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▶ To cite this version:

Alix Hervé, Isabelle Domaizon, Jean-Marc Baudoin, Tony Dejean, Pierre Gibert, et al.. Do temperature and population size structure influence the quantity of fish eDNA in water?. Frontiers in Environmental Science, 2023, 11, pp.1193393. 10.3389/fenvs.2023.1193393 . hal-04177222

HAL Id: hal-04177222 https://hal.inrae.fr/hal-04177222v1

Submitted on 4 Aug 2023

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RECEIVED 24 March 2023 ACCEPTED 14 July 2023 PUBLISHED 27 July 2023

CITATION

Hervé A, Domaizon I, Baudoin J-M, Dejean T, Gibert P, Valentini A and Logez M (2023), Do temperature and population size structure influence the quantity of fish eDNA in water? *Front. Environ. Sci.* 11:1193393. doi: 10.3389/fenvs.2023.1193393

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Do temperature and population size structure influence the quantity of fish eDNA in water?

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Various factors can influence the concentration of eDNA found in the environment and blur the link between eDNA results and *in situ* fish abundances, biomasses or even occurrences. We studied the influence of one abiotic factor, the temperature, and one biotic factor, the fish size class, on the amount of fish eDNA detectable in water. To do so, we conducted two controlled experiments using tench (*Tinca tinca, L*) placed into outdoor mesocosms during 1 week. The quantification of fish eDNA abundances was performed using quantitative PCR after filtrating a large volume of water (30 L). The Bayesian hierarchical ANOVAs performed on qPCR results did not detect the effect of size class on eDNA amount, but fish eDNA concentration was shown to increase significantly when temperature rose by 6°C (no effect detected at 3°C). This suggests that fish assemblage quantification of two thermally contrasting sites through eDNA is not directly comparable.

KEYWORDS

mesocosm, Tinca tinca, eDNA quantification, teleo, metabarcoding

Introduction

Environmental DNA (eDNA) analysis is increasingly used for aquatic biomonitoring, either to monitor the presence/absence of a given species or to assess the composition of fish assemblages. In parallel several methods are used to efficiently quantify species eDNA concentrations (used as a proxy of fish abundances) such as the quantitative PCR (qPCR; Doi et al., 2017; Fukaya et al., 2020; Kamoroff and Goldberg, 2018; Takahara et al., 2012) or digital droplet PCR (Doi et al., 2015a; Doi et al., 2015b; Capo et al., 2019). Recent developments have combined eDNA metabarcoding method with qPCR quantification techniques to assess the abundances of different species composing fish assemblages (van Bleijswijk et al., 2020), by matching the total eDNA quantity per sample and the relative abundance retrieved from the metabarcoding results (van Bleijswijk et al., 2020; Pont et al., 2023). Such methods offer the possibility to simultaneously quantify all species from a given community. Nonetheless, before using such an approach, it is necessary to assess how biotic and abiotic factors affect this quantification, with a simplified model.

Indeed, several factors can influence the eDNA concentration in the environment and among them fish biomass is a key biological factor (Doi et al., 2017; Itakura et al., 2020). Additionally, as eDNA released is linked to metabolism (Lacoursiere-Roussel et al., 2016; Yates et al., 2021), it could be expected that mass-specific eDNA production is higher for

10.3389/fenvs.2023.1193393

small individuals such as juveniles than for larger ones (Maruyama et al., 2014; Yates et al., 2021). However, Jo et al. (2019) and Mizumoto et al. (2018) reported contradicting results, whereby larger individuals were the greatest eDNA releasers. In any case, the size structure of populations may influence the eDNA signal (Yates et al., 2019; Yates et al., 2021) and this constitutes a serious issue when inferring biomass from eDNA quantification.

Such estimation is also challenged by the role played by abiotic factors such as temperature on eDNA concentration in the water (Harrison et al., 2019; Yates et al., 2021). Temperature plays a major role in the activity of ectotherms, which affects the amount of DNA released by fish in the environment (Strickler et al., 2015). On the one hand, the increase in temperature during spring is the main factor for exiting winter dormancy and inducing the reproductive behaviour in most species (except winter-spawners such as salmonids) from temperate regions (de Souza et al., 2016; Avlijaš et al., 2018; Thalinger et al., 2021). In this case, temperature indirectly influences the amount of eDNA by affecting the level of activity of the species. On the other hand, it has been demonstrated that an increase in temperature favours eDNA decay by spontaneously degrading DNA and by increasing DNA decomposition through the enhanced activity of microorganisms (Lance et al., 2017; Harrison et al., 2019). Studies that have investigated the role of temperature in fish eDNA commonly used large thermal gaps, greater than 5°C between consecutive treatments (Jo et al., 2019; Caza-Allard et al., 2022). Furthermore, all of these experimental studies were conducted under controlled constant temperatures without considering the natural circadian fluctuations of temperature. Therefore, the role played by temperature over restricted thermal gradients and in conditions closer to natural conditions still needs further investigation.

