An optimised eDNA protocol for detecting fish in lentic and lotic freshwaters using a small water volume

Teja Petra Muha, Chloe Victoria Robinson, Carlos Garcia de Leaniz, Sofia Consuegra

Swansea University, Department of Biosciences, Singleton Park, Swansea, Wales, United Kingdom

¤ Current address: Labena d.o.o., BIA Separations CRO, Ljubljana, Slovenia
¤ Current address: Centre for Biodiversity Genomics and Department of Integrative Biology, 7 University of Guelph, Guelph, ON Canada
* teja.muha@biaseparationscro.com

Abstract

Environmental DNA (eDNA) is increasingly being used for assessing the presence and relative abundance of fish in freshwater, but existing protocols typically rely on filtering large volumes of water which is not always practical. We compared the effects of water volume, filtration type and eDNA extraction procedures in the detection of fish in three freshwater bodies (pond, lake and river) using a short fragment of the 12s rRNA mtDNA gene. Quantification of eDNA capture efficiency after DNA extraction, as well as amplification efficiency, were evaluated by conventional PCR and quantitative PCR. No significant differences on eDNA capture yield were found among freshwater bodies, but increasing water volume had a positive effect on eDNA capture and amplification efficiency. Although highest eDNA capture rates were obtained using 2 L of filtered water, 100 mL syringe filtration in combination with ethanol- sodium acetate precipitation proved to be more practical and increased quantitative PCR amplification efficiency by 6.4%. Our results indicate that such method may be optimal to detect fish species effectively across both lotic and lentic freshwater environments.

Introduction

Environmental DNA (eDNA) is increasingly being used in freshwater environments to detect the presence of target invertebrate and vertebrate species, based on the detection of short extracellular DNA fragments released into the environment [1–3]. eDNA detection can be used for management purposes, such as monitoring of species’ presence/absence [1], invasive species detection [4], relative abundance estimates [5] and use of space [6]. In some cases it can offer more efficient estimations of relative abundance than conventional sampling techniques [7] as it can provide higher detection sensitivity [8]. Examples of accurate eDNA presence/absence detection rates include the American bullfrog *Lithobates catesbeianus* [9], the smooth newt *Lissotriton vulgaris* [10] and the great crested newt *Triturus cristatus* [11].

The analysis of eDNA allows to identify particular target species or even the composition of an entire community, by using either barcoding or metabarcoding approaches. Species specific
eDNA assays can be used to estimate the density of species, through the relationship between their abundance and the detection rate of their eDNA [12–14], mainly by quantitative PCR (qPCR) [15, 16]. On the other hand, eDNA metabarcoding takes advantages of the ability of next generation sequencing (NGS) techniques to sequence short fragments of DNA [17] and assign the sequences to its corresponding taxon, potentially allowing to assess relative abundance based on within target species comparison [5].

Several studies have focused on the benefits and limitations of different eDNA techniques [1, 18–21], and a number of comparative approaches have tested the efficiencies of eDNA capture by ethanol precipitation or filtration [20, 22], methods of preservation [23, 24], filter types and extraction kits [25]. It has been established, for example, that the efficiency of eDNA sampling techniques differs between lentic and lotic bodies, based on a study of four different invertebrate species using species specific primers [20]. Yet, a consistent application of the same eDNA protocol across water bodies for species detection [26, 27] or relative abundance purposes [28, 29] is still lacking.

Two of the most widely employed techniques of eDNA capturing are the ethanol- sodium acetate precipitation [30] and the filtration method [31–33]. Ethanol precipitation allows a wider size range of eDNA detection, whereas filtering largely depends on the pore size [34]. Glass fibre filters [26, 31, 35] and cellulose nitrate filters [29, 36, 37] with different pore sizes are the most commonly used filter materials in eDNA studies. Both methods (precipitation and filtering) have shown variable success rates in comparative studies [22, 34], depending on the volume of water, pore size, filter material and extraction methods used, in addition to environmental and physical conditions [20, 22, 38]. For example, ethanol- sodium acetate precipitation is unfeasible for large water volumes whereas the efficiency of the filtration largely depends on the turbidity of the water filtered water, resulting in different eDNA capture success rate. Environmental conditions in lotic bodies, particularly the acidity, can accelerate eDNA decay, affecting its abundance [39]. Thus, eDNA extraction using ethanol- sodium acetate precipitation tends to be done on small (15 mL) water samples [7, 9, 30] and appears suitable when target species are highly abundant (and hence there is a lot of eDNA) in small or closed freshwater systems [34], whereas filtration of larger volumes of water seems to be more efficient in larger systems. The type of eDNA extraction kit also determines overall eDNA capture rate efficiency [20, 40–42] but this can vary depending on the presence of inhibitors [42, 43] and pollutants that can increase the number of extraction steps, and unintentionally provide false positives by increasing exposure to potential contamination [44].

A fully optimised method should have low contamination risk and ideally allow the sampling of different water bodies. Using syringe filters in combination with ethanol precipitation, for example, could reduce the risk of contamination at the start of the eDNA processing pipeline [44]. We carried out a comparison of different methods of eDNA collection and extraction in both lentic and lotic freshwaters, using vertebrate specific primers (avoiding species specific variation in eDNA shedding, detection and amplification and ensuring amplification of fish which were expected in all of them) to assess the importance of three key factors that determine eDNA capture efficiency, namely water volume, filtration method and DNA extraction kit.

**Material and methods**

**Study sites**

Water samples of various volumes were collected in April 2017 from three freshwater bodies (two lentic and one lotic) in Wales (UK): a small (15 m wide, 1 m deep) pond located at Swansea University, an artificial freshwater lake at Cardiff Bay and the River Tawe. Cardiff Bay is
situated at the confluence of the Rivers Taff and Ely, it is approximately 200 ha and was impounded in 1999 [45]. Water from the River Tawe was collected at the headwaters, close to the river source (latitude 51˚46'0.276" N, longitude 3˚46'35.514" W), and also at the river mouth (latitude 51˚42'08.9" N, longitude 3˚53'57.2" W). In the pond, water was collected at two different sampling points on opposite sides (latitude 51˚36'26.5" N, longitude 3˚58'52.5" W). The water samples in Cardiff Bay were collected from three different stations; the barrage (longitude 51˚26'48.7" N, latitude 3˚10'01.1" W) and Cardiff International White Waters (longitude 51˚26’52.6’N, latitude 3˚10’57.1’W).

**eDNA sampling procedure**

Three replicates were obtained from each water body, collected approximately 30 cm under the surface. Water samples were kept refrigerated and transported to the laboratory for filtration within four hours of collection. To minimize the risk of cross-contamination, disposable nitrile gloves were used and Nalgene polyethylene bottles were treated with 10% bleach, left for 5 min and thoroughly rinsed with sterile distilled water before sampling at each station. All filtration was conducted on the day of the sampling. Water was thoroughly mixed between sampling stations before filtration in order to have one uniform representation for each specific water body.

**eDNA capture and amplification efficiency experiment**

The study evaluated the effects of different filtered volume, filter type, and extraction kits (Fig 1) individually. The efficiency of the experiments was assessed by eDNA capture yield (ng/μL), PCR amplification (ng/μL) and amplification using qPCR (Cq values). DNA yield as well as the efficiency of PCR amplification were measured using a Qubit 1.0 fluorometer (Thermo Fisher Scientific Inc., UK) applying the high-sensitivity assay for DNA capture yield efficiencies and broad range assay for PCR products (Life Technologies, Carlsbad, CA, USA). Standard recommendations for work with eDNA were applied through all the study [1]. In total, 108 field samples were extracted, plus six and nine filtration and extraction control samples, respectively. The extraction and pre-PCR handling of eDNA water samples was carried out in a fume hood dedicated to eDNA analyses only. Before individual extractions, 10% bleach was used to clean up the fume hood as well as 45 min exposure to UV light.

