#### ORIGINAL RESEARCH





# Needle in a haystack? A comparison of eDNA metabarcoding and targeted qPCR for detection of the great crested newt (*Triturus cristatus*)

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#### **Abstract**

Environmental DNA (eDNA) analysis is a rapid, cost-effective, non-invasive biodiversity monitoring tool which utilises DNA left behind in the environment by organisms for species detection. The method is used as a species-specific survey tool for rare or invasive species across a broad range of ecosystems. Recently, eDNA and "metabarcoding" have been combined to describe whole communities rather than focusing on single target species. However, whether metabarcoding is as sensitive as targeted approaches for rare species detection remains to be evaluated. The great crested newt Triturus cristatus is a flagship pond species of international conservation concern and the first UK species to be routinely monitored using eDNA. We evaluate whether eDNA metabarcoding has comparable sensitivity to targeted real-time quantitative PCR (qPCR) for T. cristatus detection. Extracted eDNA samples (N = 532) were screened for T. cristatus by qPCR and analysed for all vertebrate species using high-throughput sequencing technology. With qPCR and a detection threshold of 1 of 12 positive qPCR replicates, newts were detected in 50% of ponds. Detection decreased to 32% when the threshold was increased to 4 of 12 positive qPCR replicates. With metabarcoding, newts were detected in 34% of ponds without a detection threshold, and in 28% of ponds when a threshold (0.028%) was applied. Therefore, qPCR provided greater detection than metabarcoding but metabarcoding detection with no threshold was equivalent to qPCR with a stringent detection threshold. The proportion of T. cristatus sequences in each sample was positively associated with the number of positive qPCR replicates (qPCR score) suggesting eDNA metabarcoding may be indicative of eDNA concentration. eDNA metabarcoding holds enormous potential for holistic biodiversity assessment and routine freshwater monitoring. We advocate this community approach to freshwater monitoring to guide management and conservation, whereby entire communities can be initially surveyed to best inform use of funding and time for species-specific surveys.

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#### KEYWORDS

environmental DNA, great crested newt, high-throughput sequencing, metabarcoding, ponds, real-time quantitative PCR, Triturus cristatus

#### 1 | INTRODUCTION

Species monitoring has rapidly evolved with the advent of environmental DNA (eDNA) analysis (Lawson Handley, 2015). eDNA analysis allows highly sensitive detection of rare and invasive species and is increasingly being used for surveys of aquatic species (Biggs et al., 2015; Davy, Kidd, & Wilson, 2015; Evans, Shirey, Wieringa, Mahon, & Lamberti, 2017; Smart et al., 2016; Thomsen et al., 2012). This noninvasive approach uses intracellular and extracellular DNA (e.g., mucus, skin cells, urine/faeces, gametes, hair, deceased remains) released into the environment by organisms to survey for species and assess their distribution (Goldberg et al., 2016; Lawson Handley, 2015; Rees, Maddison, Middleditch, Patmore, & Gough, 2014). Typically for eDNA analysis, DNA is extracted from environmental samples (water, soil, air) and analysed using a targeted or passive approach. The targeted approach uses species-specific primers with conventional PCR (PCR), real-time quantitative PCR (gPCR), or droplet digital PCR (ddPCR), to determine presence-absence and estimate abundance of single species (Goldberg et al., 2016; Shaw, Weyrich, & Cooper, 2016). Conversely, the passive approach uses conserved primers (i.e., primers with binding sites that are shared across multiple taxa, and flank a region of highly variable DNA sequence that enables discrimination between these taxa) and PCR to sequence whole communities with high-throughput sequencing (HTS), termed eDNA metabarcoding (Deiner et al., 2017; Shaw, Weyrich, et al., 2016; Taberlet, Coissac, Pompanon, Brochmann, & Willerslev, 2012; Valentini et al., 2016). Passive eDNA monitoring is particularly attractive to ecologists for biodiversity assessment as a means to detect entire species assemblages alongside rare or invasive species (Blackman et al., 2017; Lacoursière-Roussel, Dubois, Normandeau, & Bernatchez, 2016). However, this gain in community understanding may come at the cost of accuracy and sensitivity. Direct comparisons of these two approaches are essential to determine whether they have comparable power and yield similar results.

Although in its relative infancy, eDNA metabarcoding has proven effective for community biodiversity assessment across a range of taxa in varying environments, particularly freshwater herpetofauna and fish (Bálint et al., 2017; Civade et al., 2016; Evans, Li, et al., 2017; Hänfling et al., 2016; Lacoursière-Roussel et al., 2016; Lopes et al., 2016; Shaw, Clarke, et al., 2016; Valentini et al., 2016). However, eDNA metabarcoding is confounded by potential amplification bias during PCR, preventing capture of all species present in a given area (Kelly, Port, Yamahara, & Crowder, 2014). Species' DNA in community samples is also in competition to bind to metabarcoding primers during PCR, where more common templates are more likely to be amplified. High abundance species may thus prevent the detection of low abundance species, whether by fewer individuals or less DNA

shed, resulting in "species masking" (Brandon-Mong et al., 2015; Evans et al., 2016; Kelly et al., 2014). eDNA metabarcoding may therefore be less capable of identifying eDNA of rare species within a community than species-specific qPCR (Evans et al., 2016).