To test which factor may influence fish community eDNA quantification, we conducted two controlled experiments focusing on only one species, as a simplified experimental model. The first experiment compared the eDNA concentration of the species placed in three different thermal conditions but with natural circadian fluctuations of temperature, while the second experiment investigated the effect of population size structures on the eDNA quantity released in fish populations with equivalent biomasses.

Materials and methods

Tinca tinca L. was chosen as a model species because this is a native species that is widely distributed in European lakes; characterised by a calm temper, this species shows no hostility towards its congeners, which is relevant for experimental studies (Caza-Allard et al., 2022).

For the temperature experiment, we chose to test the impact of temperature on the tolerated range of this species, with low variations between treatments to submit individuals to a plausible variation of temperature in a natural environment, rather than expose them to extreme ranges in temperature as previously evaluated in several studies (Jo et al., 2019; Caza-Allard et al., 2022). Moreover, this gradient was non-stressful for this thermotolerant species. Indeed, tench can tolerate temperatures from 0°C to 38°C, with a preference range between 20°C and 26°C (Avlijaš et al., 2018). Populations of *T. tinca* were composed of six small

individuals (94.0 mm \pm 6.85 mm, 10.9 g \pm 3 g), one mediumsized individual (239.9 mm \pm 19.1 mm, 211 g \pm 51.0 g) and one large fish (316 mm \pm 33 mm, 452 g \pm 94.6 g). They were placed into 12 large outdoor mesocosms (about 775 L) for 6 days, under three thermal treatments (four replicates per treatment): ambient temperature (AT), AT plus 3°C (AT3) and AT plus 6°C (AT6). Each mesocosm was equipped with a thermal regulation system that set the temperature based on the temperature measured in a control mesocosm, which defined the AT conditions. This system maintained a constant (controlled) deviation in temperature between mesocosms (i.e., AT3, AT6) and maintained the natural circadian fluctuations within each mesocosm (between 5.5°C and 14°C for AT, 8.5°C and 17°C for AT3 and 11.5°C and 20°C for AT6) (Supplementary Figure S1).

To test the effect of fish size classes on eDNA signal, three treatments were choose along a gradient of adult size classes and thus of fish weight: small (281 mm \pm 24.5 mm, 321 g \pm 83.8 g), medium (346.7 mm ± 10.2 mm, 594.3 g ± 52.1 g) and large (396.4 mm \pm 16.1 mm, 877.5 g \pm 63.8 g). Within each mesocosm T. tinca population was composed of fish from one size class but with comparable biomass (mean = 1714 g, standard deviation = 125 g). The experiment lasted 8 days, using the same facilities (~725 L mesocosms, 649-764 L) but with similar temperatures between mesocosms (average difference of 0.125°C, Supplementary Figure S2) which varied from 6°C to 14.8°C. During both experiments, from a few days before the start of the experiment until the last day of the experiment, fish were starved because nutrition can influence DNA production (Klymus et al., 2015), and under these temperatures, fasting was not considered stressful. Even if pH and ultraviolet B radiations are known to influence eDNA degradation (Strickler et al., 2015), their influence on the results were supposed to be similar among mesocosms. Indeed, pH values were comparable among the mesocosms (maximum deviation value of 0.2 between the lowest and highest pH values), and the outdoor mesocosms are located in open landed area of the south of France (43.523°N, 5.511°E) with expected similar Sun radiations.

All macro-organisms were removed from the water before the experiments to ensure that potential fish DNA was naturally degraded in the water for at least 1 week through UV and microorganism activity (Thomsen et al., 2012). Three to four pre-treatment eDNA samples were collected before fish were placed into the mesocosms to verify the fish eDNA concentration prior fish introduction. At the end of each experiment, in each mesocosm, 30 L of water was filtered with a VigiDNA 0.45 µm filter capsule (SPYGEN, France), following the protocol described in Cantera et al. (2019). At the end of each filtration, the water inside the capsule was emptied and the capsule was filled with 80 mL of CL1 Conservation buffer (SPYGEN) and stored at room temperature. The DNA extraction and the amplification with teleo primers (Valentini et al., 2016) by quantitative PCR was performed following the protocol described in Pont et al. (2023). Two DNA extractions followed by a qPCR were performed to verify the absence of eDNA contamination. All controls were negative; the two DNA extraction controls were sequenced in Hervé et al. (2022) and only human DNA was found. To be comparable, the qPCR results were expressed in number of DNA copies per Gram of fish, after being corrected by the volume and the fish biomass of each mesocosm.