The water volume experiment included 15 mL of volume without filtration based on ethanol- sodium acetate precipitation with the rest of volumes (100 mL, 250 mL, 1000 mL and 2000 mL) based only on filtering. All filtration was carried out using glass fibre filter and DNA was extracted using Qiagen kits. The filtration type experiment was based on 100 mL of water and DNA was extracted using the Qiagen extraction kit, comparing ethanol- sodium acetate precipitation with cellulose filtration and two other filter types, cellulose nitrate and glass fibre, without precipitation. The extraction kit experiment compared the Nexxtc Bacteria, Tissue and cell kit, the Nexxtc Blood kit and the Qiagen DNeasy Blood & Tissue kits, filtering 250 mL of water with glass fibre filter.

**Water filtration volume comparison**

For comparisons of water filtration volumes, three replicates of 15 mL, 100 mL, 250 mL, 1000 mL and 2000 mL water were collected at each site (Fig 1). For the 15 mL water samples we followed the protocol for ethanol- sodium acetate precipitation described in [30] by adding 1.5 mL sodium acetate (3 M) and 33 mL of absolute ethanol. The mixture was centrifuged at 5000 g for 45 min at 6˚C and the supernatant was discarded [46]. The precipitation itself was
Fig 1. Graphical representation of filtration volume, filtration type and extraction kit experiments.

https://doi.org/10.1371/journal.pone.0219218.g001
conducted on the day of water sampling, by centrifuging no more than four hours after collection. The falcon tubes with the DNA pellets were then stored at -20˚C without preservatives until the DNA extraction one week later. Negative control nuclease-free water was included.

The larger water volumes (100 mL, 250 mL, 1000 mL and 2000 mL) were filtered through the Advantec GA55 Borosilicate Glass Fibre Filters with 0.6 μm pore size (47 mm) (Fig 1). Each water sample was filtered through a filter funnel attached to a collection bottle and connected to the electronic vacuum pump Welch N. 2522C-02, with strength of 20 kpa for 15 s up to 75 s per sample. To avoid contamination, the filter funnel and handling tweezers were cleaned with a 10% bleach solution, rinsed with 99% molecular grade ethanol and then with sterile nuclease-free water between samples. For each different volume, a negative control consisting of nuclease-free water was used. For eDNA extraction, the Qiagen DNeasy Blood & Tissue DNA extraction kit (Qiagen GmbH, Hilden, Germany) was used. For the 15 mL method the Qiagen protocol for blood (spin protocol) was used whereas for the filtration methods we used the protocol for dried blood spots. The manufacturer’s protocol was followed in all cases, with the single modification of reducing the final elution volume to 50 μL in all three experimental designs. In total, 45 samples were used for water filtering comparison, 15 per water body, 3 for each of the five volume categories, with all samples amplified in duplicate for PCR and qPCR (Table B in S1 File).

**Filtration type comparison**

For the filtration type comparison, we used 100 mL of water and two different DNA capture methods: a pump filtration only method and syringe filtration with additional ethanol- sodium acetate precipitation. For the filtration only method, we used two different filter materials, Whatman Cellulose Nitrate Membrane Circle filters with 0.45 μm pore size (47 mm) and Advantec GA55 Borosilicate Glass Fibre Filters with 0.6 μm pore size (47 mm) (Fig 1). The second capture method was based on a combination of filtration using closed syringe filters Minisart cellulose syringe filters with 0.45 μm pore size (Sartorius, Germany) with additional ethanol- sodium acetate precipitation. For the syringe filtration, the water was pushed through by hand at an approximate flow rate of 50 mL per 30 s. After filtration, a mixture of 1350 μL absolute ethanol and 150 μL of sodium acetate was passed through the filters which were then centrifuged at 5000 g for 45 min at 6˚C. For the other two types of filters, filtration was carried out as above. DNA was purified with the Qiagen DNeasy Blood & Tissue DNA extraction kit. For the 100 mL syringe filtration method, the Qiagen DNA purification protocol for blood (spin protocol) was used, whereas for the other two filtration techniques we applied the protocol for dried blood spots, designed for the DNA isolation out of filter paper. In total, 27 samples were extracted for comparison, 9 per water body, 3 per each of the three filtration comparison types, all samples analysed in duplicates at both PCR and qPCR (Table C in S1 File).

**Extraction kit comparison**

Two hundred and fifty mL of water were collected and filtered through Advantec GA55 Borosilicate Glass Fibre Filters with 0.6 μm pore size (47 mm) for the extraction kit comparison (Fig 1). The 250 mL water volume was selected for practical reasons, as larger volumes result in reduced estimate error for all replicates [29]. The Qiagen DNeasy Blood & Tissue DNA extraction kit (protocol for dried blood spots) was compared to three additional kits all from Nexttec (Nexttec Biotechnologie GmbH, Germany): the 1-step DNA Isolation Kit for Tissues & Cells, 1-step DNA Isolation Kit for Blood (200 μL) and 1-step DNA Isolation Kit for Bacteria. The reason for selecting Nexxttec kits was based on the potential advantages of reduced
potential contamination, having a single step between the digestion of the sample and the final DNA elution. All extractions were carried out following the manufacturers’ instructions, with the only modification of reducing the elution volume to 50 μL. 36 samples in total were used for extraction kit comparison, 12 per each of the three water bodies, 3 per each of the four different extraction kits tests, with two replicates for PCR and qPCR amplification (Table D in S1 File).

**PCR amplification**

In order to overcome the potential specificity bias of species-specific primers [47] and to avoid differences based on single species representation in different lentic and lotic bodies, we used the vertebrate specific primer pair 12S-V5 developed by Riaz et al. [48], which amplifies a 144-bp long fragment of the 12s rRNA mtDNA gene and has been widely used [49–51]. The amplification reaction was performed in a total volume of 30 μL with, 12.5 μL Bioline BioMix Red PCR Mastermix (2X), 3 μL template, 1.5 μL of each primer (10 μM), adding sterile nuclease- free water to the final total volume. PCR conditions were as follow, 10 min at 95˚C, followed by 40 cycles of 10 s at 95˚C and 30 s at 52˚C 30 s and 72˚C for 30 s, with a final extension step at the 72˚C for 5 min. PCR products were visualised on a 2% agarose gel and quantified with Qubit 1.0 fluorometer. Positive controls were used for the evaluation of primer pair efficiency with DNA extracted from two different fish species commonly found in Tawe and Cardiff Bay, brown trout (*Salmo trutta*) and Atlantic salmon (*Salmo salar*). DNA was extracted from muscle or fin tissue from these target species using the Qiagen DNeasy Blood & Tissue DNA extraction kit. A negative control PCR with no DNA template was added at all PCR amplifications.

SYBR Green technology (Bio-Rad, US) was used in real-time PCR in a combination with 12S-V5 primer pair in a final reaction volume of 20 μL which included, 10 μL SsoAdvanced Universal SYBR Green Supermix (1x), 3 μL template, 0.4 μL of each 12S-V5 primer (10 μM) and 6.2 μL sterile nuclease- free water. The qPCR amplification was performed under the following conditions: 7 min at 95˚C, followed by 40 cycles of 10 s at 95˚C and 30 s at 95˚C and 30 s at 59˚C. Each of one of the three sampling replicates was amplified twice on a plate and final average Cq values of the duplicates was used for the statistical analysis. Each qPCR plate included three negative controls consisting of sterile nuclease- free water instead of the template. A standard curve with 8- point 10- fold dilutions with starting concentration of 1 ng/ μL of genomic *Salmo trutta* DNA was used. *S. trutta* was chosen for the standard curve as it represents one of the most common fish species in Welsh freshwater bodies [52].