The sensitivity of eDNA metabarcoding has been evaluated against conventional biodiversity monitoring methods in freshwater ecosystems (Bálint et al., 2017; Civade et al., 2016; Evans, Li, et al., 2017; Hänfling et al., 2016; Lopes et al., 2016; Shaw, Clarke, et al., 2016; Valentini et al., 2016), yet specific investigations comparing the sensitivity of eDNA metabarcoding and targeted qPCR are sparse. Similarly, comparisons of qPCR and conventional survey for species monitoring have included cost projections (Biggs, Ewald, Valentini, Gaboriaud, & Griffiths, 2014; Davy et al., 2015; Evans, Shirey, et al., 2017; Smart et al., 2016), but cost has not been thoroughly assessed in qPCR and eDNA metabarcoding comparisons (Lacoursière-Roussel et al., 2016; Schneider et al., 2016). Schneider et al. (2016) achieved improved detection of invasive mosquito species (IMS) with qPCR and eDNA metabarcoding as opposed to conventional sampling. Although qPCR provided higher detection probability for two species, metabarcoding achieved comparable results for a third species and allowed simultaneous detection of IMS and other taxa in a single sequencing run without development of multiple speciesspecific markers. In another study, eDNA metabarcoding failed to detect wood turtle Glyptemys insculpta (LeConte, 1830) in four rivers where gPCR and conventional visual surveys detected the species (Lacoursière-Roussel et al., 2016). Amplification of longer fragments during metabarcoding versus qPCR could account for difference in sensitivity of the two methods, with the shorter qPCR assay being more capable of detecting heavily degraded DNA (Lacoursière-Roussel et al., 2016). Further research is clearly needed to determine whether these two approaches are comparable.

The great crested newt Triturus cristatus (Laurenti, 1768) (Figure 1) is a model organism for eDNA-based monitoring. T. cristatus secrete mucus, breed in water, and produce aquatic eggs and larvae-all sources of DNA deposition in ponds. The species is rare in parts of the UK and Europe, and as such, all life stages are protected by UK and European legislation (Buxton, Groombridge, Zakaria, & Griffiths, 2017; Rees, Bishop, et al., 2014). eDNA analysis using targeted qPCR has been repeatedly verified against conventional surveying (bottle trapping, torchlight counts, larval netting, egg searches) for T. cristatus and found to achieve comparable or improved species detection (Biggs et al., 2015; Rees, Bishop, et al., 2014; Thomsen et al., 2012). eDNA sampling can be undertaken with relative ease, is cost-efficient (Biggs et al., 2014), and can be implemented in large-scale citizen science monitoring programs without loss of species detection (Biggs et al., 2015). T. cristatus is the first species to be routinely monitored using eDNA in the UK



**FIGURE 1** Adult male great crested newt *Triturus cristatus*. Photograph by Brett Lewis (Lewis Ecology, Brett Lewis Photography)

(Natural England, 2015) and targeted eDNA assays are now offered as a commercial service by ecological consultancies. The targeted eDNA assay is highly effective for *T. cristatus* detection; however, should metabarcoding have comparable sensitivity, this approach would allow detection of *T. cristatus* alongside pond communities and potentially enable more cost-effective monitoring of entire ecosystems and ecological hypothesis testing.

Here, we perform a large-scale comparison (N = 532 ponds) of eDNA metabarcoding and targeted qPCR for T. cristatus detection to compare method sensitivity. A single primer pair that is vertebratespecific for mitochondrial DNA (mtDNA) and requires no a priori knowledge of species composition, was employed for eDNA metabarcoding. The metabarcoding results were then compared to results obtained using the standard T. cristatus qPCR assay (Biggs et al., 2015). Our hypotheses are as follows: (1) eDNA metabarcoding will give equivalent results to qPCR for T. cristatus detection, (2) eDNA metabarcoding sequence read count for T. cristatus will increase as qPCR score (the number of positive qPCR replicates) increases, indicative of eDNA concentration, and (3) metabarcoding primers will amplify DNA from all taxa equally well and no bias toward amplification of T. cristatus will occur (bias would be indicated by a positive association between the proportion of *T. cristatus* sequence reads and PCR product concentration). We also examined cost and investigator effort required by each approach to determine whether a trade-off between cost, time, and amount of data generated exists.

# 2 | MATERIALS AND METHODS

## 2.1 | Sampling

Samples from 532 ponds distributed across three UK counties (Cheshire, Kent and Lincolnshire) were analysed for this project. Of these, 508 ponds (ranging from 9 to 9375 m<sup>2</sup>) were sampled as part of *T. cristatus* surveys through Natural England's Great Crested Newt Evidence Enhancement Programme. *T. cristatus* egg searches were performed once during the daytime at 506 of 508 ponds. Any other

life stages seen were also recorded. A further 24 ponds were sampled for eDNA by ecological consultants for private contracts but egg searches were not undertaken. All water samples were collected using methodology outlined by Biggs et al. (2015). Water samples were then sent to Fera (Natural England) and ADAS (private contracts), where one eDNA sample per pond was produced and analysed according to laboratory protocols established by Biggs et al. (2015). Details of sampling methodology and laboratory protocols are provided in Appendix S1.

# 2.2 | Targeted qPCR for T. cristatus

Targeted qPCR was conducted as part of the *T. cristatus* monitoring programmes mentioned above in Fera and ADAS laboratories during 2015. Both laboratories used a standardised protocol, which tests for PCR inhibitors and sample degradation prior to testing for *T. cristatus* (Biggs et al., 2015). Extracted DNA was amplified by TaqMan probe qPCR using published primers and probe (Thomsen et al., 2012) to amplify an 81 bp fragment of the cytochrome b gene. For each sample, 12 qPCR replicates were performed and a sample recorded as positive for *T. cristatus* if one or more qPCR replicates were positive. Following qPCR, the eDNA samples were placed in storage at -80°C.

