Next-generation monitoring of aquatic biodiversity using environmental DNA metabarcoding

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Abstract

Global biodiversity in freshwater and the oceans is declining at high rates. Reliable tools for assessing and monitoring aquatic biodiversity, especially for rare and secretive species, are important for efficient and timely management. Recent advances in DNA sequencing have provided a new tool for species detection from DNA present in the environment. In this study, we tested whether an environmental DNA (eDNA) metabarcoding approach, using water samples, can be used for addressing significant questions in ecology and conservation. Two key aquatic vertebrate groups were targeted: amphibians and bony fish. The reliability of this method was cautiously validated in silico, in vitro and in situ. When compared with traditional surveys or historical data, eDNA metabarcoding showed a much better detection probability overall. For amphibians, the detection probability with eDNA metabarcoding was 0.97 (CI = 0.90–0.99) vs. 0.58 (CI = 0.50–0.63) for traditional surveys. For fish, in 89% of the studied sites, the number of taxa detected using the eDNA metabarcoding approach was higher or identical to the number detected using traditional methods. We argue that the proposed DNA-based approach has the potential to become the next-generation tool for ecological studies and standardized biodiversity monitoring in a wide range of aquatic ecosystems.

Keywords: amphibian, detection probability, environmental DNA, fish, monitoring, wildlife management

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Introduction

Global biodiversity loss represents one of the most serious environmental crises of the 20th and 21st centuries, with considerable impact on both ecosystem services and the health of our planet (Pimm et al. 2014). An overall biodiversity decline of 52% was recorded between 1970 and 2010, and this loss was even higher for freshwater populations than for marine or terrestrial ecosystems (WWF 2014). While scientists are struggling to find explanations and to understand the gravity of this loss (Monastersky 2014), there is an urgent need to improve effectiveness of the strategies employed to halt global biodiversity loss and render them more integrative (Beumer & Martens 2013). It is commonly acknowledged that biodiversity is threatened by a variety of anthropogenic factors (Barnosky et al. 2011); however, the most critical issue is the sheer lack of efficient and reliable tools to document the remaining species and to assess biodiversity trends. Indeed, to monitor certain taxonomic groups, the existing methods appear to be inefficient, selective, destructive or strictly dependent on a declining taxonomic expertise (Wheeler et al. 2004). Recent technological advances have provided an alternative tool for species detection using DNA present in aquatic or terrestrial environments (environmental DNA or eDNA; Taberlet et al. 2012). Taxonomical inventories and assessment of geographical distribution of species based on eDNA analysis may help to improve environmental monitoring and influence management and policy decisions (Kelly et al. 2014a; Thomsen & Willerslev 2015). In contrast to conventional survey methods, eDNA offers the advantages of being noninvasive (no macroorganisms are caught, disturbed or killed during monitoring) and reduces the risk of unintentional secondary dispersal of alien species and diseases. Overall, eDNA methods used for species detection demonstrated higher detection capability and cost-effectiveness compared to traditional methods (e.g. Darling & Mahon 2011; Deiner et al. 2012). Two main approaches using eDNA have been proposed: eDNA barcoding (or species-specific approach), which aims at detecting a single species in the environment, and eDNA metabarcoding (or multispecific approach), which simultaneously identifies several taxa from an environmental sample without ‘a priori’ knowledge of the species likely to be present in the sampled ecosystem (Taberlet et al. 2012). The latter method is based on the use of primers specific for a given taxonomic group (thereafter referred as group-specific primers) coupled with next-generation sequencing (NGS). It has been largely applied in terrestrial ecosystems and for diet analysis (e.g. Pompanon et al. 2012; Yoccoz et al. 2012). However, applications of eDNA metabarcoding in aquatic environments are still in their infancy, with only six studies published to date (Thomsen et al. 2012a,b; Kelly et al. 2014b; Deiner et al. 2015; Evans et al. 2015; Miya et al. 2015), of which only the ones from Thomsen, Deiner and Miya were performed in natural environments. These studies have emphasized the challenges posed by the method, for example sampling strategy optimization and markers reliability, but also the great perspectives yet to be investigated.

The objective of this study was to test whether an eDNA approach using water samples can be used for addressing significant questions in ecology and conservation. More specifically, our goal is to assess the potential of an integrated eDNA metabarcoding approach for ecological studies and for monitoring aquatic biodiversity, and to demonstrate its reliability considering two key aquatic vertebrate groups: amphibians and bony fish. Characterized by many discrete, rare and recently extinct species, amphibians (Batrachia) represent one of the most vulnerable animal groups (Stuart et al. 2004). Their distributions are often poorly known because the detection probabilities using traditional survey methods can be very low and may vary because of local environmental conditions (Kéry & Schmidt 2008; Tanadini & Schmidt 2011). Bony fish (Teleostei) are generally less cryptic and are often used as indicators of the ecological status of rivers (Roset et al. 2007) and lakes (Argillier et al. 2013) or of the function of water bodies (Copp et al. 1991). Nevertheless, traditional fish inventories show their limits in large water bodies (e.g. large rivers or lakes) where stratified sampling methods and invasive fishing gear (e.g. gill nets) must be used. Furthermore, those methods are usually species and/or size-selective (Hudy 1985; Hubert et al. 2012) and suffer from a relatively high proportion of fish species misidentification, particularly for young stages (Daan 2001).