**Cloning**

For species confirmation, four randomly selected samples from each water body and experimental design (twelve in total) were chosen and amplified with the 12S-V5 vertebrate primer pair using the same PCR protocol as above. The amplified PCR products (144 bp) were cloned into a pDRIVE Cloning Vector using Qiagen PCR cloning plus kit (Qiagen GmbH, Hilden, Germany) following the manufacturer’s recommendations. Three different concentrations of ligation- reaction mixture were plated on agar plates: 20 μL, 50 μL and 100 μL. Plasmid DNA was extracted using the Wizard Plus SV Minipreps DNA Purification kit (Promega, Madison, WI, USA). Sequencing was then carried out with T7 and Sp6 primers at the Institute of Biological, Environmental and Rural Sciences (IBERS), Aberystwyth. For sequencing 12, 7 and 12 clones were randomly selected from the river Tawe, the pond and Cardiff Bay respectively, with lower representation of pond samples due to low number of colonies.
Statistical analysis

For each response variable, eDNA capture yield (ng/μL), PCR (ng/μL) and qPCR (Ct values) amplification yields, three individual linear models were applied for each of the three experiments separately. For the volume experiment, volume and water body, including the interaction between them were used as predictors. For the filtration type experiment, water body and filtration type, were used as predictors including the interactions among them. For the extraction kit experiment, water body and extraction kit were used as predictors, including their interaction between them. Models with and without interactions between predictors for each of the three response variables (eDNA capture, PCR and qPCR) were compared based on AIC criteria using the ‘mass’ package. For the post-hoc analysis the ‘lsmeans’ package was used [53] based on Tukey contrasts for each experiment individually. Technical PCR and qPCR replicates were averaged before the analysis. Only samples with two technical working replicates were considered for further statistical analyses. Positive PCR and qPCR reactions without quantified DNA capture yield were only used for further comparison based on amplification efficiencies, excluding DNA yield. All statistical analyses were done with R, version 3.3.2.

Results

In total 108 samples were extracted from all three freshwater bodies excluding negative and positive controls. No amplification appeared in filtration and extraction negative controls during PCR and qPCR. All positive controls performed as expected, and species were confirmed by Sanger sequencing of 144 bp length products. R² values for the qPCR standard curve ranged from 0.95 to 1.00, and the efficiency ranged from 97 to 104%, with a slope between —3.3, -3.2. Average capture and amplification success rate including standard deviations for all three experiments presented separately for the DNA capture yield, PCR and qPCR efficiency are in Tables 1–3.

Linear models found differences among water bodies, based on DNA capture efficiency (ng/μL), for two out of the three individual experiments, water volume (F (2, 38) = 4.441, p = 0.020) and extraction kit (F (2, 26) = 7.065, p = 0.005), but not for the filtration type (F (2, 24) = 1.402, p = 0.274) (Table 1). Amplification rates in general did not differ between the water bodies for all of the linear models, with the following exception: PCR amplification in the extraction kit experiment (Table 1, F (2, 28) = 6.412, p = 0.006). Significant interactions were identified between water body and volume (F (8, 40) = 3.781, p = 0.003), with 2000 mL of water resulting in higher capture efficiency in the pond compared to all lower filtering volumes in the Tawe (Tukey’s Post-hoc test, p < 0.001), and (based on qPCR) between filter type and water body (F (4, 25) = 3.667, p = 0.024), with much higher efficiency of syringe filtration combined with ethanol precipitation compared to cellulose nitrate filtering in the pond (Tukey’s Post-hoc test, p < 0.001) (Table 1). Full linear models are reported in Table 1.

Water filtration volume comparison

In total 45 samples (15 per water body corresponding to three sampling replicates for each one of the five-volume categories) were processed (Table 2), of which all samples were used for the statistical analyses for DNA capture yield, 42 samples for the PCR and 39 samples for the qPCR. eDNA capture yield increased with increase in filtered volume (F (4, 40) = 2.137, p < 0.001), with the highest DNA yield obtained from 2000 mL water sampled (Tukey’s Post-hoc test, p < 0.001). There were significant differences between volumes for both amplifications (PCR, F (4, 37) = 6.049, p < 0.001; qPCR, F (4, 34) = 3.330, p = 0.020) with the most efficient qPCR amplification, being at 2000 mL of water filtered compared to 100 mL and 250 mL (Tukey’s Post-hoc test, p = 0.010). The largest water volume filtered for the duration of experiment (2 L) showed the highest DNA capture efficiency (0.406 ± 0.497 ng/μL), about tenfold
higher compared to the other methods, followed by the 1 L (Fig 2). The capture yield for the 15 mL category was low compared to filtration with only 0.027 ± 0.009 ng/μL yield (Table 2). There was a gradual increase in the eDNA capture efficiency from smallest 100 mL category up to largest, 2 L. The PCR amplification rate was the highest for the largest filtered volume tested with an average of 15.111 ± 2.473 ng/μL (Table 2). The amplification rate for the 15 mL ethanol-sodium acetate precipitation method was high compared to other filtered volumes (12.738 ± 4.203 ng/μL). The qPCR amplification efficiency for the 2 L category resulted in an average of 31.242 ± 0.699 cycles, comparatively similar to the 15 mL category with an average of 32.978 ± 1.896 cycles (Table 2).

### Table 1. Linear models analysing effects of filtration volume, filtration type and extraction kit in correlation to water body type on successful eDNA extraction and amplification for each of the experimental category separately, including comparison between models with and without an interaction term between the tested categories and water bodies.

| Model                     | Dependent variable | Predictors                              | Model output statistics | AIC  |
|---------------------------|--------------------|------------------------------------------|-------------------------|------|
| **Volume experiment**     | DNA capture efficiency = Volume * Water body | DNA capture yield (ng/μL) Volume x Water body | F (8, 30) = 3.781, p = 0.003 | -51.37 |
|                           | DNA capture yield (ng/μL) Volume x Water body | Water body | F (2, 38) = 4.441, p = 0.020 |
|                           | DNA capture yield (ng/μL) Volume | Volume | F (4, 40) = 2.137, p < 0.001 |
|                           | PCR efficiency = Volume * Water body | PCR efficiency (ng/μL) Volume x Water body | F (8, 37) = 1.327, p = 0.275 | 237.8 |
|                           | PCR efficiency (ng/μL) Volume | Water body | F (2, 35) = 1.073, p = 0.356 |
|                           | PCR efficiency = Volume | Volume | F (4, 37) = 6.447, p < 0.001 |
|                           | qPCR efficiency = Volume * Water body | qPCR (Cq values) Volume x Water body | F (8, 24) = 1.167, p = 0.359 | 160 |
|                           | qPCR (Cq values) Volume | Water body | F (2, 32) = 1.722, p = 0.2 |
|                           | qPCR efficiency = Volume | Volume | F (4, 34) = 3.602, p = 0.019 |

**Filtration type experiment**

| DNA capture efficiency = Filtration type * Water body | DNA capture yield (ng/μL) Filtration type X Water body | Filtration type X Water body | F (4,24) = 2.287, p = 0.105 | -87.53 |
| Filtration type | Filtration type X Water body | Water body | F (2,24) = 4.294, p = 0.032 |
| Filtration type | Filtration type X Water body | Filtration type | F (2,24) = 1.402, p = 0.274 |
| DNA capture efficiency = Filtration type | DNA capture yield (ng/μL) Filtration type | Filtration type | F (2,24) = 3.379, p = 0.05 | 140.23 |
| PCR efficiency = Filtration type * Water body | PCR efficiency (ng/μL) Filtration type X Water body | Filtration type X Water body | F (4,25) = 0.737, p = 0.580 |
| PCR efficiency = Filtration type | PCR efficiency (ng/μL) Filtration type | Filtration type | F (2,25) = 3.990, p = 0.037 |
| qPCR efficiency = Filtration type * Water body | qPCR (Cq values) Filtration type X Water body | Filtration type X Water body | F (4,25) = 3.667, p = 0.024 |
| qPCR efficiency = Filtration type | qPCR (Cq values) Filtration type | Filtration type | F (2,25) = 5.845, p = 0.011 |
| qPCR efficiency = Filtration type | qPCR (Cq values) Filtration type | Filtration type | F (2,25) = 3.501, p = 0.047 |