## 2.3 | Metabarcoding of vertebrate communities

eDNA samples were stored at -20°C until PCR amplification. Metabarcoding was performed using published vertebrate-specific primers (Riaz et al., 2011) which amplify a 73-110 bp fragment of the 12S ribosomal RNA gene (rRNA). The assay was first validated in silico using ecoPCR software (Bellemain et al., 2010; Ficetola et al., 2010) against a custom, phylogenetically curated reference database for UK vertebrates. Full details of reference database construction are provided in Appendix S2. The complete reference database compiled in GenBank format has been deposited in a dedicated GitHub repository for this study, permanently archived at: https://doi. org/10.5281/zenodo.1188709. Parameters set allowed a 50-250 bp fragment and maximum of three mismatches between the primer pair and each sequence in the reference database. Primers were then validated against tissue DNA extracted from UK amphibian species (Appendix S3) having been previously validated in vitro for UK fish communities by Hänfling et al. (2016). After primer validation, a two-step PCR protocol was used to construct metabarcoding libraries from the eDNA samples. During the first PCR, the target region was amplified using metabarcoding primers, comprised of the aforementioned specific locus primer, random hexamers, sequencing primer, and pre-adapter (Illumina 2011). DNA from the cichlid Rhamphochromis esox (Boulenger, 1908) was used for PCR positive controls (six per PCR plate; N = 114), whilst sterile molecular grade water (Fisher Scientific UK Ltd, UK) substituted template DNA for no template controls (NTCs, six per PCR plate; N = 114). In the second PCR, molecular identification (MID) tags (unique 8-nucleotide sequences) and Illumina MiSeq adapter sequences were added to

the amplified product. Two independent libraries were constructed, each containing 266 eDNA samples, 57 NTCs, and 57 positive controls. Sequencing was performed on an Illumina MiSeq using  $2\times300~\text{bp}$  V3 chemistry at Fera. The first sequencing run revealed human contamination across samples and in some PCR controls; therefore, reactions prepared for the second sequencing run were sealed with mineral oil to minimise PCR contamination. Full details of the eDNA metabarcoding workflow are provided in Appendix S3.

# 2.4 | Bioinformatic processing

Illumina data were converted from raw sequences to taxonomic assignment using a custom pipeline for reproducible analysis of metabarcoding data: metaBEAT (metaBarcoding and eDNA Analysis Tool) v0.8 (https://github.com/HullUni-bioinformatics/metaBEAT). Bioinformatic data processing/analysis largely followed the workflow outlined by Hänfling et al. (2016), with minor modifications (see Appendix S3 for details). To ensure reproducibility of analyses, the workflow has been deposited in the GitHub repository.

## 2.5 | Data analysis

All downstream analyses were performed in the statistical programming environment R v.3.3.2 (R Core Team, 2016). Data and R scripts have been deposited in the GitHub repository. Manipulation of the dataset produced by metaBEAT is described in Appendix S4.

#### 2.5.1 Detection thresholds and contamination

At present, there are no standard guidelines for eDNA analysis to indicate minimum number of positive eDNA samples or replicates required to class sites as species positive (Goldberg et al., 2016). Samples analysed by qPCR in this study were previously considered T. cristatus positive if one or more qPCR replicates gave a positive result (Biggs et al., 2015). We term this analysis qPCR NT (No Threshold). This inference of species presence is employed across many studies but may not be reliable or reproducible (Goldberg et al., 2016). More stringent qPCR thresholds reduced detection sensitivity for palmate newt Lissotriton vulgaris (Razoumowsky, 1789) (Smart et al., 2016), but may be necessary to ensure consistency and prevent false positives (Rees, Maddison, et al., 2014). To facilitate comparison with current qPCR scoring (our NT interpretation) and eDNA metabarcoding, we applied a stringent qPCR threshold of ≥4/12 positive qPCR replicates to infer species presence and termed the new analysis qPCR TA (Threshold Applied).

The raw eDNA metabarcoding dataset with no detection thresholds applied was termed metabarcoding NT (No Threshold). A second dataset was constructed to reduce the potential for false positives by application of a species-specific threshold: a species was only classed as present at a given site if its sequence frequency exceeded a species-specific threshold. Thresholds for each species were defined by analysing sequence data from PCR positive controls (N = 114) and identifying the maximum sequence frequency

for a given species across all PCR positive controls (Table S2). For example, the species-specific false positive sequence threshold for *T. cristatus* was 0.028% to omit all false detections in the PCR positive controls. The resultant dataset was termed metabarcoding TA (Threshold Applied).

We tested whether mineral oil reduced contamination by analysing the distribution of positive control sequences (R. esox) and human DNA in eDNA samples, and any DNA in NTCs, across both sequencing runs using binomial generalised linear mixed effects models (GLMMs) within the R package "Ime4" v1.1-12 (Bates, Mächler, Bolker, & Walker, 2015). The response variable was presence-absence of contamination and explanatory variables were PCR plate (random effect) and sequencing run, i.e. mineral oil sealed versus non-sealed (fixed effect). Human DNA may be present in eDNA samples as a real environmental signal or contaminant prior to PCR and thus may not be a true PCR contaminant. Consequently, contamination in eDNA samples was examined using several model permutations, where contamination comprised both cichlid and human DNA, cichlid DNA alone, and human DNA alone. An information-theoretic approach using Akaike's Information Criterion (AIC) to evaluate model fit was employed, where low AIC models are more parsimonious than high AIC models (Akaike, 1973). Significance of the fixed effect in the model was tested by a likelihood ratio test (LRT).

# 2.5.2 | Comparison of eDNA methods for *T. cristatus* detection

We tested the null hypothesis of no significant difference in sensitivity of qPCR and metabarcoding. Overall agreement between eDNA metabarcoding and qPCR for *T. cristatus* detection was measured using Cohen's kappa coefficient (Cohen, 1960), following which Pearson's Chi-squared Test of Independence was used to test equality of *T. cristatus* detection between eDNA approaches.

Previously, Biggs et al. (2015) found qPCR score was an inconsistent predictor of *T. cristatus* abundance, where ponds with low scores had low newt counts but high scores did not correspond to large populations. qPCR score may only be proxy for the amount of DNA present rather than the number of individuals. The relationship between read count and qPCR score has not been examined previously, and whether read production is indicative of DNA concentration remains unknown. We hypothesised samples with higher qPCR score would have increased *T. cristatus* read count. First, the average number of *T. cristatus* reads produced by eDNA metabarcoding per qPCR score (1–12 of 12) was calculated. A Spearman Rank Correlation was then used to test for a relationship between average read count and qPCR score.