The eDNA metabarcoding approach proposed here was cautiously validated in silico (i.e. bioinformatically), in vitro (i.e. using DNA extracted from tissue samples) and in situ, on a wide range of aquatic ecosystems (including both stagnant and running waters) by comparing the eDNA metabarcoding results with traditional survey data, historical data and results obtained using a previous eDNA metabarcoding approach (Thomsen et al. 2012b).

Materials and methods

Design and in silico validation of group-specific primers

Group-specific primers were designed on mtDNA for Teleostei and Batrachia using the ecoprimers software
(Riaz et al. 2011) which identifies barcode markers and their associated primers, considering several constraints, such as (i) the target taxonomic group, (ii) the minimum and maximum length of the amplicons, (iii) the maximum number of mismatches between the primers and their target sequences and (iv) the number of nucleotides with a perfect match on the 3′ end of the primers. Because DNA is susceptible to degradation in the environment, metabarcodes of <100 bp were targeted (Taberlet et al. 2012). ECOPRIMERS optimizes both the versatility of the primers for the target group and the taxonomic resolution of the amplified region. The designed primers were then tested by in silico PCR (ECOPCR program; Bellemain et al. 2010; Ficetola et al. 2010) against the entire set of DNA sequences available from the EMBL-European Nucleotide Archive (release 117, standard sequences). This step allowed the evaluation of the match between the designed primers and all potentially amplifiable sequences. The primers were then manually optimized, when necessary, by degenerating some nucleotide positions within the primers to better match the desired taxonomic group and by adding a few nucleotides on the 5′ end to homogenize the annealing temperatures of both the forward and reverse primers.

The designed primers were then extensively tested in silico using the ECOPCR program: (i) a collection of all mitochondrial DNA sequences, and (ii) all the publicly available DNA sequences. Both data sets were based on release 117 (standard sequences) of the EMBL database. First, for each primer, a sequence logo was generated (Crooks et al. 2004) based on the in silico PCR results for the target group without enforcing a perfect match at the 3′-end nucleotides and with a maximum of three mismatches in the entire primer sequence. This logo illustrates the match between the primer and its target sequence within the target taxonomic group. Second, a mismatch analysis was performed, both for the target taxonomic group and for the nontarget group (maximum of three mismatches allowed, excluding the last two nucleotides on the 3′ end) to assess the specificity of the primer pairs. Finally, the length distribution of the amplified sequences (excluding primers) was analysed using the OBITOOLS package (http://metabarcoding.org/obitools; Boyer et al. 2015).

Additionally, the Teleostei primer performance was evaluated by comparison with the alternative primer pairs recently proposed by Kelly et al. (2014b) and by Thomsen et al. (2012b).

Reference database

Fifty-three amphibian and 86 fish species (64 European freshwater fish species and 22 marine fish species present in Danish coastal waters) were collected by experienced herpetologists and ichthyologists and used to construct the reference database (Tables S1 and S2, Supporting information, respectively). DNA was extracted from 10 mg of tissue sampled from 1 to 13 individuals per species, taken from hind leg muscles of amphibians and from dorsal or caudal fins of fish, using DNeasy Blood & Tissue kit (Qiagen GmbH, Hilden, Germany), following the manufacturer’s instructions, in a room dedicated to DNA extraction from tissues. DNA amplifications were performed in a final volume of 25 µL, using 3 µL of DNA extract as template (ranging from 1 to 30 ng/µL). The amplification mixture contained 1 U of AmpliTaq Gold DNA Polymerase (Applied Biosystems, Foster City, CA, USA), 10 mM of Tris-HCl, 50 mM of KCl, 2.5 mM of MgCl₂, 0.2 mM of each dNTP, 0.2 µM of each group-specific primer designed in this study (Table 1), 4 µM of human blocking primers (i.e. a DNA oligo that preferentially binds to human DNA and that is modified to impede its amplification; Table 1) and 0.2 µg/µL of bovine serum albumin (Roche Diagnostic, Basel, Switzerland). For both taxonomic groups, the PCR mixture was denatured at 95 °C for 10 min, followed by 35 cycles of 30 s at 95 °C, 30 s at 55 °C and 1 min at 72 °C, followed by a final elongation at 72 °C for 7 min, in a room dedicated to amplified DNA, with negative air pressure and physically separated from the DNA extraction rooms.