**Extraction kit experiment**

| DNA capture efficiency = Extraction kit * Water body | DNA capture yield (ng/μL) Extraction kit X Water body | Extraction kit X Water body | F (6, 20) = 2.363, p = 0.069 | -45.65 |
| Extraction kit | Extraction kit X Water body | Water body | F (2, 26) = 7.065, p = 0.005 |
| Extraction kit | Extraction kit X Water body | Extraction kit | F (3, 29) = 10.657, p = 0.001 |
| PCR efficiency = Extraction kit * Water body | PCR efficiency (ng/μL) Extraction kit X Water body | Extraction kit X Water body | F (6, 22) = 2.162, p = 0.086 |
| PCR efficiency = Extraction kit * Water body | PCR efficiency (ng/μL) Extraction kit | Extraction kit | F (2, 28) = 6.412, p = 0.006 |
| PCR efficiency = Extraction kit * Water body | PCR efficiency (ng/μL) Extraction kit | Extraction kit | F (3, 30) = 4.159, p = 0.018 |
| qPCR efficiency = Extraction kit * Water body | qPCR (Cq values) Extraction kit X Water body | Extraction kit X Water body | F (6, 20) = 2.042, p = 0.107 |
| qPCR efficiency = Extraction kit * Water body | qPCR (Cq values) Extraction kit | Extraction kit | F (2, 26) = 3.380, p = 0.054 |
| qPCR efficiency = Extraction kit * Water body | qPCR (Cq values) Extraction kit | Extraction kit | F (3, 28) = 0.299, p = 0.825 |

https://doi.org/10.1371/journal.pone.0219218.t001
Differences in eDNA capture yield (ng/μL), and amplification efficiencies by PCR (ng/μL) and qPCR (Cq) in the five different categorical groups for filtration volume experiment (15 mL, 100 mL, 250 mL, 1000 mL and 2000 mL), where each category is represented by three sampling replicates per three water bodies (9). For the amplification efficiency the technical duplicates of each sampling replicate were averaged before plotting. The 15 mL volume is based on ethanol- sodium acetate precipitation whereas the rest are based on water filtration. Filtration took place using glass fibre filter and eDNA was extracted with the Qiagen extraction kit.

Filtration type comparison

Twenty-seven samples were used for the comparison between the filtration types excluding three negative filtrations and one extraction controls, with nine samples representing each individual water body, as sample triplicates for each of the three individual filter types were examined (Table 3). For statistical analysis 25 samples were evaluated for DNA capture yield, 26 for PCR and 26 for qPCR. We found statistically significant differences between filtration type categories for DNA capture yield (F (2, 24) = 4.294, p = 0.032), PCR amplification (F (2, 25) = 4.362, p = 0.024) and qPCR (F (2, 25) = 5.845, p = 0.011). DNA extraction yield was the highest for the ethanol- sodium acetate precipitation in combination with filtration (0.070 ± 0.058 ng/μL) in comparison to other two solely filtration procedures (Table 3). Both, filtration only methods without ethanol–sodium acetate precipitation performed poorly measured by DNA capture yield (Fig 3). Cellulose nitrate filters were the only filtration method where some of the filters failed to yield any eDNA and those samples were excluded from further statistical analysis (Table C in S1 File). PCR amplification efficiency using the combined method of ethanol–sodium acetate precipitation and filtration was the highest as well. This may indicate that the combined method is the most suitable for the capture and amplification of eDNA from freshwater environments.

Table 2. Comparison of water volumes by eDNA capture and amplification efficiencies for volume experiment using glass fibre filter (0.6 μm) and Qiagen extraction kit for each individual response, DNA capture yield (ng/μL), PCR (ng/μL) and qPCR (Cq).

| Water body (n. of samples)          | Volume | Mean DNA capture yield (±SD) (ng/μL) | Mean PCR efficiency (±SD) (ng/μL) | Mean qPCR efficiency (±SD) (Cq) |
|-------------------------------------|--------|-------------------------------------|----------------------------------|-------------------------------|
| Cardiff Bay (15), Pond (15), Tawe (15) | 15 mL  | 0.027 ± 0.009                        | 12.738 ± 4.203                   | 32.978 ± 1.896               |
|                                     | 100 mL | 0.044 ± 0.045                        | 8.813 ± 3.383                    | 34.194 ± 1.236               |
|                                     | 250 mL | 0.040 ± 0.019                        | 8.156 ± 4.797                    | 33.960 ± 1.983               |
|                                     | 1000 mL| 0.087 ± 0.131                        | 13.386 ± 1.793                   | 33.683 ± 1.893               |
|                                     | 2000 mL| 0.406 ± 0.497                        | 15.111 ± 2.473                   | 31.242 ± 0.699               |

*The 15 mL volume within the volume experiment is based solely on ethanol- sodium acetate precipitation. Water bodies (Cardiff Bay, Tawe river and Pond) including number of sampling replicates per water body (15), total number of samples (45) and categories tested (15, 100, 250, 1000 and 2000 mL) are stated.

https://doi.org/10.1371/journal.pone.0219218.t002

Differences in eDNA capture yield (ng/μL), and amplification efficiencies by PCR (ng/μL) and qPCR (Cq) in the five different categorical groups for filtration volume experiment (15 mL, 100 mL, 250 mL, 1000 mL and 2000 mL), where each category is represented by three sampling replicates per three water bodies (9). For the amplification efficiency the technical duplicates of each sampling replicate were averaged before plotting. The 15 mL volume is based on ethanol- sodium acetate precipitation whereas the rest are based on water filtration. Filtration took place using glass fibre filter and eDNA was extracted with the Qiagen extraction kit.

Table 3. Comparison of filtration methods for eDNA capture and amplification efficiencies for filtration type experiment for 100 mL water filtered using Qiagen extraction kit for each individual response DNA capture yield (ng/μL), PCR (ng/μL) and qPCR (Cq).

| Water body (n. of samples) | Filtration type | Mean DNA capture yield (±SD) (ng/μL) | Mean PCR efficiency (±SD) (ng/μL) | Mean qPCR efficiency (±SD) (Cq) |
|----------------------------|-----------------|-------------------------------------|----------------------------------|-------------------------------|
| Cardiff Bay (9), Pond (9), Tawe (9) | Cellulose nitrate | 0.023 ±0.019                       | 8.645 ± 1.207                   | 35.626 ± 2.341               |
|                             | Glass fibre filter | 0.022 ±0.013                      | 9.280 ± 3.293                   | 34.115 ± 1.157               |
|                             | Syringe filtration + precipitation | 0.070 ±0.058                      | 12.593 ± 3.455                 | 33.253 ± 1.925               |

Water bodies (Cardiff Bay, Tawe river and Pond) including number of sampling replicates per water body (9), total number of samples (27) and categories tested (cellulose nitrate, glass fibre filter and syringe filtration + ethanol–sodium acetate precipitation) are stated.

https://doi.org/10.1371/journal.pone.0219218.t003
Fig 2. eDNA capture yield (ng/μL) and amplification efficiencies by filtration volume experiment.

https://doi.org/10.1371/journal.pone.0219218.g002
Fig 3. Filtration type experiment evaluating eDNA capture yield (ng/μL) and amplification efficiencies.

https://doi.org/10.1371/journal.pone.0219218.g003
syringe filtration and precipitation yielded the highest DNA concentrations (average value of 12.593 ± 3.45 ng/μL) (Table 3). A slightly better amplification performance was produced by glass fibre filter (average value of 9.280 ± 3.293 ng/μL) in comparison to cellulose nitrate filter with an average of 0.635 ng/μL lower amplification rate. The syringe filtration in a combination with ethanol- sodium acetate precipitation resulted in low Cq values with an average of 33.235 ± 1.925 cycles evaluated by qPCR. QPCR provided similar results to PCR regarding performance of the glass fibre filter versus the cellulose nitrate filter with an average of 1.511 cycles higher for the glass fibre filter (Table 3).