Following data exploration (see Appendix S4), a negative binomial GLMM was used to counter overdispersion and improve model fit. The GLMM examined read count in relation to qPCR score, accounting for other variables that may affect metabarcoding signal strength. Variation in *T. cristatus* read count was examined using the proportion of *T. cristatus* reads within the total number of reads produced for each eDNA sample as the response

variable. Sequencing run and PCR plate were considered random effects and all other explanatory variables as fixed effects (qPCR score, sample degradation, sample inhibition, post-PCR eDNA concentration). Presence-absence of sample degradation and inhibition was determined by qPCR in 2015 using methodology outlined by Biggs et al. (2015). Model fit was again evaluated using AIC, and significance of fixed effects in the model was tested with stepwise backward deletion of terms from the model informed by LRTs. All values were bound in a new data frame and model results plotted for evaluation using the R package "ggplot2" v 2.1.0 (Wickham, 2009).

## 2.5.3 | Cost and investigator effort

Cost of materials and investigator effort and salary (hourly rate of £21.20 assumed) were calculated for eDNA samples; however, estimates do not include travel to sampling sites, procedural controls, qPCR standards, or consumables and reagents required for assay optimisation. Time required to perform PCR for metabarcoding and qPCR was estimated assuming available machinery to run four PCR plates in parallel and one qPCR plate.

#### 3 | RESULTS

## 3.1 | Targeted qPCR and egg searches

Targeted qPCR detected *T. cristatus* in 253 (49.80%) samples analysed by Fera (*N* = 508). Of 255 (50.20%) samples that were negative, one was inhibited and nine were degraded. qPCR and egg searches produced consistent results for 297 (58.47%) ponds, with 51 (10.04%) positive and 246 (48.43%) negative ponds by both methods. Of the 211 ponds where there was disagreement between methods, 202 (39.76%) were qPCR positive but negative by egg searches, and seven (1.38%) were positive with egg searches but qPCR negative. Of 24 samples analysed by ADAS, 12 (50.00%) were qPCR negative and 12 (50.00%) were qPCR positive for *T. cristatus*. No egg search data were available for these ponds.

# 3.2 | Vertebrate metabarcoding

The *in silico* and *in vitro* primer validation confirmed that *T. cristatus*, and other native UK amphibians tested, can be reliably amplified and identified with the chosen assay (Appendix S5: Figure S1). Furthermore, the *in silico* approach showed that the majority of all UK vertebrates can be amplified (see Appendix S5 for details). Both sequencing runs had comparable yield and sequencing quality score; summary statistics for each sequencing run and read counts for taxonomic assignment levels are provided in Appendix S5 (Tables S3, S4). A full summary of sequence read count data is also given in Appendix S5 (Table S5). eDNA metabarcoding identified a combined total of 60 species (Appendix S5: Figure S2) across both sequencing libraries, with 375,954 and 508,879 sequences assigned to *T. cristatus* from each library. Analyses of overall pond species compositions

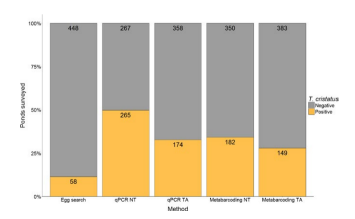
inferred by eDNA metabarcoding (Appendix S5: Figure S3, Table S6) are reported separately (Harper et al., 2018).

All samples (*N* = 532) were sequenced and of 57 samples that did not produce visible PCR bands, nine generated sequence reads. Notably, the 57 samples were not inhibited or degraded at time of qPCR. Weak PCR bands were observed in some NTCs; therefore, all PCR controls were sequenced (Appendix S5: Figures S4–6). Six NTCs contained *T. cristatus* DNA but only one exceeded 100 *T. cristatus* reads (307/330 reads). Twelve other sources occurred in NTCs (Appendix S5: Table S7); seven occurred in more than one NTC and eight had high maximum read counts (>100 reads).

Contamination of NTCs (any DNA) and environmental samples (cichlid/human DNA) was observed (Appendix S5: Figures S4-6). Read counts of NTC contaminants were reduced between sequencing runs with the addition of mineral oil to PCR reactions included on the second sequencing run (Appendix S5: Figures S4-6) but this reduction was not statistically significant (GLMM:  $\chi_1^2 = 2.083$ ,  $F_1 = 1.941$ , p > .05). Mineral oil did not reduce human DNA signal in environmental samples between sequencing runs either (GLMM:  $\chi_1^2 = 3.608$ ,  $F_1 = 3.591$ , p > .05); however, it did reduce human DNA in combination with cichlid DNA (GLMM:  $\chi_1^2 = 10.348$ ,  $F_1 = 21.143$ , p < .01), and cichlid DNA contamination alone (GLMM:  $\chi_1^2 = 5.053$ ,  $F_1 = 6.978$ , p < .05) of environmental samples.

# 3.3 | eDNA metabarcoding vs qPCR for T. cristatus detection

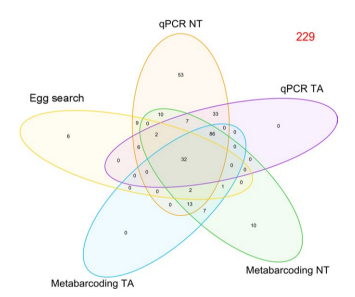
*T. cristatus* detection by metabarcoding NT (34.21%) was less sensitive than qPCR NT (49.81%) but marginally higher than qPCR TA (32.71%) (*N* = 532 ponds, Figure 2). Metabarcoding TA had lower detection efficiency (28.01%) and failed to detect *T. cristatus* in 116 and 25 ponds where the species was detected by qPCR NT and qPCR TA, respectively. Nonetheless, both molecular approaches attained higher *T. cristatus* detection than daytime egg searches (11.46%) in 506 ponds where all three approaches were implemented.