Table 1 Group-specific mitochondrial 12S primers and human blocking primers for Batrachia and Teleostei. The code corresponds to the position of the 3′ nucleotide on the Rana nigromaculata mitochondrial genome (Accession no.: NC_002805) for the batra_F and batra_R primers and on the Cyprinus carpio mitochondrial genome (Accession no.: NC_001606) for teleo_F and teleo_R primers

<table>
<thead>
<tr>
<th>Name</th>
<th>Code</th>
<th>Sequence (5′-3′)</th>
<th>Suggested annealing temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>batra_F</td>
<td>L3541</td>
<td>ACACCGCCTGTACCCCT</td>
<td>55 °C</td>
</tr>
<tr>
<td>batra_R</td>
<td>H3596</td>
<td>GTAYACCTACATGTACGCTT</td>
<td></td>
</tr>
<tr>
<td>batra_blk</td>
<td></td>
<td>TCACCCTCTCAAGTATACCTTCACGGCA-SPC3I</td>
<td></td>
</tr>
<tr>
<td>teleo_F</td>
<td>L1848</td>
<td>ACACCGCCGTCACCTC</td>
<td>55 °C</td>
</tr>
<tr>
<td>teleo_R</td>
<td>H1913</td>
<td>CTTGGGTACATACCCATG</td>
<td></td>
</tr>
<tr>
<td>teleo_blk</td>
<td></td>
<td>ACCCTCTCAAGTATACCTTCACGGAC-SPC3I</td>
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The PCR products were purified and sequenced (forward and reverse) using Sanger technology at the Eurofins MWG Operon sequencing facilities (Ebersberg, Germany). Sequences were aligned and primers were trimmed using GENEIOUS v6.0 (Biomatters, Auckland, New Zealand, http://www.geneious.com/).

**In vitro validation of the designed primers**

To validate the universality of the designed markers for the studied taxonomic groups, we assessed whether all target species used in the reference database could be amplified with these primers. For this purpose, two pools of DNA extracted from tissue samples were prepared, one for each taxonomic group (Batrachia and freshwater Teleostei) by mixing 3 μL of DNA from each species. The group-specific primers were 5’ labelled with a unique seven-nucleotide tag (with at least three differences between tags) allowing the assignment of sequences to the respective samples during the sequence analysis. Tags for forward and reverse primers were identical for each sample. A negative control (ultrapure water) was added during the PCR step. After amplification, the two samples were titrated using capillary electrophoresis (QIAxcel; Qiagen GmbH) and purified using a MinElute PCR purification kit (Qiagen GmbH). Before sequencing, purified DNA was titrated again using capillary electrophoresis. Several purified PCR products were pooled in equal volumes, to achieve an expected sequencing depth of 100 000 reads per amphibian DNA sample and 300 000 reads per fish DNA sample. Library preparation and sequencing were performed at Fasteris facilities (Geneva, Switzerland).

Libraries were prepared using TruSeq Nano DNA genomic kit (Illumina, San Diego, CA, USA) and a pair-end sequencing (2 × 100 bp) was carried out using an Illumina MiSeq sequencer (Illumina) using the Pair-end MiSeq Reagent Kit V2 (Illumina) following the manufacturer’s instructions. In total, eight MiSeq runs were performed.

The sequence reads were analysed using the programs implemented in the OBITOOLS package (http://metabarcoding.org/obitools; Boyer et al. 2015) as described in De Barba et al. (2014). Program ILLUMINA-PAIREDEND was used to assemble forward and reverse reads corresponding to a single molecule. No special threshold was applied after the alignment step, the bad alignments being removed implicitly during the following filtration steps. Subsequently, NGSFILTER identified primers and tags and assigned the sequences to each sample. This program was used with its default parameters tolerating two mismatches for each of the two primers and no mismatch for the tags. A separate data set was created for each sample by splitting the original data set in several files using OBISPLIT. After this step, each sample was analysed individually before merging the taxon list for the final ecological analysis. Strictly identical sequences were clustered together using OBLUNIQ. Sequences shorter than 20 bp, or with occurrence lower than 10 were excluded using the OBIGREP program. The OBICLEAN program was then run to assign the status of ‘head’, ‘internal’ or ‘singleton’ to each sequence, within a PCR product. All sequences labelled ‘internal’ that correspond most likely to PCR substitutions and indel errors were discarded. The taxonomic assignment of MOTUs was performed using the program ECOTAG, with both the local reference database of Batrachia and Teleostei built for this study and the sequences extracted from the release 118 (standard sequences) of the EMBL database using the ECOPCR program (Bellemain et al. 2010; Ficetola et al. 2010). MOTUs showing <98% similarity with either the local or the EMBL reference databases were removed. Taxa were preferentially assigned based on the local reference database, except if the similarity was higher for the EMBL reference database. Finally, to take into account bad assignation of a few numbers of sequences to the wrong sample (Schnell et al. 2015), all sequences with a frequency of occurrence below 0.001 per taxon and per run for amphibians and below 0.003 per taxon and per run for fish were discarded. These threshold were empirically determined to clear the controls included in our global data production procedure (De Barba et al. 2014).