Differences in eDNA capture as well as amplification by PCR and qPCR using three different filtration methods (C- Cellulose nitrate filter, G- Glass fibre filter, S- Syringe filter with a combination of ethanol- sodium acetate precipitation). Each experimental category is represented by three sampling replicates per three water bodies (9). For the amplification efficiencies the technical duplicates of each sampling replicate were averaged before plotting. The lowest Cq value corresponds to the highest efficiency.

**Extraction kit comparison**

Thirty-six samples were extracted for the comparison between the filtration types, as well as one negative filtration and four negative extraction controls for each of the extraction kits tested (Table 4). Of these, twelve samples were used for each individual water body as sample triplicates were used for each individual extraction kit. In total, the statistical analysis based on DNA capture yield used 32 samples, 34 for the PCR and 32 for the qPCR. A model including both, the experimental groups and water bodies, identified significant differences between extraction kits for DNA capture yield (F (3, 29) = 10.657, p = 0.001) and PCR amplification (F (3, 30) = 4.159, p = 0.018), with the highest capture and amplification rate corresponding to the Nexxtec Blood kit (Tukey’s Post-hoc test, p < 0.001), but there were no significant differences for qPCR (F (3, 28) = 0.299, p = 0.825) (Table 1). All Nexxtec kits were generally more efficient with regards to DNA capture in comparison to Qiagen (Fig 4). Between the Nexxtec kits the most efficient one appears to be the kit designed for blood samples with much higher efficiency compared to other two, 0.206 ng/μL higher DNA capture yield on average. The 1—step Nexxtec DNA Isolation Kit for Blood proved particularly efficient with samples from Cardiff Bay with DNA capture yields of 0.511 ± 0.229 ng/μL and had on average 4.438 ng/μL higher amplification rate compared to other Nexxtec kits (Table 4).

Estimation of DNA capture extraction efficiency and amplification evaluated by PCR and qPCR, comparing between the following extraction kits Nexxtec bacteria, Nexxtec blood, Nexxtec Tissue and Qiagen. Each experimental category is represented by three sampling replicates per three water bodies (9), number of sampling replicates per water body (12), total number of samples = 36 and extraction kits tested (Nexxtec Blood, Nexxtec Bacteria, Nexxtec Tissue and Qiagen).

**Table 4. Comparison of extraction kits for eDNA capture and amplification efficiencies for extraction kit experiment with 250 ml of water filtered using a glass fibre filter (0.6 μm) for each individual response DNA capture yield (ng/μL), PCR (ng/μL) and qPCR (Cq).**

| Water body (n. of samples) | Extraction kit     | Mean DNA capture yield (±SD) (ng/μL) | Mean PCR efficiency (±SD) (ng/μL) | Mean qPCR efficiency (±SD) (Cq) |
|---------------------------|--------------------|--------------------------------------|----------------------------------|---------------------------------|
| Cardiff Bay (12), Pond (12), Tawe (12) | Nexxtec Blood | 0.284 ±0.232 | 10.080 ±1.603 | 33.929 ±2.045 |
|                           | Nexxtec Bacteria  | 0.095 ±0.068 | 4.784 ±4.133 | 34.392 ±1.841 |
|                           | Nexxtec Tissue    | 0.061 ±0.051 | 6.607 ±4.721 | 33.551 ±1.848 |
|                           | Qiagen            | 0.039 ±0.018 | 8.156 ±4.797 | 33.949 ±1.975 |

https://doi.org/10.1371/journal.pone.0219218.t004
Fig 4. eDNA capture yield (ng/μL) and amplification efficiencies by extraction kit comparison.

https://doi.org/10.1371/journal.pone.0219218.g004
Species composition

Sequencing of the cloned PCR products indicated that the three dominant fish species found in each individual water body were European bullhead (*Cottus gobio*) in the Tawe River, three-spinned stickleback (*Gasterosteus aculeatus*) in the pond and European carp (*Cyprinus carpio*) in Cardiff Bay, irrespective of the sampling technique used (Fig 5, Table A in S1 File). As 12S-V5 are vertebrate primers, there were also human (*Homo sapiens*), domestic pig (*Sus scrofa domesticus*) and common mallard (*Anas platyrhynchos*) sequences among the results. From 11 clones in the River Tawe, seven belonged to European bullhead, two were identified as *Anas platyrhynchos* and two remained unidentified. In the pond, three sequences belonged to *Gasterosteus aculeatus* and four remaining cloning sequences remained unidentified. In Cardiff Bay, five sequences belonged to *Cyprinus carpio*, five to *Homo sapiens*, and one to *Sus scrofa domesticus*.
Graphical representation of the most successful sampling method for all response variables tested for each water body separately. Pie charts indicate species proportion from total number (n) of sequenced cloned samples, River Tawe (11), lake Cardiff Bay (11) and Swansea University pond (7). The sequences that were not identified are marked as (unknown). Only one fish species was identified by Sanger sequencing in each water body using general vertebrate primers. The 2000 mL filtered water volume and syringe filtering with ethanol- sodium acetate precipitation technique appeared to be the most successful accounting for both, DNA capture and amplification. The selected images of fish were collected from the websites (Wikimedia, Titu. do, Flickr) marked with permission for reuse and modifications. The images of water bodies are personal copyright material.

Discussion

The results from three different comparisons testing the effects of filtration volume, filtration type and extraction procedure, evaluated by DNA capture yield and amplification efficiencies highlight the importance of selecting the appropriate sampling method due to their variable efficiencies. Our results showed differences in capture yield among the three freshwater bodies depending on the volume and extraction kit used. This study indicates that a consistent eDNA sampling procedure is needed to be used in different freshwater bodies for species detection and quantitative assessment to be comparable across them.

It has been previously shown that a combination of different capture and extraction methods can result in different success rates of eDNA metabarcoding for different target groups [20], using universal primers [3, 5, 54]. Our approach, a novel combination of ethanol- sodium acetate precipitation with filtration, worked well in lentic and lotic water bodies with a high efficiency, easiness of handling, low cost, low chances of contamination and practicality. This method can be a reliable tool for eDNA species-specific assessments, confirming presence/absence of certain species [55], as well as a sampling approach to determine community composition based on metabarcoding.