**FIGURE 2** Comparison of survey methodology for *T. cristatus* detection in freshwater ponds across the UK. Bars represent proportion of positive and negative *T. cristatus* ponds by each method with frequency displayed on bars

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Overlap between survey methods for positive T. cristatus ponds (N = 277), and unique detections by each method are summarised in Figure 3. Negative T. cristatus ponds (N = 229) were examined in combination with species positive ponds in Appendix S5 (Table S8). Each survey method detected the species in ponds where other methods failed. Despite lower T. cristatus detection efficiency, egg searches detected the species in six ponds where it went undetected by qPCR and metabarcoding. Metabarcoding NT and metabarcoding TA revealed T. cristatus in seven ponds which other methods did not, whilst gPCR NT and gPCR TA detected T. cristatus in 33 ponds unique to other methods. All methods detected T. cristatus in 32 ponds, and both metabarcoding and qPCR identified T. cristatus in 86 ponds. Disagreement between molecular methods was more likely when samples were positive rather than negative by qPCR. Without thresholds, 39.25% of qPCR-positive ponds (N = 265) were negative by metabarcoding, but 7.87% of qPCR-negative ponds (N = 267) were



**FIGURE 3** Venn diagram which summarises the number of positive *T. cristatus* detections (*N* = 277) by each method (egg search, qPCR NT, qPCR TA, metabarcoding NT, and metabarcoding TA), and overlap in *T. cristatus* detection between methods for 506 ponds where all methods were applied. Negative *T. cristatus* detections (*N* = 229) are highlighted in red

positive by metabarcoding. With thresholds, 29.31% of qPCR-positive ponds (N = 174) were negative by metabarcoding, whereas 7.26% of qPCR-negative ponds (N = 358) were positive by metabarcoding.

Agreement between eDNA approaches is summarised in Table 1. Agreement was strongest between eDNA approaches when the aPCR detection threshold was applied, irrespective of whether the metabarcoding detection threshold was applied. Metabarcoding (NT or TA) and qPCR TA did not significantly differ in their detection of T. cristatus (Table 1). An identical positive correlation was observed between qPCR score and the average number of T. cristatus reads obtained for samples belonging to each qPCR score ( $r_c = .648$ , df = 11, p < .05), regardless of threshold application to the metabarcoding data. Despite some inconsistency across gPCR scores, samples with a higher qPCR score generally had more T. cristatus reads, supportive of a relationship between metabarcoding and abundance of eDNA from single species. Notably, metabarcoding produced T. cristatus reads for qPCR NT and qPCR TA negative samples, but the T. cristatus metabarcoding signal of these (qPCR  $NT_{negative}$  = 2,639 reads max., qPCR  $TA_{negative} = 3,075$  reads max.) was much lower than samples with higher qPCR score (max. 65,325 reads; Appendix S5). Further examination of the relationship between gPCR score and metabarcoding TA revealed qPCR score and post-PCR eDNA concentration of samples also influenced the proportion of T. cristatus reads, i.e. relative T. cristatus sequence read production (Table 2). A significant positive relationship was observed between qPCR score and the proportion of T. cristatus reads within total reads per sample (p < .001) (Figure 4a). Conversely, post-PCR eDNA concentration had a significant negative influence on the proportion of T. cristatus reads (p < .001), where read proportion decreased as post-PCR eDNA concentration increased (Figure 4b).

# 3.4 | Comparison of method cost and investigator effort

Cost and investigator effort for both eDNA approaches were comparable. Metabarcoding was marginally more expensive (£3 per pond) than qPCR, but used 1 day less of investigator effort. A full breakdown of expenditure per pond is given in Appendix S5 (Table S9) and summarised in Figure 5.

**TABLE 1** Summary of analyses testing for agreement between eDNA approaches, with threshold applied (TA) and no threshold (NT), for *T. cristatus* detection. Cohen's kappa coefficient (*k*) represents strength of agreement between methods (1 = 100%). Pearson's Chi-squared Test of Independence tested whether methods significantly differed for *T. cristatus* detection

Comparison	Probability of observed agreement	Probability of expected agreement	k	Overall agreement	$\chi^2$	df	р
Metabarcoding NT qPCR NT	.77	.50	0.53	Moderate	25.940	1	<.001
Metabarcoding TA qPCR NT	.74	.50	0.48	Moderate	52.291	1	<.001
Metabarcoding NT qPCR TA	.84	.56	0.63	Good	0.207	1	>.05
Metabarcoding TA qPCR TA	.86	.58	0.66	Good	2.561	1	>.05

**TABLE 2** Summary of analyses testing for variation in proportion of *T. cristatus* sequence reads in a sample produced by eDNA metabarcoding, attributable to qPCR score or post-PCR eDNA concentration. Test statistic is for LRT used

Model variables	N (ponds)	df	AIC	Effect size	Standard error	$\chi^2$	F	р
qPCR score	532	1	1,578.3	0.373	0.032	150.682	147.117	<.001
post-PCR eDNA concentration	532	1	1,441.9	-0.056	0.015	14.272	12.457	<.001

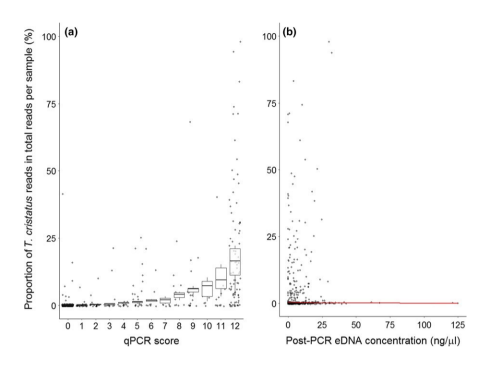


FIGURE 4 Relationship between fixed effects (qPCR score, post-PCR eDNA concentration) and response variable (proportion of *T. cristatus* reads) in eDNA samples, as predicted by the negative binomial GLMM. The 95% CIs, as calculated using the predicted proportions and standard error for these predictions, are given for each relationship. The observed data (points) are also displayed against the predicted relationships (boxes, line). The proportion of *T. cristatus* reads within eDNA samples increased as qPCR score increased (a). Conversely, the proportion of T. cristatus reads decreased as post-PCR eDNA concentration increased (b)