**In situ validation of the eDNA metabarcoding approach**

To assess the efficiency and reliability of this proposed eDNA metabarcoding approach, a comparative study was performed on 62 sites corresponding to a wide range of aquatic ecosystems (Tables S3 and S4, Supporting information). The eDNA sampling for all sites was conducted on the same day as the traditional method with some exceptions (see Tables S3 and S4, Supporting information). At each site, the list of species obtained using eDNA was compared to that from traditional surveys (n = 57) or historical data (n = 4). Six sites were chosen to test the presence of false positives: two small channels in a saltwater marsh along the Mediterranean Sea in 2012 because they were unsuitable for amphibian life (sites 18 and 19, Table S3, Supporting information), and four mountain lakes, two without fish (Sites 1 and 2, Table S4, Supporting information) and two where the brown trout _Salmo trutta_ was the only species present (sites 3 and 4, Table S4, Supporting information). To evaluate the performance of the metabarcoding approach proposed here (primers, sequencing device...
and analysis of NGS data) without the effect of sampling or extraction protocol, the same DNA extracts used by Thomsen et al. (2012b; sea samples) were also analysed.

Traditional field survey. For the comparative study, 39 sites were surveyed for amphibian species in 2012 (16 sites) and 2013 (23 sites), consisting of cattle ponds, ditches and natural field depressions (i.e. temporary ponds). During the day, a visual encounter survey was performed on the borders of each pond. Amphibians (larvae and adults) were also caught using a 4-mm mesh hand-net. Sampling duration varied from 10 to 30 min depending on pond area. From approximately 10:00 pm (sunset) to 12:00 pm, the ponds were visited again and a calling survey was performed. The ponds were approached quietly to a distance of approximately 20 m and, after waiting for 5 min, a 10-min auditory survey was conducted. Depending on the site typology, sometimes dip netting and visual survey with a 340-lumens torch was also performed to detect newts. All these methods were grouped as ‘traditional survey methods’ in the following analysis.

Fish surveys were performed from 2012 to 2014 by experienced fish biologists in different aquatic ecosystems (ponds, ditches, streams, lakes and rivers) in France and in the Netherlands. According to the ecosystem and the Water Framework Directive (European Council 2000), recommendations for fish monitoring, electrofishing and/or netting protocols (fyke, seine, gill) were implemented (Table S4, Supporting information).

eDNA metabarcoding survey. For small still-water ecosystems (ponds, ditches and mountain lakes), sampling was performed following the protocol described in Biggs et al. (2015). For running water (streams and rivers) or large ecosystems (lakes) where DNA is diluted in large water volumes, a new sampling strategy was tested. This strategy is based on the filtration of up to 100 L of water on site, using a filtration capsule (Environchek HV 1 μm; Pall Corporation, Ann Arbor, MI, USA), sterile tubing and a peristaltic pump (1.67 L/min) for each sample. The number of filtrations per site depended on the size of the water ecosystem (Table S4, Supporting information).

All sampling kits were prepared in a dedicated ‘DNA-free’ room. This laboratory was equipped with positive air pressure, UV treatment and frequent air renewal. Laboratory personnel wore full protective clothing (disposable coveralls, hood, mask, laboratory-specific shoes, overshoes and two pair of gloves) that was put on in an airlock foyer before entering the processing room.

For ponds, ditches and mountain lake samples, DNA extraction was performed following the method described in Tréguier et al. (2014) in a room dedicated to the analysis of ‘rare DNA’ samples, with the same requirements as the ‘DNA-free’ room. For the other sites, the filtration capsules were transported at 4 °C to the DNA extraction room and then stored at −20 °C. Later, they were rinsed with 125 mL of resuspension buffer (50 mM Tris, 10 mM EDTA) and shaken by hand for 5 min. The buffer was emptied into three 50-mL tubes, and the procedure was repeated once. In total, 250 mL was retrieved in five tubes and was centrifuged at 15 000 g for 15 min, and the supernatant was removed with a sterile pipette, leaving 15 mL of liquid at the bottom of the tube. Subsequently, 33 mL of ethanol and 1.5 mL of 3 M sodium acetate were added to each tube. After manual shaking, the tube was immediately placed at −20 °C overnight. The remaining DNA extraction steps were performed following the protocol described in Tréguier et al. (2014). Negative extraction controls were performed at each DNA extraction session (2 for amphibian and 6 for fish samples). Those controls were amplified and sequenced in the same way and in parallel to the samples to monitor possible contaminations.

DNA amplification, sequencing and analysis of NGS results were performed following the protocol described in the ‘In vitro validation of the designed primers’ section, using 12 PCR replicates per DNA sample and 50 PCR cycles instead of 35. PCR-negative controls (with 12 replicates as well) were performed and sequenced to detect potential contamination. DNA from each filter capsule was extracted, amplified and sequenced separately, but sequencing results were pooled for each site. The three DNA extracts used by Thomsen et al. (2012b) were also pooled before the amplification, and then, they were amplified and analysed as described above.