We examined the influence of filtration volume, filtration type, filtration method and type of extraction kit on capture yield and amplification efficiencies. DNA capture as well as amplification appeared to be very sensitive to changes in the volume of water filtered, as previously reported [20]. The efficiency of eDNA capture yield and amplification success rate largely differed between volumes. It would thus be recommended to filter as much of the water as possible, although the size distribution of particles in the aquatic environment can be a crucial factor determining the filter pore size and consequently the most feasible volume of water to be filtered [56]. Size of filtered particles [56], contamination risk [55] and feasibility of the proposed sampling (depending on location and proximity to the laboratory), can result in practical limitations in the maximum amount of water that is possible to filter [56]. DNA capture efficiency is an important evaluator of sampling technique used as it reflects the presence of the whole DNA within the sample. Another important factor is the number of replicates used for each individual evaluation, as the differences between the sampling triplicates were the most obvious in the DNA capture yield, where the whole extracted DNA, not just vertebrate eDNA, was quantified. In relation to this, while the highest eDNA capture rate corresponded to Cardiff Bay during the extraction kit experiment, it had the lowest amplification success (PCR and qPCR), potentially explained by high abundance of non-vertebrate DNA. This variability could be due to lack of power and more replicates would be recommended to increase reproducibility.

The most commonly used filter materials in eDNA studies are glass fibre filters [26, 31, 35] and cellulose nitrate filters [29, 32, 37] with different pore sizes, where larger pore sizes allow
larger filtered water volumes and smaller pore sizes capture more particles but limit volume and speed of filtration [56]. Glass fibre filter with larger pore size resulted in slightly higher amplification efficiency compared to cellulose nitrate filter with smaller pore size, a contrasting result to previous ones, where normally larger filter pore size required larger volumes of water filtered [22]. The choice for the material of the filter type used depends as well on the practicality of usage during DNA extraction as filtration materials differ greatly. Judging by the easiness of filter handling and sample preparation for extraction, glass fibre material was preferred here. The difference between material types was not as high as the difference between capture techniques, with much higher efficiency of ethanol-sodium acetate precipitation in combination with filtration compared to filtration only.

The smallest water volume tested, based only on ethanol-sodium acetate precipitation (15 mL) provided solid amplification rates despite small volume. Thus, the newly proposed 100 mL syringe filtration with ethanol-sodium acetate precipitation method combines the strength of both techniques: the portability and easiness of the ethanol-sodium acetate precipitation while increasing the volume filtered and decreasing contamination risk by minimising filter handling. The proposed syringe filtration method appears to be highly efficient, affordable and reliable and it is thus an upgraded method from the one proposed by [30]. High efficiency of syringe filters compared to other filtration techniques has been shown with the use of Sterivex-GP polyethersulfone syringe filters, but it is a more costly alternative to the syringe filters used in this study [22].

The extraction kit seems to be the least important factor when it comes to selection of sampling techniques for eDNA capture. On several occasions extraction procedures based on usage of commercial kits resulted in no differences [25], whereas in other cases there has been shown significant variations [20, 23, 25]. The higher DNA capture efficiency and PCR amplification rates were provided by Nexxtec Blood kit, for which a single step before DNA elution highly minimises risk for the contamination. There was no difference between the extraction kits based on capture rate and the qPCR assessment.

Species-specific sequencing identified one dominant fish species per water body independently of the sampling technique used. This is likely due to the number of samples cloned only detecting the most abundant species in each area (Wharf Angling Club, 2018). Cardiff Bay is highly associated to human activities and the presence of human DNA is therefore not surprising. Mammal (including human) and avian DNA presence is common in eDNA studies utilising universal primers [57, 58] and all our negative filtration, extraction and PCR controls ensured that its origin was not laboratory contamination.

Our study contributes towards the understanding of the role of different sampling and extraction methods on the efficiencies of eDNA capture techniques. Focusing on well-known vertebrate primers [49–51, 59, 60] to avoid species-specific bias allowed us to compare efficiencies in three different water bodies with distinctive community composition, that can potentially introduce drawbacks assessing eDNA presence/absence using qPCR, with a preferred species specific assay design. There was no difference between the PCR or qPCR success rate for the two most evident differential factors, the water bodies and volume, whereas filtration type and extraction kit differed greatly. Dissimilarities between capture and extraction techniques between pond, lake and river, highlight the importance of other abiotic aspects affecting eDNA yield such as acidity, substrate material and hydrological dynamics [39, 61, 62], including seasonality [63], which can majorly affect eDNA detection rates in different situations.

In summary, our study indicates that the main source of variation in the eDNA capture and amplification efficiencies is the sampling technique. Our results indicate that a careful sampling plan selecting the most efficient eDNA sampling protocol is essential and suggest that sampling and filtering the largest feasible volume is the best strategy. However, a syringe
filtration through a 0.45 μm cellulose filter, combined with ethanol- sodium acetate precipitation is an alternative with low contamination risk/ high yield that can be easily used both in lotic and lentic environments with minimal sampling effort.

Supporting information

S1 File. qPCR melt curve plots of all three experiments volume, filter type and extraction kit carrier out in Tawe river (Figure A). qPCR melt curve plots of all three experiments volume, filter type and extraction kit carrier out in Cardiff Bay (Figure B). qPCR melt curve plots of all three experiments volume, filter type and extraction kit carrier out in Swansea University pond (Figure C). Identification of species in each of the water bodies pond, lake and river defined by capture and extraction technique, based on cloning and Sanger sequencing (Table A). Dataset for the filtration volume experiment combining all sampling triplicates from all three water bodies used for the statistical analysis, based on glass fibre filtration and Qiagen extraction kit (Table B). Dataset for the filtration type experiment combining all sampling triplicates from all three water bodies used for the statistical analysis, based on 100 mL filtered volume and Qiagen extraction kit (Table C). Dataset for the extraction kit experiment combining all sampling triplicates from all three water bodies used for the statistical analysis, based on 250 mL filtered volume using glass fibre filter (Table D).

(DOCX)

Acknowledgments

We would like to thank to Cardiff Harbour Authority, especially to David Hall for organisation of water collections within Cardiff Bay and a provision of boat facilities. We also thank Deiene Rodriguez Barreto and Tamsyn M. Uren Webster for their generous advice concerning laboratory procedures.

Author Contributions

Conceptualization: Teja Petra Muha, Sofia Consuegra.
Data curation: Teja Petra Muha.
Formal analysis: Teja Petra Muha.
Funding acquisition: Carlos Garcia de Leaniz, Sofia Consuegra.
Methodology: Teja Petra Muha, Chloe Victoria Robinson.
Supervision: Carlos Garcia de Leaniz, Sofia Consuegra.
Validation: Teja Petra Muha, Chloe Victoria Robinson.
Writing – original draft: Teja Petra Muha.
Writing – review & editing: Carlos Garcia de Leaniz, Sofia Consuegra.

References

1. Goldberg CS, Turner CR, Deiner K, Klymus KE, Thomsen PF, Murphy MA, et al. Critical considerations for the application of environmental DNA methods to detect aquatic species. Methods in Ecology and Evolution. 2016; 7(11):1299–307.
2. Deiner K, Bik HM, Mächler E, Seymour M, Lacoursière-Roussel A, Alternatt F, et al. Environmental DNA metabarcoding: transforming how we survey animal and plant communities. Molecular ecology. 2017; 26(21):5872–95. https://doi.org/10.1111/mec.14350 PMID: 28921802
3. Valentini A, Taberlet P, Miaud C, Civade R, Herder J, Thomsen PF, et al. Next-generation monitoring of aquatic biodiversity using environmental DNA metabarcoding. Molecular Ecology. 2016; 25(4):929–42. https://doi.org/10.1111/mec.13428 PMID: 26479867

4. Nathan LM, Simmons M, Wegleitner BJ, Jerde CL, Mahon AR. Quantifying environmental DNA signals for aquatic invasive species across multiple detection platforms. Environmental science & technology. 2014; 48(21):12800–6.