# 4 | DISCUSSION

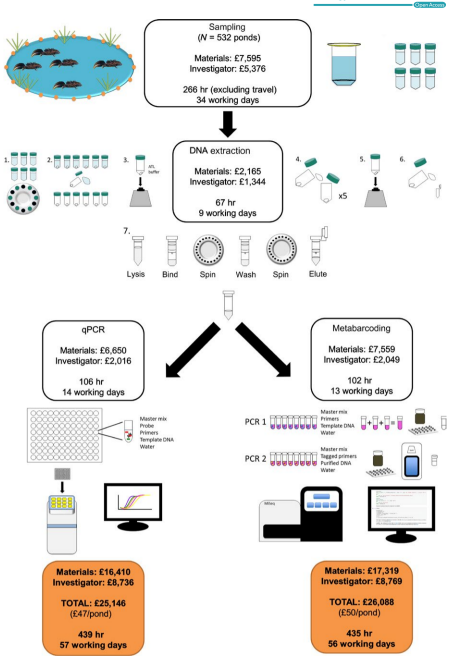
We have demonstrated eDNA metabarcoding is a highly sensitive tool for monitoring *T. cristatus* alongside the wider biological community, corroborating other comparisons of eDNA metabarcoding and qPCR for single-species monitoring (Lacoursière-Roussel et al., 2016; Schneider et al., 2016). Despite reduction in single-species detection, eDNA metabarcoding revealed a wealth of biodiversity information and could enable more effective freshwater monitoring networks and better understanding of community structure and ecosystem function alongside *T. cristatus* monitoring (Biggs, von Fumetti, & Kelly-Quinn, 2016). However, both eDNA approaches have advantages and drawbacks which must be considered for design and implementation of biodiversity monitoring programs.

# 4.1 | Single-species detection by qPCR and metabarcoding

A direct comparison of sensitivity between qPCR and metabarcoding is not straightforward: stochasticity in qPCR largely occurs during amplification (volume of template DNA and technical replication), whereas stochastic variation during metabarcoding arises through PCR amplification and sequencing (depth and replication) (Deiner

et al., 2017; Kelly et al., 2014; Thomsen et al., 2016). In our study, 12 independent qPCR replicates were performed for each sample but due to limited resources, metabarcoding was based on three pooled PCR replicates which were sequenced once only. Therefore, to enable a fair comparison between methods in terms of PCR effort, a threshold of ≥4/12 positive replicates (aPCR TA) was applied to the qPCR data. Detection sensitivity was most similar between methods with the qPCR threshold and without the metabarcoding threshold. Both eDNA metabarcoding and qPCR displayed reduced T. cristatus detection when thresholds were applied; however, this may reflect reduced false positive detections rather than decreased sensitivity. Lower sensitivity of the eDNA metabarcoding approach used here may also stem from sample degradation during long-term storage. The samples used were stored for more than 12 months at -80°C before metabarcoding. However, long-term storage and continual freeze-thawing of samples may allow aggregation of inhibitory substances which impair PCR amplification and cause false negatives (Takahara, Minamoto, & Doi, 2015).

Despite lower sensitivity, strength of eDNA signal produced by metabarcoding was correlated with that of qPCR, where both *T. cristatus* average read count and read proportion broadly increased with qPCR score of eDNA samples. The correlation was inconsistent though, where high average or proportional *T. cristatus* read count did not always correspond to high qPCR score. Biggs et al. (2015)



**FIGURE 5** Cost and investigator effort required for targeted qPCR of *T. cristatus* and eDNA metabarcoding of vertebrate communities from pond water samples

also found a variable positive association between qPCR and *T. cristatus* counts, where high qPCR score did not always correlate with high counts. Quantitative data on eDNA concentration are needed to examine the performance of each eDNA approach in relation to the amount of eDNA present, and whether these tools can reliably estimate species abundance. This data can be obtained with highly sensitive qPCR assays, and inclusion of internal DNA standards in sequencing runs for metabarcoding (Ushio et al., 2018). Nonetheless, our results suggest performance of metabarcoding and qPCR is linked and influenced by external factors. Evans et al. (2016) suggested the relative abundance and biomass of a species interact to exert a combined effect on eDNA production rate and subsequent

metabarcoding detection. The abundance, biomass, and distribution of *T. cristatus* (Biggs et al., 2015), as well as shedding rate, environmental factors, and eDNA transport (Buxton et al., 2017; Goldberg et al., 2016), may all influence detection and concentration of eDNA, and inferences made using qPCR and metabarcoding.

The comparison between qPCR and metabarcoding must also be examined in context of the sequencing effort. Here, we sequenced a large number of samples (380 including PCR controls) per run to provide a realistic cost scenario for routine monitoring. Yet, metabarcoding sensitivity would likely improve with an increase in read depth per sample (Kelly et al., 2014). In order to directly compare eDNA signal production by these approaches, it may be necessary

to perform sequencing replicates to verify true positives where rare species are expected and generate an "eDNA metabarcoding score" system similar to qPCR (Brandon-Mong et al., 2015; Civade et al., 2016; Port et al., 2016; Thomsen et al., 2016). PCR and sequencing replication in metabarcoding may enhance species detection probability through improved amplification of low abundance or highly degraded DNA (Ficetola et al., 2015; Port et al., 2016) that is readily amplified by qPCR (Lacoursière-Roussel et al., 2016).