Results of the experiments on temperature (A–C) and fish size class (D–F). The qPCR results are presented as number of copies standardized to fish biomass quantified for each temperature treatment (A) and each size class treatment (D) (qPCR replicates from each mesocosm are represented with the same shape). The posterior distributions (from the Bayesian hierarchical linear model) of the average number of copies per treatment are illustrated for temperature treatment (B) and size class treatment (E). The differences between the average values of each treatment are compared two by two for temperature treatment (C) and size class treatment (F). Differences are considered significant if the 95% credible interval (between the grey horizontal bars) does not include 0.

The quantifications were not biased by the previous existence of eDNA in the mesocosms. Six of the seven pre-treatment samples were negatives and one displayed an eDNA concentration that was 444-fold lower than what was observed at the end of the experiments, and then considered negligible (Supplementary Table S1).

T. tinca were held in captivity under permit number A1300101 granted by the French authority of the "Direction départementale de la protection des populations des Bouches-du-Rhône" under articles R 214-87, R 214-122, and R 215-10 of the French law. At the end of the experiments, fish were euthanized using anaesthetic overdose (eugenol) in accordance with the European Directive 2010/63/EU.

To test the effect of each factor on eDNA amount, we used Bayesian hierarchical ANOVAs with non-informative priors, to take into account the variability between mesocosms within the same treatment. For each mean treatment parameter (μ_i), we used a normal non-informative prior for the mean value of each treatment and of each mesocosm (within-treatment) and inverse gamma priors for the precision parameters (inverse of variances) (see Supplementary Material for the codes of the two models). To compare the mean values of each treatment, we computed the deviation between mean values of each treatment two by two (e.g., small vs. medium, small vs. large, and medium vs. large) and the associated credible interval (CI, 95%) from their posterior distributions that were obtained from 3 Monte-Carlo Markov Chains. After an initial burn-in of 10,000 values, one value on 10 was conserved along the 500,000 iterations (to limit the autocorrelation between estimations). All these computations were performed using the Bayesian software JAGS and the statistical software R (v. 4.1.0).

Results

The effect of temperature on fish eDNA concentration was marked and significant (as revealed by the posterior distribution of the deviation between mean values) when comparing AT and AT6 (Figures 1A–C), with a higher number of DNA copies (on average 1.71 times greater) observed for the highest temperature. The pattern observed for AT3 showed a high variation of DNA concentrations between mesocosms (Figures 1A, B), while the variation between PCR replicates was still low (points from the same mesocosm are very close on Figures 1A, B; posterior distribution of within mesocosm variances in Supplementary Figure S3).

No effect of size classes on the amount of eDNA was detected when standardised to the fish biomass within each mesocosm (Figures 1D–F). Each CIs (95%) derived from the posterior distribution of the deviation between the average values of two size classes (e.g., medium vs. small) included zero (Figure 1F).

Discussion

It has been shown that fish eDNA concentration in the water is a balance between the activity of individuals (DNA released is thought to increase with the level of fish activity) and eDNA degradation by microorganisms and free enzymes (Harrison et al., 2019). AT6 conditions were close to the preferential range of *T. tinca* (Avlijaš et al., 2018), suggesting that in these conditions the eDNA release increased and outweighed the effect of degradation due to microbial activity. This pattern is consistent with majority of the previously observed temperature–eDNA relationship (i.e., Caza-Allard et al., 2022) but is much more marked than what was observed in previous studies, even when considering a wider thermal gradient (Jo et al., 2019). Some authors also found no relationship between temperature and eDNA amount, even in laboratory conditions, with contrasted thermal conditions (Rourke et al., 2023).