The results of the traditional surveys were compared with those from the eDNA metabarcoding approach. For this purpose, a unit was considered as an identified taxon (family, genus or species) for eDNA metabarcoding and as a species for traditional surveys.

Site occupancy analysis

Detection probabilities were calculated for each amphibian species and both methods using site occupancy models (MacKenzie et al. 2002). All statistical analyses were performed using R 3.0.2 (R Core Team 2014) and the package ‘UNMARKED’ (Fiske & Chandler 2011). Site occupancy models were performed for the 39 amphibian sites sampled in 2012–2013 (Table S3, Supporting information) with the two approaches (traditional, eDNA metabarcoding) considered as temporal replicates and
the 39 water bodies as spatial replicates. Analyses were performed for all species merged into a single data set and for each species separately to assess whether or not the relative performance of each method differed among species and for all species combined. Models that systematically included differences in occupancy probability between species (‘species’ effect on occupancy probability in the model) were fitted. On the detection probability, four different models were fitted (constant, differences between species, differences between species and methods in addition, differences between species and methods in interaction). The relative performance of these five models was compared using Akaike information criteria (Burnham & Anderson 2004).

The number of sites per ecosystem type included in the fish survey was found to be insufficient; therefore, these data were excluded from this site occupancy analysis.

Historical data. To confirm the amphibian detectability observed in this study using conventional survey methods (one visit only), historical survey data for this group were examined (55 sites). From 1983 to 2013, in the Languedoc-Roussillon region, 983 water bodies were submitted to repeated amphibian inventories (at least one visit in two different years or two visits during the same year) using traditional methods (e.g. diurnal or nocturnal visual encounters, dip netting and calling surveys, with any combination of these methods). All data are stored in a regional database ‘MALPOLON’ hosted at CEFE UMR 5175, Biogeography and Vertebrate Ecology team, in Montpellier (France) and used for the regional atlas (e.g. Geniez & Cheylan 2012). Among these 983 inventoried water bodies, 55 sites were selected near the 30 water bodies surveyed in 2012–2013 for the comparative study (Site 1–30, Table S3, Supporting information). Those 55 sites were visited at least 10 times since 1995, and at least two amphibian species were detected per site. A classical site occupancy data set was constructed in which each visit was considered to be a temporal replicate and each water body/year as a spatial replicate. The data set was analysed for all species separately and constructed only using the constant model on occupancy and detection probability. Once the detection probability was obtained, the number of visits required to detect each species with a 95% probability was calculated.

Results

Design, in silico and in vitro validation of group-specific primers

Given the defined constraints, a single primer pair was identified by eCOPRIMERS for Batrachia and Teleostei, hereafter referred as ‘batra’ and ‘teleo’ (Table 1). They both amplify a short fragment of the 12S rRNA region. For each primer pair, Fig. 1 shows (i) sequence logos illustrating the match between the different primers and their target sequences for the considered target group, (ii) statistics about the mismatches for the target and nontarget groups and (iii) the length of the amplified fragment (excluding primers). The in silico analysis demonstrates that the designed primers show a high taxonomical coverage (i.e. the proportion of species amplified in the target group; 0.9928 for Batrachia and 0.9855 for Teleostei) and high taxonomical discrimination (Table S5, Supporting information). Furthermore, the ‘teleo’ primers perform better concerning taxonomical coverage and discrimination than the alternative primer pairs recently proposed in other metabarcoding studies (Thomsen et al. 2012b; Kelly et al. 2014b). For comparing the different primer pairs, both the Bc index and the taxonomic discrimination have to be considered. For example, despite having a better taxonomic discrimination (4295 different sequences for 3811 species) when compared to the ‘teleo’ primers, the primers ‘Thomsen2’ have a low coverage (Bc) leading to the nonamplification of more than half of the target fish sequences. The comparison between the ‘Kelly’ and the ‘teleo’ primers is also interesting: the ‘teleo’ primers not only have a slightly higher coverage, but also show a better taxonomic discrimination, despite amplifying a shorter fragment (Table S5, Supporting information).

The in vitro validation demonstrated that the two primer pairs successfully amplified the DNA of all amphibian and fish species in the pooled samples, confirming their universality for the studied taxonomic group. Species-level identification was possible for all analysed amphibians except for species of the genus Pelophylax, which are well known to hybridize (e.g. PLoëtner et al. 2008). For freshwater fish species, 81% were unambiguously identified to species level, 14% to genus level (i.e. Carassius, Cottus, Leuciscus, Salvelinus) and 5% to family level (i.e. Chondrostoma nasus, Chondrostoma toxostoma and Telestes souffia). A taxon (species, genus or family) was ‘unambiguously identified’ if all the sequences associated with this taxon were not found in any other taxon. For marine fish species analysed, 68% were identified to species level and 32% to family level (i.e. Hippoglossoides platessoides, Limanda limanda, Platichthys flesus, Pleuronectes platessa, Ammodytes marinus, Ammodytes tobianus, Hyperolus lanceolatus).