Similarly, sequencing of independent biological replicates, opposed to pseudoreplicates from a single water sample, may improve detection and minimise false negatives produced by eDNA metabarcoding (Andruszkiewicz et al., 2017; Bálint et al., 2017). Currently, 90 ml (6 × 15 ml sampled from 600 ml) water is sampled during *T. cristatus* eDNA survey, followed by ethanol precipitation (Biggs et al., 2015). Whilst this may be appropriate for highly sensitive targeted qPCR, larger water volumes and filtration may be required to capture eDNA from less abundant vertebrates and characterise community diversity (Shaw, Weyrich, et al., 2016). Additionally, eDNA from different species, and individuals within species, can be unevenly distributed throughout water bodies and may be concentrated in particular areas (Biggs et al., 2015; Evans, Li, et al., 2017; Hänfling et al., 2016), thus sampling strategies must be carefully designed to ensure eDNA samples are representative of biodiversity present.

Metabarcoding assays are also susceptible to problems from taxon bias, DNA swamping and bioinformatics related problems (Kelly et al., 2014; Shaw, Weyrich, et al., 2016; Taberlet et al., 2012). Potential reduction in sensitivity of passive community sequencing versus targeted qPCR may relate to the performance of metabarcoding primers for target species. During metabarcoding, DNA from rare species may be masked by highly abundant species (Schneider et al., 2016), or under-represented due to disproportionate eDNA shedding rates across species and preferential amplification of other species (Kelly et al., 2014). PCR-free workflows (i.e., shotgun sequencing) eliminate this bias through indiscriminate sequencing; however, this is unsuitable for conservation projects with target species as a mass of uninformative data are produced, and too costly for routine monitoring schemes (Shaw, Weyrich, et al., 2016; Valentini et al., 2016). We found T. cristatus read proportion was negatively associated with post-PCR concentration of eDNA samples. As a positive relationship was not observed, this would suggest PCR amplification with our selected marker and primers was not biased toward our focal species. However, we cannot conclude that our metabarcoding assay was free of primer bias as post-PCR concentration of eDNA samples can be influenced by PCR stochasticity.

Multiple markers (e.g., COI, CytB, 12S, 16S) are increasingly used in eDNA metabarcoding to cast a wider net of species detection and minimise primer bias (Evans, Li, et al., 2017; Evans et al., 2016; Hänfling et al., 2016; Shaw, Clarke, et al., 2016; Valentini et al., 2016). Using markers from both mitochondrial and nuclear genes may reduce bias associated with specific genes or primers, and provide greater taxonomic resolution (Kelly et al., 2014). Furthermore, multiple markers of different lengths may enhance understanding of eDNA persistence and state, and species location. Long barcodes

bind to stable DNA that has been recently deposited by species (Hänfling et al., 2016) and may reduce false negatives whilst increasing taxonomic resolution and accuracy (Kelly et al., 2014; Shaw, Clarke, et al., 2016; Valentini et al., 2016). In contrast, short barcodes (such as 12S used here) challenge sequencers and bioinformatics tools (Shaw, Clarke, et al., 2016; Taberlet et al., 2012), but readily amplify short, degraded DNA fragments that persist longer and possibly disperse further in water bodies, improving probability of detection (Hänfling et al., 2016). It is possible that metabarcoding detection rates could be improved using group-specific metabarcoding primers for amphibians, such as the "batra" set recently designed by Valentini et al. (2016). More specific primers could increase relative coverage of *T. cristatus*, providing more comparable detection rates to qPCR. This is worth investigating, but with the caveat that group-specific primers obviously restrict the biodiversity information that can be gained from an ecosystem.

## 4.2 | False negatives

This study did not aim to evaluate sensitivity of molecular methods against standard T. cristatus survey methodologies. Egg searches were used to detect false negatives produced by qPCR and metabarcoding and in doing so, revealed some interesting results. Biggs et al. (2015) previously demonstrated qPCR had higher detection rate than egg searches (as well as torchlight, netting, and bottle trapping), but here we show this also holds true for metabarcoding. Importantly, absence of eggs does not infer absence of adults, and this method is highly dependent on weather conditions and water clarity (Biggs et al., 2015; Rees, Bishop, et al., 2014). Despite considerably higher detection rate of both eDNA approaches, eggs were recorded in a small number of ponds that were eDNA negative. eDNA analysis can incorrectly infer absence or low abundance of species if inhibition or interference from non-target DNA has occurred (Goldberg et al., 2016). Alternatively, eDNA false negatives may have been a by-product of sampling strategy and effort for T. cristatus. Larger water volumes and/or more biological replication instead of pseudoreplication (established T. cristatus eDNA sampling strategy) may improve detection (Andruszkiewicz et al., 2017; Bálint et al., 2017; Lopes et al., 2016). All methods revealed T. cristatus in ponds where other approaches failed, emphasising that these species monitoring tools are complementary and should be used in combination to achieve maximum detection probability. However, integrative strategies combining molecular and conventional tools are often not costefficient for most applications.

## 4.3 | False positives

False positives may arise from field contamination and eDNA transport in the environment—particularly by waterfowl (Shaw, Clarke, et al., 2016). eDNA is retained by predators, discarded in feces, and transported by anthropogenic activity, combined with natural water currents and flow (Hänfling et al., 2016). In the laboratory, PCR-accumulated and sequencing error, including primer mismatch

(Andersen et al., 2012) and "tag jumps" (Schnell, Bohmann, & Gilbert, 2015), can induce misassignment leading to false positives, cross-contamination between samples, or laboratory contamination (Andruszkiewicz et al., 2017).

False positives can be modeled and estimated using Site Occupancy Modelling of metabarcoding data (Ficetola et al., 2015), or risk of false positives minimised using a sequencing threshold, that is the number of sequence reads required for a sample to be species positive (Civade et al., 2016; Evans, Li, et al., 2017; Hänfling et al., 2016). However, such thresholds can reduce detection of rare species, a primary goal of this study, and may fail where false and true positives occur at similar frequency (Hänfling et al., 2016). Instead, we calculated species-specific sequence thresholds to more accurately control for false positives in our dataset without compromising *T. cristatus* detection.