Regarding the effect of fish size classes on the amount of eDNA found in the water using universal fish primers, no clear and significant pattern was detected when standardised to the fish biomass. The amount of eDNA per Gram was comparable between larger fish and smaller ones even if they were 2.7 fold heavier. Most of the previous studies dealing with the link between fish biomass (or density) and eDNA amount have already highlighted a positive correlation between them (Rourke et al., 2022) especially in controlled conditions (Yates et al., 2019). Once standardised by fish weight, the pattern could be different with smaller fish shedding more DNA than larger ones (Maruyama et al., 2014). This was not the pattern observed here, which can be due to the different life stages used in the two studies; in our study we chose to explore the effect of a gradient of adult size classes, with less contrast in fish sizes than in the Maruyama et al. (2014) study were adults and juvenile were used. It would therefore be interesting to conduct a similar experiment considering a larger number of size classes, especially among smaller fish. The absence of significant response observed in our study could also be partly due to the quantification method that we used. The great majority of studies working on one species used specific primers to run the qPCRs, while we used universal primers designed for the amplification of teleost fishes and elasmobranchs species (Valentini et al., 2016). We chose these primers since they are the ones used in the study that tested the combination of metabarcoding and qPCR to demonstrate the possibility to quantify the abundance of species at the community level (van Bleijswijk et al., 2020; Pont et al., 2023). With the teleo primers, there is a risk to amplify DNA from organisms other than fish (Bylemans et al., 2018); however, the verifications we performed prior to fish introduction showed that such non-target DNA was at extremely low number of copies or below detection level; additionally the qPCR controls were found to be negative.

Another possibility is that the between individual variability was too high (Rourke et al., 2023) to detect an effect of size class or that the link between biomass and eDNA amount is not linear (Coulter et al., 2019; Souma et al., 2023). In their study, Coulter et al. (2019) observed that once a certain size is reached there is no more variation in the eDNA quantification (the curve reached an asymptote). Such nonlinear relationship between eDNA and fish biomass could explain the already weak link (correlation) already observed in some studies (Capo et al., 2019; Guivas and Brammell, 2020) especially different biomass could lead to quite similar eDNA concentrations. The lower link observed in situ (Yates et al., 2019) could also be due to the selectivity of the traditional methods used to assess fish biomass that serve as a comparison point. For instance, gillnets are size-selective (Prchalová et al., 2009), with larger fish with the greatest biomass poorly caught. Our results and previous observations (Maruyama et al., 2014) suggested that eDNA collected on the field is mainly due to large individuals as they represent the biggest proportion of the biomass. Even if juvenile shedding rate is greater than those of large fish (Maruyama et al., 2014), and even if they constitute the majority of the captures, they represent only a small proportion of the total biomass (tench smaller than 20 cm represent less than 2% of the biomass of the fish caught in French lacustrine biomonitoring program, OFB unpublished data). Therefore, the selectivity of traditional methods, with all the other parameters that affect eDNA in the field, could blur the potential biomass-eDNA relationship. The relationship between eDNA and fish biomass needs further investigation (Yates et al., 2021).

Finally, these results suggest the temperature effect may be an issue if biomass or abundance estimations from eDNA are compared, e.g., between two fish assemblages/communities from different lakes or in different environmental conditions. The effect of temperature may hinder comparisons because two comparable communities living in different thermal conditions would lead to different biomass estimations from eDNA quantifications. An extreme case could arise if communities with distinct biomasses would result in a comparable eDNA quantification because of the counterbalancing effect of temperature. Further investigations under contrasting environmental conditions, for a larger diversity of fish species, are needed before using eDNA as a quantifying tool in routine practice.

Data availability statement

The raw data supporting the conclusion of this article are available at 10.6084/m9.figshare.23708145.

Ethics statement

The animal study was reviewed and approved by French authority of the "Direction départementale de la protection des populations des Bouches-du- Rhône," permit number A1300101, under articles R 214-87, R 214-122, and R 215-10 of the French law.

Author contributions

AH, ML, TD, ID, and J-MB designed the experiment. AH, ML, and PG realised the experiment. AH and ML analysed the data. AH, ML, ID, TD, AV, and J-MB participated to the conceptualization. AV and TD, participated to the formal analysis. TD, J-MB, and ML participated to the project supervision. All authors contributed to the article and approved the submitted version.

Funding

This study was supported by the French Biodiversity Agency (OFB), the French National Research Institute for Agriculture, Food and Environment (INRAE), the company SPYGEN and the French National Agency for Research and Technology (ANRT).

Conflict of interest

Teleo primers and the use of the amplified fragment for identifying fish species from environmental samples are patented by the CNRS and the Universite' Grenoble Alpes. This patent only

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restricts commercial applications and has no implications for the use of this method by academic researchers. SPYGEN owns a licence for this patent. AH, TD, and AV are research scientists at a private company specialising in the use of eDNA for species detection.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fenvs.2023.1193393/ full#supplementary-material

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