5. Evans NT, Olds BP, Renshaw MA, Turner CR, Li Y, Jerde CL, et al. Quantification of mesocosm fish and amphibian species diversity via environmental DNA metabarcoding. Molecular ecology resources. 2016; 16(1):29–41. https://doi.org/10.1111/1755-0998.12433 PMID: 26032773

6. Stewart K, Ma H, Zheng J, Zhao J. Using environmental DNA to assess population-wide spatiotemporal reserve use. Conservation Biology. 2017; 31(5):1173–82. https://doi.org/10.1111/cobi.12910 PMID: 28221696

7. Doi H, Uchii K, Takahara T, Matsushashi S, Yamanaka H, Minamoto T. Use of droplet digital PCR for estimation of fish abundance and biomass in environmental DNA surveys. PloS one. 2015; 10(3):e0122763. https://doi.org/10.1371/journal.pone.0122763 PMID: 25799582

8. Biggs J, Ewald N, Valentini A, Gaboriaud C, Dejean T, Griffiths RA, et al. Using eDNA to develop a national citizen science-based monitoring programme for the great crested newt (Triturus cristatus). Biological Conservation. 2015; 183:19–28.

9. Dejean T, Valentini A, Miquel C, Taberlet P, Bellemain E, Miaud C. Improved detection of an alien invasive species through environmental DNA barcoding: the example of the American bullfrog Lithobates catesbeianus. J Appl Ecol. 2012; 49(4):953–9.

10. Smart AS, Tingley R, Weeks AR, van Rooyen AR, McCarthy MA. Environmental DNA sampling is more sensitive than a traditional survey technique for detecting an aquatic invader. Ecological applications. 2015; 25(7):1944–52. PMID: 26591459

11. Harper LR, Lawson Handley L, Hahn C, Boonham N, Rees HC, Gough KC, et al. Needle in a haystack? A comparison of eDNA metabarcoding and targeted qPCR for detection of the great crested newt (Triturus cristatus). Ecology and evolution. 2018; 8(12):6330–41. https://doi.org/10.1002/ece3.4013 PMID: 29988445

12. Pilliod DS, Goldberg CS, Arkle RS, Waits LP. Factors influencing detection of eDNA from a stream-dwelling amphibian. Molecular Ecology Resources. 2014; 14(1):109–16. https://doi.org/10.1111/1755-0998.12159 PMID: 24034561

13. Lacoursière-Roussel A, Rosabal M, Bernatchez L. Estimating fish abundance and biomass from eDNA concentrations: variability among capture methods and environmental conditions. Molecular ecology resources. 2016; 16(6):1401–14. https://doi.org/10.1111/1755-0998.12522 PMID: 26946353

14. Dyer RJ, Roderique BA. Development and Testing of Environmental DNA (eDNA) Protocols for the Endangered James Spiny mussel (Pleurobema collina). Virginia Transportation Research Council, 2017.

15. Demeke T, Jenkins GR. Influence of DNA extraction methods, PCR inhibitors and quantification methods on real-time PCR assay of biotechnology-derived traits. Analytical and bioanalytical chemistry. 2010; 396(6):1977–90. https://doi.org/10.1007/s00216-009-3150-9 PMID: 19789856

16. Rees HC, Maddison BC, Midlleditch DJ, Patmore JR, Gough KC. The detection of aquatic animal species using environmental DNA—a review of eDNA as a survey tool in ecology. Journal of Applied Ecology. 2014; 51(5):1450–9.

17. Shokralla S, Spall JL, Gibson JF, Hajibabaee M. Next-generation sequencing technologies for environmental DNA research. Molecular Ecology. 2012; 21(8):1794–805. https://doi.org/10.1111/j.1365-294X.2012.05538.x PMID: 22466820

18. Bohmann K, Evans A, Gilbert MTP, Carvalho GR, Creer S, Knapp M, et al. Environmental DNA for wildlife biology and biodiversity monitoring. Trends in Ecology & Evolution. 2014; 29(6):358–67.

19. Shokralla S, Gibson J, King I, Baird D, Janzen D, Hallwachs W, et al. Environmental DNA barcode sequence capture: targeted, PCR-free sequence capture for biodiversity analysis from bulk environmental samples. bioRxiv. 2016:087437.

20. Deiner K, Walser J-C, Mächler E, Altermatt F. Choice of capture and extraction methods affect detection of freshwater biodiversity from environmental DNA. Biological Conservation. 2015; 183:53–63. http://dx.doi.org/10.1016/j.biocon.2014.11.018,

21. Turner CR, Miller DJ, Coyne KJ, Corush J. Improved methods for capture, extraction, and quantitative assay of environmental DNA from Asian bigheaded carp (Hypophthalmichthys spp.). Plos one. 2014; 9(12):e114329. https://doi.org/10.1371/journal.pone.0114329 PMID: 25474207

22. Spens J, Evans AR, Halfmaerten D, Knudsen SW, Sengupta ME, Mak SS, et al. Comparison of capture and storage methods for aqueous macrobial eDNA using an optimized extraction protocol: advantage of enclosed filter. Methods in Ecology and Evolution. 2017; 8(5):635–45.
23. Hinlo R, Gleeson D, Lintemans M, Furlan E. Methods to maximise recovery of environmental DNA from water samples. PloS one. 2017; 12(6):e0179251. https://doi.org/10.1371/journal.pone.0179251 PMID: 28604830

24. Williams KE, Huyvaert KP, Piaggio AJ. No filters, no fridges: a method for preservation of water samples for eDNA analysis. BMC research notes. 2016; 9(1):298.

25. Djurhuus A, Port J, Closek CJ, Yamahara KM, Romero-Maraccini O, Walz KR, et al. Evaluation of Filtration and DNA Extraction Methods for Environmental DNA Biodiversity Assessments across Multiple Trophic Levels. Frontiers in Marine Science. 2017; 4:314.

26. Wilcox TM, McKelvey KS, Young MK, Jane SF, Lowe WH, Whiteley AR, et al. Robust detection of rare species using environmental DNA: the importance of primer specificity. PloS one. 2013; 8(3):e59520. https://doi.org/10.1371/journal.pone.0059520 PMID: 23555689

27. Takahara T, Minamoto T, Doi H. Using Environmental DNA to Estimate the Distribution of an Invasive Fish Species in Ponds. PLoS ONE. 2013; 8(2):e56584. https://doi.org/10.1371/journal.pone.0056584 PMID: 23437177

28. Takahara T, Minamoto T, Yamanaka H, Doi H, Kawabata Zi. Estimation of fish biomass using environmental DNA. PloS one. 2012; 7(4):e35868. https://doi.org/10.1371/journal.pone.0035868 PMID: 22563411

29. Pilliod DS, Goldberg CS, Arkle RS, Waits LP. Estimating occupancy and abundance of stream amphibians using environmental DNA from filtered water samples. Canadian Journal of Fisheries and Aquatic Sciences. 2013; 70(8):1123–30.

30. Ficetola GF, Miaud C, Pompanon F, Taberlet P. Species detection using environmental DNA from water samples. Biology letters. 2008; 4(4):423–5. https://doi.org/10.1098/rsbl.2008.0118 PMID: 18400683

31. Jerde CL, Mahon AR, Chadderton WL, Lodge DM. “Sight-unseen” detection of rare aquatic species using environmental DNA. Conservation Letters. 2011; 4(2):150–7.

32. Goldberg CS, Sepulveda A, Ray A, Baumgardt J, Waits LP. Environmental DNA as a new method for early detection of New Zealand mudsnails (Potamopyrgus antipodarum). Freshwater Science. 2013; 32 (3):792–800.

33. Turner CR, Barnes MA, Xu CC, Jones SE, Jerde CL, Lodge DM. Particle size distribution and optimal capture of aqueous macrobial eDNA. Methods in Ecology and Evolution. 2014; 5(7):676–84.

34. Minamoto T, Naka T, Moji K, Maruyama A. Techniques for the practical collection of environmental DNA: filter selection, preservation, and extraction. Limnology. 2016; 17(1):23–32.