In our study, human DNA occurred at high frequency and abundance; this may have been a true environmental signal from pond water, or real contaminant as encountered in other metabarcoding research (Port et al., 2016; Thomsen et al., 2016; Valentini et al., 2016). Blocking primers can prevent amplification of abundant nontarget DNA like human (Valentini et al., 2016) but may fail (Thomsen et al., 2016) or prevent amplification of target taxa (Port et al., 2016). Alongside human, other aquatic and terrestrial vertebrate DNA occurred at high frequency in NTCs, although these were not removed by addition of mineral oil. An even stricter forensic laboratory set-up, such as that employed for ancient DNA (aDNA), should be adopted to ensure data robustness. Positive and negative controls should be included at each stage of metabarcoding workflows to monitor contamination (Deiner et al., 2017). However, preventive measures inevitably increase research cost and some degree of contamination is unavoidable in metabarcoding (Brandon-Mong et al., 2015; Kelly et al., 2014; Port et al., 2016; Thomsen et al., 2016).

Our results also highlight the importance and impact of qPCR thresholds when inferring species presence-absence. Similar to Smart et al. (2016), we found a stringent qPCR threshold reduced detection sensitivity. As yet, no guidance exists to indicate how many samples or replicates must be positive to class a site as species positive (Goldberg et al., 2016; Smart et al., 2016) but clearly this must be addressed to improve standardisation and reproducibility of eDNA research. Importantly, less stringent thresholds (and false positives inherent to these) are somewhat precautionary and may better protect T. cristatus by preventing development. Therefore, whilst reduction or removal of false positives is desirable, detection thresholds must not compromise protection of threatened species either. Until a suitable threshold can be established, it may be more appropriate to re-analyse samples which yield one positive qPCR replicate to prevent false positives (Goldberg et al., 2016; Rees, Maddison, et al., 2014).

# 4.4 | Cost and investigator effort

Cost efficiency combined with the overarching aim of a monitoring or conservation program should always be considered. We found

eDNA metabarcoding was slightly more costly than qPCR but both approaches required similar investigator effort, qPCR scales to the number of samples being processed (Schneider et al., 2016) whereas metabarcoding has fixed costs, including reagent kit for HTS platform (Bálint et al., 2017). eDNA metabarcoding becomes more costefficient as more samples are processed (Bálint et al., 2017) but fewer replicates would reduce gPCR cost (Davy et al., 2015; Smart et al., 2016). Cost of eDNA monitoring is influenced by sample size, methods, replication, laboratory, statistical power, and occupancy modeling (Davy et al., 2015; Evans, Shirey, et al., 2017). Consequently, cost is proportional to project requirements (Davy et al., 2015) and will vary depending on choice of qPCR or metabarcoding workflow. Whilst qPCR is established technology that has reached its price ceiling, HTS is relatively new technology and prices will continue to drop, meaning higher sample throughput and more technical replication will be possible. We therefore argue that metabarcoding will become more cost-efficient in the long-term, providing more data at lower cost and comparable sensitivity to qPCR. However, where samples cannot be processed in large batches, qPCR may retain cost efficiency.

#### 5 | CONCLUSION

eDNA metabarcoding holds promise for holistic biodiversity monitoring of freshwater ponds as opposed to targeted qPCR for flagship or indicator species such as T. cristatus. Metabarcoding can reveal entire species assemblages from environmental samples without prior ecosystem information and provide broad-scale distribution data for multiple species simultaneously. Nonetheless, the method at present appears to be less sensitive than qPCR for single-species monitoring, and species detection by molecular and conventional methods was incongruent. Comprehensive study of the influence of water volume, eDNA capture, and extraction method, and sample storage on single-species and community detection in lentic and lotic systems is required. Minimising the risk of false positives and contamination remains a pressing issue in metabarcoding, and standard contamination measures (Goldberg et al., 2016) may be insufficient for analysis of vertebrate assemblages. Currently, cost and investigator effort required for metabarcoding and qPCR are broadly equivalent, but reduced sequencing costs may level the playing field. We conclude that eDNA metabarcoding is not yet a replacement for targeted qPCR and conventional survey, but rather another tool in the ecologist toolbox. Ultimately, choice of monitoring tool(s) is specific to the aims of each conservation project. At present, qPCR retains sensitivity for T. cristatus populations of all sizes, regardless of sample number processed. Under a realistic conservation monitoring scenario, where funding is limited and samples must be processed in large batches, metabarcoding may suffer from false negatives due to reduced sequencing depth and replication. However, in many cases, the biodiversity information generated by this approach, and its implications for community ecology and conservation, will eclipse lower sensitivity. This passive screening approach would be most effective for initial survey of water bodies to generate broad-scale multi-species distribution data. This holistic data can then inform best use of funding and time for targeted species-specific survey.

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#### **CONFLICT OF INTEREST**

None declared.

#### **AUTHOR CONTRIBUTIONS**

B.H., L.R.H., L.L.H and N.B. conceived and designed the study. H.C.R., K.C.G., and N.B. contributed samples for processing. H.C.R. performed extractions under licence from Natural England and qPCR for eDNA samples from private contracts at The University of Nottingham. N.B. performed qPCR for eDNA samples from Natural England's Great Crested Newt Evidence Enhancement Program. L.R.H. performed remaining laboratory work and analysed the data. I.P.A. and E.L. offered advice on and supervised sequencing. C.H. assisted with bioinformatics analysis. P.B. and S.P. contributed datasets for analysis. L.R.H. wrote the manuscript, which all authors revised.

#### **DATA ACCESSIBILITY**

Raw sequence reads have been archived on the NCBI Sequence Read Archive (Bioproject: PRJNA417951; SRA accessions: SRR6285413–SRR6285678). Jupyter notebooks, R scripts and corresponding data are deposited in a dedicated GitHub repository (https://github.com/HullUni-bioinformatics/Harper\_et\_al\_2018) which has been permanently archived (https://doi.org/10.5281/zenodo.1188709).

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#### SUPPORTING INFORMATION

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