In situ validation of the eDNA metabarcoding approach

Sequencing data analysis. In total, 20 211 018 reads were obtained (7 655 154 for amphibian samples and
12 369 765 for fish samples), corresponding to an average of 253 626 per sample. After filtering, 11 994 052 sequences were retrieved (3 300 203 for amphibian samples and 8 527 193 for fish samples). After the filtering, all extraction and PCR controls analysed were negative. The raw number of reads obtained per sample, the number of reads retrieved after bioinformatics filtering per sample as well as the associated run number are shown in Table S6 (Supporting information).

**Amphibians.** Amphibian species were detected in all studied water bodies using the eDNA metabarcoding approach and in 89.2% (33/37) using traditional survey methods (Table S3, Supporting information). The eDNA approach missed the detection of two species (Triturus marmoratus and Pelophylax sp.) identified using the traditional method in one and two ponds, respectively. By contrast, in 64 cases, the traditional method missed the detection of species identified using eDNA (all species and water bodies together). The detection probability for each species using site occupancy models was always 1.00 using the eDNA metabarcoding approach (Fig. 2 and Table S7, Supporting information), except for Pelophylax sp. ($P = 0.89$) and T. marmoratus ($P = 0.91$). The detection probability for the traditional method was highly variable among species ($P = 0.20–1.00$). Both methods show the same detection probability ($P = 1.00$) for Alytes obstetricans and Discoglossus pictus, two species inhabiting a single water body. For the remaining nine species, the detection probability using the eDNA metabarcoding method was always higher ($P = 0.00013$, Student’s $t$-test). When all species were considered together, the best fitted model included the sampling method covariate in addition to the species effect. This additive effect demonstrates that, considering all sites, eDNA metabarcoding improves the detection of all species. Overall, the detection probability with eDNA metabarcoding was 0.97 ($CI = 0.90–0.99$) vs. 0.58 ($CI = 0.50–0.63$) for traditional surveys. Thus, four successive visits at a water body are required using traditional methods to achieve the same detection probability obtained with a single visit using the eDNA metabarcoding approach ($1 – (1 – 0.58)^4$). The analysis of historical data using occupancy models demonstrated that the detection probabilities
for each species during a single visit varied between 0.25 and 0.65 (Table S7, Supporting information). When the species detection probability is 0.25, 11 visits are required to reach a 95% chance of detection, whereas four visits are required when the species detection probability reaches 0.65. No amphibian species were detected using the eDNA metabarcoding and traditional approaches in the ponds that were not suitable for amphibians.

**Fish.** In 89% of the studied sites, the number of fish taxa detected using the eDNA metabarcoding approach was higher (~47% of the sites) or identical (~42% of the sites) to the number species detected using traditional methods. When a taxon was not detected using the genetic approach, only a few individuals of this taxon were generally retrieved using the traditional survey. However, on many occasions, eDNA detected the presence of the taxon even when it was present at low effectives (~66% when n < 6). Sometimes, the combination of the two methods led to a higher number of taxa detected (Fig. 3 and Tables S8–S15, Supporting information). When comparing our results with the ones from (Thomsen et al. 2012b), using the same DNA samples, 24 taxa were identified vs. 15, respectively (Fig. 4 and Table S15, Supporting information).

In the fishless mountain lakes, no fish species were detected using eDNA, and only *Salmo trutta* was detected in the two other control sites. Therefore, no false-positive records were generated using the eDNA metabarcoding approach.

**Discussion**

Biodiversity assessments are at the basis of numerous ecology and conservation issues. In the present study, a novel eDNA metabarcoding approach was tested for bony fish and amphibian monitoring in a variety of water systems. The reliability and efficiency of this new method was assessed using in silico, in vitro and in situ validations, which emphasized its strengths and limitations.

**A powerful tool for aquatic species detection**

Several advantages of the eDNA metabarcoding approach proposed here make it a powerful tool for ecological studies and aquatic biodiversity monitoring. First, species detectability was shown to be superior to traditional surveys: the number of species detected per site using eDNA was identical or higher than conventional survey methods in all cases for amphibians and in 89% of the cases for fish. Second, the monitoring effort required to infer the amphibian and fish communities is lower using the eDNA approach, allowing an increased monitoring efficiency. For instance, the fish sampling effort required to reach an identical number of detected species in lakes was 3 days using 88 gill nets vs. 4 h using the eDNA metabarcoding approach.
(three water-filtration samples). For amphibian monitoring in Mediterranean ponds, four visits would be necessary to obtain similar detectability than with a single eDNA analysis. Third, this approach causes no disturbance to the ecosystem or to the target species, a top concern regarding conservation or restoration measures and for social and ethic valuation of scientific surveys of biodiversity. This noninvasive method, using sterile and disposable sampling material, also limits the risk of translocations of invasive alien species and pathogens during the field step. Fourth, it allows the detection of virtually all species of a target taxonomic group without a priori knowledge of their presence in the water body. This emphasizes the role of eDNA metabarcoding for environmental monitoring, including the early detection of alien species. Fifth, eDNA metabarcoding can also prove very advantageous in habitats where traditional methods cannot be implemented because of logistic constraints (e.g. low accessibility to the aquatic site) or have limited effectiveness (e.g. electric fishing cannot be performed in low conductivity waters; Allard et al. 2014). Sixth, as the primers designed in this study are universal for all amphibians and all bony fish species in the world, they can be used for global biodiversity assessment, allowing the large-scale temporal and spatial standardization of the method.
Importance of the choice of eDNA metabarcoding markers

