
observationally negative bromeliads were both found within the existing known range of the species. Overall, we had a naïve occupancy rate of 66.7% using the eDNA method (12 positives out of 18 samples), compared to 61.1% for direct observational surveys (11 positives out of 18 surveyed sites).

From direct observational survey data, the occupancy rate was estimated to be 0.61 and detectability was 0.77 (Table 3). From eDNA analysis, a bromeliad occupancy rate of 0.61 was estimated. A false positive rate at sample collection (θ10) of 0.04 was found, with a true positive rate at sample collection (θ11) of 0.97, equivalent to a 3% false negative rate (Table 3). This compared to the false positive rate at the laboratory analysis stage (p10) of 0.02 and true positive rate at the laboratory analysis stage (p11) of 0.98, equivalent to a 2% false negative rate (Table 3). The conditional probability of detection analysis given by the number of amplified qPCR replicates (Fig. 2) shows the probability that an occupied site is wrongly classified as unoccupied in relation to the

Table 2. Water samples collected in the field for 19 bromeliads and two field negatives (FN) with sample number, characteristics of bromeliads (elevation given in metres above sea level, size given as bromeliad height in centimetres), and results from direct observations detection history (0 = absence and 1 = presence) and environmental DNA (eDNA) analysis (P = positive and N = Negative).

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Bromeliad</th>
<th>Direct observation</th>
<th>eDNA</th>
<th>eDNA positive replicates</th>
<th>Signs of inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Elevation</td>
<td>Size</td>
<td>2014</td>
<td>2015</td>
<td>2016</td>
</tr>
<tr>
<td>8131</td>
<td>2063.4</td>
<td>63</td>
<td>0</td>
<td>1</td>
<td>1</td>
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<tr>
<td>6940</td>
<td>2047.6</td>
<td>52</td>
<td>1</td>
<td>1</td>
<td>1</td>
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<tr>
<td>1149</td>
<td>2013.7</td>
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<td>0</td>
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<td>1</td>
<td>1</td>
<td>1</td>
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<tr>
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<tr>
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<td>1885.9</td>
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<td>1</td>
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<tr>
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<td>FN 1</td>
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<td>FN 2</td>
<td>1558.9</td>
<td>-</td>
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</tr>
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</table>

Table 3. Parameter estimates from occupancy models derived from direct observational surveys and eDNA samples with multiple qPCR replicates (for observational surveys using occupancy models, CI = confidence interval; for eDNA samples using Bayesian framework, CI = credible intervals).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate</th>
<th>Upper</th>
<th>Lower</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ψ</td>
<td>0.61</td>
<td>0.38</td>
<td>0.80</td>
<td>Occupancy estimated from observational survey data</td>
</tr>
<tr>
<td>p</td>
<td>0.77</td>
<td>0.62</td>
<td>0.88</td>
<td>Detection probability for visual night encounters</td>
</tr>
<tr>
<td>Ψ</td>
<td>0.61</td>
<td>0.38</td>
<td>0.81</td>
<td>Occupancy estimated from eDNA samples</td>
</tr>
<tr>
<td>θ10</td>
<td>0.04</td>
<td>0.00</td>
<td>0.20</td>
<td>Stage 1 (sample collection) false positive rate</td>
</tr>
<tr>
<td>θ11</td>
<td>0.97</td>
<td>0.85</td>
<td>1.00</td>
<td>Stage 1 true positive rate</td>
</tr>
<tr>
<td>1-θ11</td>
<td>0.03</td>
<td></td>
<td></td>
<td>Stage 1 false negative rate (given by 1-θ11)</td>
</tr>
<tr>
<td>p10</td>
<td>0.02</td>
<td>0.00</td>
<td>0.07</td>
<td>Stage 2 (laboratory analysis) false positive rate</td>
</tr>
<tr>
<td>p11</td>
<td>0.98</td>
<td>0.94</td>
<td>1.00</td>
<td>Stage 2 true positive rate</td>
</tr>
<tr>
<td>1-p11</td>
<td>0.02</td>
<td></td>
<td></td>
<td>Stage 2 false negative rate (given by 1-p11)</td>
</tr>
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</table>
number of samples which amplify. Our results suggest a high probability of false positive detection for any sample amplifying with fewer than three positive qPCR replicates. Additionally, when five or more of the eight replicates amplify, we can be confident that the site is indeed occupied. We also observed that there is little gain in occupancy estimate with this additional effort (Fig. 2).