35. Janosik AM, Johnston CE. Environmental DNA as an effective tool for detection of imperiled fishes. Environmental biology of fishes. 2015; 98(8):1889–93.

36. Goldberg CS, Sepulveda A, Ray A, Baumgardt J, Waits LP. Environmental DNA as a new method for early detection of New Zealand mudsnails (Potamopyrgus antipodarum). 2013.

37. Goldberg CS, Pilliod DS, Arkle RS, Waits LP. Molecular detection of vertebrates in stream water: a demonstration using Rocky Mountain tailed frogs and Idaho giant salamanders. PloS one. 2011; 6(7):e22746. https://doi.org/10.1371/journal.pone.0022746 PMID: 21818382

38. Rius M. Optimising the detection of marine taxonomic richness using environmental DNA metabarcoding: the effects of filter material, pore size and extraction method. Metabarcoding and Metagenomics. 2018; 2(e28963).

39. Seymour M, Durance I, Cosby BJ, Ransom-Jones E, Deiner K, Ormerod SJ, et al. Acidity promotes degradation of multi-species environmental DNA in lotic mesocosms. Communications biology. 2018; 1 (1):4.

40. Renshaw MA, Olds BP, Jerde CL, McVeigh MM, Lodge DM. The room temperature preservation of filtered environmental DNA samples and assimilation into a phenol–chloroform–isoamyl alcohol DNA extraction. Molecular ecology resources. 2015; 15(1):168–76. https://doi.org/10.1111/1755-0998.12281 PMID: 24834966

41. Roh C, Villatte F, Kim B-G, Schmid RD. Comparative study of methods for extraction and purification of environmental DNA from soil and sludge samples. Applied biochemistry and biotechnology. 2006; 134 (2):97–112. PMID: 16943632

42. Eichmiller JJ, Miller LM, Sorensen PW. Optimizing techniques to capture and extract environmental DNA for detection and quantification of fish. Molecular ecology resources. 2016; 16(1):56–68. https://doi.org/10.1111/1755-0998.12421 PMID: 25919417

43. Whitehouse CA, Hottel HE. Comparison of five commercial DNA extraction kits for the recovery of Francisella tularensis DNA from spiked soil samples. Molecular and cellular probes. 2007; 21(2):92–6. https://doi.org/10.1016/j.mcp.2006.08.003 PMID: 17011748
44. Ficetola GF, Taberlet P, Coissac E. How to limit false positives in environmental DNA and metabarcoding? Molecular ecology resources. 2016; 16(3):604–7. https://doi.org/10.1111/1755-0998.12508 PMID: 27062589

45. Burton NH, Rehfisch MM, Clark NA. Impacts of disturbance from construction work on the densities and feeding behavior of waterbirds using the intertidal mudflats of Cardiff Bay, UK. Environmental Management. 2002; 30(6):0865–71.

46. Valiere N, Taberlet P. Urine collected in the field as a source of DNA for species and individual identification. Molecular Ecology. 2000; 9(12):2150–2. PMID: 11123627

47. Tedersoo L, Anslan S, Bahram M, Põlme S, Rii T, Liiv I, et al. Shotgun metagenomes and multiple primer pair-barcode combinations of amplicons reveal biases in metabarcoding analyses of fungi. MycoKeys. 2015; 10:1.

48. Riaz T, Shehzad W, Viani A, Pompanon F, Taberlet P, Coissac E. ecoPrimers: inference of new DNA barcode markers from whole genome sequence analysis. Nucleic Acids Research. 2011; 39(21):e145–e. https://doi.org/10.1093/nar/gkr732 PMID: 21930509

49. Kelly RP, Port JA, Yamahara KM, Crowder LB. Using environmental DNA to census marine fishes in a large mesocosm. PloS one. 2014; 9(1):e86175. https://doi.org/10.1371/journal.pone.0086175 PMID: 24454960

50. Miya M, Sato Y, Fukunaga T, Sado T, Poulsen J, Sato K, et al. MiFish, a set of universal PCR primers for metabarcoding environmental DNA from fishes: detection of more than 230 subtropical marine species. Royal Society open science. 2015; 2(7):150088. https://doi.org/10.1098/rsos.150088 PMID: 26687265

51. Port JA, O’Donnell JL, Romero-Maracccini OC, Leary PR, Litvin SY, Nickols KJ, et al. Assessing vertebrate biodiversity in a kelp forest ecosystem using environmental DNA. Molecular ecology. 2016; 25 (2):527–41. https://doi.org/10.1111/mec.13481 PMID: 26685644

52. Elliott J. Wild brown trout Salmo trutta: an important national and international resource. Freshwater Biology. 1989; 21(1):1–5.

53. Lenth RV. Least-squares means: the R package lsmeans. Journal of statistical software. 2016; 69 (1):1–33.

54. Hänfling B, Lawson Handley L, Read DS, Hahn C, Li J, Nichols P, et al. Environmental DNA metabarcoding of lake fish communities reflects long-term data from established survey methods. Molecular ecology. 2016; 25(13):3101–19. https://doi.org/10.1111/mec.13660 PMID: 27095076

55. Muha TP, Rodriguez-Rey M, Rolla M, Tricarico E. Using environmental DNA to improve Species Distribution Models for freshwater invaders. Frontiers in Ecology and Evolution. 2017; 5:158.

56. Barnes MA, Turner CR. The ecology of environmental DNA and implications for conservation genetics. Conservation Genetics. 2016; 17(1):1–17.

57. Boessenkool S, Epp LS, Haile J, Bellemain E, Edwards M, Coissac E, et al. Blocking human contaminant DNA during PCR allows amplification of rare mammal species from sedimentary ancient DNA. Molecular ecology. 2012; 21(8):1806–15. https://doi.org/10.1111/j.1365-294X.2011.05306.x PMID: 21988749

58. Thomsen PF, Møller PR, Sigsgaard EE, Knudsen SW, Jørgensen OA, Willerslev E. Environmental DNA from seawater samples correlate with trawl catches of subarctic, deepwater fishes. PloS one. 2016; 11(11):e0165252. https://doi.org/10.1371/journal.pone.0165252 PMID: 27851757

59. Harper LR, Handley LL, Hahn C, Boonham N, Rees HC, Gough KC, et al. Needle in a haystack? A comparison of eDNA metabarcoding and targeted qPCR for detection of great crested newt (Triturus cristatus). bioRxiv. 2017:215897.

60. Calvignac-Spencer S, Merkel K, Kutzner N, Kühl H, Boesch C, Kappeler PM, et al. Carrion fly-derived DNA as a tool for comprehensive and cost-effective assessment of mammalian biodiversity. Molecular ecology. 2013; 22(4):915–24. https://doi.org/10.1111/mec.12183 PMID: 23298293

61. Jerde CL, Olds BD, Shogren AJ, Andruszkiewicz EA, Mahon AR, Bolster D, et al. Influence of stream bottom substrate on retention and transport of vertebrate environmental DNA. Environmental science & technology. 2016; 50(16):8770–9.

62. Goldberg CS, Strickler KM, Fremier AEK. Degradation and dispersion limit environmental DNA detection of rare amphibians in wetlands: increasing efficacy of sampling designs. Science of the Total Environment. 2018; 633:695–703. https://doi.org/10.1016/j.scitotenv.2018.02.255 PMID: 29602110

63. Bista I, Carvalho GR, Walsh K, Seymour M, Hajibabaei M, Lallias D, et al. Annual time-series analysis of aqueous eDNA reveals ecologically relevant dynamics of lake ecosystem biodiversity. Nature communications. 2017; 8:14087. https://doi.org/10.1038/ncomms14087 PMID: 28098255