The importance of marker choice in eDNA metabarcoding has recently been emphasized (Coissac et al. 2012; Deagle et al. 2014). Because there is no ideal universal metabarcode (Riaz et al. 2011), marker choice should be specific to the target taxonomic group, and validation is required before application of the metabarcoding analysis in situ (Deagle et al. 2014). Mitochondrial rRNA genes have been recommended for animal identification because they have a similar taxonomic resolution as the COI marker and they present conserved regions that flank variable regions, which allows the design of primers with high-resolution power for the target taxonomic group (Deagle et al. 2014).

The comparison of results obtained using different metabarcodes (this study) also underlines the importance of selecting an appropriate marker. For example, the primers proposed by Kelly et al. (2014b) or Thomsen et al. (2012b) show a lower taxonomic coverage and resolution compared to the ‘teleo’ primer pair (Table S5, Supporting information). However, it should be noted that the Thomsen primers were specifically designed to estimate local biodiversity in Danish coastal waters. Recently, Miya et al. (2015) proposed an alternative universal primer pair for the amplification of fish species. The amplified fragment, located in the 12S gene, is nearly twice the size of the amplified fragment using ‘teleo’ primers for only a slightly higher taxonomic resolution. We also identified a nearly identical universal primer pair using the ECOPRIMERS software during the course of this study. However, to optimize the robustness of the amplification without losing significant taxonomic resolution, we opted for the shorter ‘teleo’ marker, and not for primers similar to those published by Miya et al. (2015).

The analysis performed using the same DNA extracts from Danish coastal marine fishes as used in Thomsen et al. (2012b) detected more fish species than the previous study (24 vs. 15). These results are not simply because of the primers used but more generally because of a better performance of the metabarcoding approach, including the number of PCR replicates (12 vs. 8), the sequencing technology used (Illumina Miseq vs. Roche 454), the sequencing depth (144 851 vs. 20 315 reads) and sequence data analysis.

Limitations of the eDNA metabarcoding approach

Population structure and size. As with every biodiversity assessment technique, the eDNA metabarcoding approach proposed here presents some limitations. First, the use of this approach does not permit information of size, developmental stage and sex of the target organisms to be obtained. Second, considering that mitochondrial DNA is maternally inherited in most cases (Giles et al. 1980), it is not possible to distinguish hybrids from their maternal species when using this eDNA approach. Third, the eDNA approach does not easily provide quantitative estimates for the surveyed species, whereas density and/or biomass information are often required to comply with legal statutes (e.g. European Water Framework Directive; European Council 2000). Some studies have demonstrated a relationship between the amount of eDNA and the biomass for some species (e.g. Takahara et al. 2012; Thomsen et al. 2012a), but further studies are required to evaluate this relationship in various environments, with a range of species densities on different taxonomic groups and especially when different life stages excrete DNA to the water column simultaneously (juveniles may produce more eDNA per biomass than adults because of increased metabolism during growth; Klymus et al. 2015). Therefore, the number of sequences obtained per taxon may not be interpreted as quantitative but rather as semi-quantitative and can differentiate between abundant and rare species (e.g. Pompanon et al. 2012; Kelly et al. 2014b). It should be noted, however, that a reliable quantification is also difficult using traditional
methods depending on the studied ecosystem characteristics because of sampling and identification bias for certain species (Miranda & Schramm 2000).

Taxonomic resolution and target group detectability. Taxonomic resolution does not always reach the species level. To improve taxonomic resolution, one approach is to complement the group-specific primers with one or several additional primer pairs specifically designed to amplify more discriminant genetic regions for families with many closely related species (e.g. Cyprinidae or Gadidae).

Additionally, the primers may amplify nontarget groups (Fig. 1). To overcome this issue, the use of blocking primers for abundant nontarget species (e.g. human) is recommended and a high sequencing depth is needed to detect rare species of the target group.

Dealing with errors. One of the main challenges associated with eDNA methods is the risk of false-positive and false-negative detections (Darling & Mahon 2011). In the present study, precautions were taken in the field and in the laboratory to prevent the occurrence of such errors (e.g. Dejean et al. 2012; Biggs et al. 2015), and no false positives were encountered at the six control sites. The reliability of the proposed method was demonstrated using in silico, in vitro and in situ validation tests. Both the set-up of the laboratory, which should employ similar rigorous standards as those described for ancient DNA laboratories (Cooper & Poinar 2000), and the experience of laboratory personnel working with rare and degraded DNA are key factors that strongly influence the reliability of eDNA results.