![Figure 2. Posterior conditional probabilities of species absence (1-Ψ(χ)) given by the number of amplifying qPCR replicates.](image)

## DISCUSSION

We found that eDNA from bromeliad tanks, including sample collection (stage 1) and laboratory analysis (stage 2), is highly reliable, with very low levels of error for both false negatives and false positives (false negative: stage 1 = 3%; stage 2 = 2%; false positive: stage 1 = 4%; stage 2 = 2%). Higher error rates were observed for commercial eDNA surveys for great crested newts in ponds within the UK, from both sample collection (stage 1: false positive rate = 15%; false negative rate = 27%) and laboratory analysis (stage 2: false positive = 5%; false negative = 19%) (Griffin et al., 2020). Phytotelm-breeding species are often elusive and difficult to detect using visual surveys, and occasionally require destructive sampling methods such as removal of plants (Brozio et al., 2017; Torresdal et al., 2017). In these cases, eDNA is a reliable non-invasive method that detects species presence with very low error rates (i.e., low false positive/negative rates). With a single set of eDNA samples we were able to accurately detect species presence in every bromeliad confirmed to be occupied through repeated direct observations, as well as in one bromeliad where species occupancy had not been identified. However, this additional detection had only a single positive qPCR replicate which is highly likely a case of false positive detection, as indicated by the conditional probabilities analysis.

We have demonstrated the feasibility of collecting eDNA samples in a remote setting and transporting them to a laboratory for analysis. Despite one sample being damaged in transit and the potential false positive result described above, it was possible to maintain a contamination-free environment during sample collection and transport as demonstrated by the absence of amplification in the samples of water collected from streams close to the study site. It is also evident from the high proportion of qPCR replicates amplifying in the confirmed positive samples that for a species which spends a large part of its life cycle within the phytotelm (such as a bromeliad), sampling only a small volume of water is not a limiting factor for the recovery of target DNA. Whilst we present the results from a relatively small sample size, we successfully demonstrate the feasibility of extracting and amplifying DNA from water samples as little as 120 ml. The conditional probability of species detection analysis showed that above five qPCR positive replicates, there is limited gain in the estimate of occupancy. Therefore, the number of qPCR replicates may be excessive and a reduction in laboratory replication may be possible without reducing the occupancy estimate; however further analysis would need to be undertaken to confirm this observation which would require a larger data set.

A major advantage of the use of eDNA methods in remote locations is that it can reliably confirm the presence of species in bromeliads, even when bromeliads have high UV exposure, such as at the mountaintop described here. In this scenario, eDNA surveys would have wide-ranging benefits compared to multiple observational visits. Firstly, it is challenging and costly to access remote areas, requiring an experienced team of observers within an expedition that lasts a number of days. Secondly, our described method reduces bias caused by variation in observer experience (Barata et al., 2017) and is sensitive to different life stages (e.g., eggs and larvae; Zinger et al., 2020) that can be missed during visual surveys. Thirdly, eDNA is a non-invasive method that can reduce environmental impact associated with direct observations (Brozio et al., 2017; Torresdal et al., 2017). Most importantly, because high prevalence of chytrid fungus can be found in bromeliads occupied by frogs (Ruano-Fajardo et al., 2016), reduced visits could also decrease the potential risks associated with the spread of wildlife pathogens by the survey team, while eDNA samples can also be reanalysed for the pathogen (Schmidt et al., 2013).

Despite the very high detection rates from eDNA samples, estimated occupancies using eDNA and direct observations were similar. This has implications for the use of eDNA methods for occupancy monitoring since eDNA analysis has laboratory and consumable costs above those incurred in direct observational surveys. For monitoring purposes, estimates of species occupancy can be available from a detection history, which would require multiple visits, and/or eDNA samples with laboratory replication. For our target species, an increase in detection does not improve statistical power and four visits are required to detect a change of 30% in the population using direct observation (Barata et al., 2017). In our case study, the mean costs per bromeliad sampled by non-invasive observational surveys (four nights with a team of two people, 21 bromeliads = £21.29 per bromeliad) is lower than the costs per sample for a single set of eDNA analysis (a one-day visit by one person, two days of lab work by one person, and laboratory supplies,
uncover the extent of the distribution of many elusive phytotelm-breeding species with reduced expedition costs and environmental impacts. eDNA surveys are considered a promising method for amphibian monitoring regardless of species rarity (Burns et al., 2020) or population density (Lopes et al., 2017). However, the use of eDNA method for monitoring occupancy of phytotelm-dependent species will likely improve as the method becomes more cost-effective and we have a better understanding of the factors affecting detection probability in such environments. Our conclusions are applicable to other phytotelm-dependent species, but the feasibility of the method could vary with species’ life history and the volume of water available for analysis.

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