Another challenge is to properly address errors caused by DNA degradation or that are produced during the PCR and sequencing steps that can lead to taxonomic misidentification. The choice of reliable bioinformatics tools (Coissac et al. 2012) is crucial in order to deal with this kind of errors. The use of a carefully produced local and exhaustive reference database for the target group allows avoiding a high level of sequencing errors (Harris 2003), mislabelled species (Santos & Branco 2012) and a lack of sequences for the target genetic region in public databases. Additionally, the use of a local reference database allows increasing the taxonomic resolution of the assigned eDNA sequences (e.g. Taberlet et al. 2007).

To avoid PCR errors, the amplification step could be omitted, and the entire eDNA retrieved from the sample could be sequenced directly using a shotgun approach, as proposed by Mahon et al. (2014). However, in natural aquatic ecosystems, the proportion of target group DNA (e.g. fish, amphibians) is most often minimal compared with nontarget DNA (bacteria, phytoplankton, etc.). Therefore, to detect rare species in environmental samples, several millions of reads per sample would be required. However, even if NGS becomes more affordable with an ever-increasing number of produced sequences, the cost of shotgun sequencing eDNA currently remains too high to be used routinely in biodiversity monitoring.

Conclusion

This study demonstrated the performance and reliability of the novel eDNA metabarcoding approach proposed for ecological studies and for aquatic biodiversity. It also underlined that eDNA metabarcoding and traditional approaches can be considered complementary. If the objective is to obtain a list of species present in an aquatic ecosystem, including rare or secretive species, then eDNA metabarcoding is the most efficient tool. If additional data are required (development stage, sex, etc.), traditional survey methods should be used in parallel. In the present study, the eDNA metabarcoding method was validated for fish and amphibian surveys, and similar approaches could be developed for other taxonomic groups. In a time of global biodiversity loss and substantial environmental change, we firmly believe that the proposed approach represents a next-generation tool for efficient, precise and standardized monitoring of aquatic biodiversity in various ecosystems. This approach can therefore deliver key data for addressing many fundamental and applied research questions in ecology.

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Conflicts of interest

PT and EC are co-inventors of a patent on ‘batra’ and ‘teleo’ primers and on the use of the amplified fragment for identifying amphabions and fish species from environmental samples. This patent only restricts commercial applications and has no impact on the use of this method by academic researchers. AV, EB, CG, PJ and TD are research scientists in a private company, specialized on the use of eDNA for species detection.

References


Barnosky AD, Matzke N, Tomiya S et al. (2011) From molecules to management: a statistical framework for identifying amphibians and fish species from environmental samples. This patent only restricts commercial applications and has no impact on the use of this method by academic researchers. AV, EB, CG, PJ and TD are research scientists in a private company, specialized on the use of eDNA for species detection.


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Data accessibility

Sequences for the reference databases are uploaded as online supporting information (Tables S1 and S2, Supporting information) and on Dryad doi: 10.5061/dryad.jm58p. All Illumina raw sequences data and OTU tables are available on Dryad doi:10.5061/dryad.jm58p.

Supporting information

Additional supporting information may be found in the online version of this article.

Table S1 Amphibian tissue samples used for the reference database construction.

Table S2 Fish tissue samples used for the reference database construction.

Table S3 Geographical coordinates, sampling date and results of traditional and eDNA metabarcoding surveys for amphibians.

Table S4 Location, site characteristics and sampling methods used for eDNA metabarcoding and traditional surveys for the fish comparative study.
Table S5 In silico assessment of different primer pairs targeting Batrachia and Teleostei.

Table S6 Number of reads obtained from the NGS runs per sample before and after bioinformatic filtering.

Table S7 Number of detection events of amphibian species using eDNA metabarcoding and traditional surveys; species detection probability with confidence intervals between brackets for eDNA metabarcoding, traditional surveys and historical data and number of visits required to achieve a 95% chance of species detection.

Table S8 Results from the eDNA metabarcoding survey and historical data for fish in control sites (sites 1–4).

Table S9 Results from eDNA metabarcoding and traditional surveys for fish in pond ecosystems (sites 5–8).

Table S10 Results from eDNA metabarcoding and traditional surveys for fish in ditch ecosystems (sites 9–12).

Table S11 Results from eDNA metabarcoding and traditional surveys for fish in lake ecosystems (site 13).

Table S12 Results from eDNA metabarcoding and traditional surveys for fish in stream ecosystems in the Netherlands (sites 14–17).

Table S13 Results from eDNA metabarcoding and traditional surveys for fish in stream ecosystems in France (sites 18–19).

Table S14 Results from eDNA metabarcoding and traditional surveys for fish in river ecosystems (sites 20–23).

Table S15 Results from eDNA metabarcoding for fish in the marine ecosystem (site 23).