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Author

Rourke, Meaghan L, Broadhurst, Matt K, Fowler, Ashley M, Hughes, Julian M, DiBattista, Joseph D, Furlan, Elise M

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EDITED BY

Géraldine Lassalle,
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l'agriculture, l'alimentation et l'environnement
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United States
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United States in collaboration with reviewer EL
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Swedish University of Agricultural
Sciences, Sweden
Steven Weiss Joseph Weiss,
University of Graz, Austria

*CORRESPONDENCE

Meaghan L. Rourke
✉ meaghan.duncan@dpi.nsw.gov.au

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Correlated environmental DNA and electrofished estimates of relative biomass for golden perch, *Macquaria ambigua*

Meaghan L. Rourke^{1,2*}, Matt K. Broadhurst^{3,4}, Ashley M. Fowler⁵,
Julian M. Hughes⁵, Joseph D. DiBattista^{6,7} and Elise M. Furlan⁸

¹Department of Primary Industries, Narrandera Fisheries Centre, Narrandera, NSW, Australia, ²Gulbali Institute for Agriculture, Water, and Environment, Charles Sturt University, Albury, NSW, Australia, ³Fisheries Conservation Technology Unit, NSW Department of Primary Industries, National Marine Science Centre, Southern Cross University, Coffs Harbour, NSW, Australia, ⁴School of the Environment, The University of Queensland, Brisbane, QLD, Australia, ⁵New South Wales Department of Primary Industries, Sydney Institute of Marine Science, Mosman, NSW, Australia, ⁶School of Environment and Science, Griffith University, Southport, QLD, Australia, ⁷Australian Museum Research Institute, Australian Museum, Sydney, NSW, Australia, ⁸Centre for Conservation Ecology and Genomics, Institute for Applied Ecology, University of Canberra, University Drive, Bruce, ACT, Australia

In response to the need to augment electrofishing surveys of golden perch, *Macquaria ambigua* (an important endemic freshwater fish in southeastern Australia), with other non-lethal survey methods, the utility of environmental DNA (eDNA) as an index of relative fish abundance/biomass was investigated. From three adjacent rivers, five sites in each were sampled for eDNA immediately before electrofishing during sequential years. Up to six individuals or 10 kg of electrofished golden perch (200–548 mm total length) were caught or observed site⁻¹. Analyses of concurrent eDNA concentrations revealed no significant relationship with the relative abundance of golden perch, but there was with relative biomass—manifesting as increasing eDNA concentrations between 1.1 and approximately 5.5 kg biomass site⁻¹, after which concentrations stabilized at greater biomasses. Future research warrants assessing the viability of sampling eDNA for spatio-temporally monitoring rivers where low biomasses of golden perch (1–5 kg site⁻¹) are likely to occur.

KEYWORDS

aquatic, biomonitoring, fishery-independent survey, freshwater, Murray–Darling Basin, Percichthyidae

Introduction

Globally, river regulation frequently impedes some teleosts from completing vital life-history movements and so it poses challenges for managing freshwater ecosystems (1, 2). Disrupted river flows fragment populations and/or increase vulnerability to genetic repercussions of diminished population size and can ultimately lead to local extinctions (3). Quantifying population sizes within rivers is therefore essential for prioritizing management strategies but requires careful consideration of methods to minimize impacts while remaining cost-effective, robust, and reliable (4).

Traditional, non-lethal fishery-independent freshwater monitoring often incorporates electrofishing, which involves administering a regulated electrical charge to the water and stunning fish before censusing and releasing them (5). Although useful, electrofishing is expensive and labor-intensive and requires highly trained staff. The method is also

relatively invasive with sub-lethal effects on some species (6). A newer, more benign survey method involves monitoring concentrations of DNA shed into the environment by aquatic organisms (termed 'eDNA') as an index of their relative abundance/biomass (7). Many recent studies have identified useful correlations between abundance/biomass and eDNA concentrations (but not always; e.g., (8, 9)). However, relationships are often stronger in controlled, rather than natural environments—owing to the influence of confounding biotic and abiotic variables among the latter and difficulties with estimating absolute population sizes using conventional methods (10). Such variability supports ongoing species-specific field-based studies to determine if eDNA concentrations may be useful for monitoring spatio-temporal changes in population sizes (11).

One large system of regulated rivers in Australia where native fish populations are censused by electrofishing is the Murray–Darling Basin (MDB). Modified river flows, along with other anthropogenic impacts, have resulted in population declines among various endemic species, including golden perch, *Maquaria ambigua*, a large-bodied (up to 23 kg and 76 cm total length (TL)) (12) recreationally important freshwater species. Golden perch have a complex life history that encompasses large spatial scales (often > 1000 km) and is closely linked to river hydrology (13, 14).

As a result, golden perch are highly susceptible to river regulation (12). Nevertheless, owing to both natural recruitment and management, including restocking, the abundance of golden perch in the MDB has increased over recent decades (15). Ongoing population monitoring is required to assess future stocks and prioritize management.

In the absence of commercial fishing or regular angler surveys, assessing golden perch stock status in the MBD has relied on annual electrofishing. There is growing interest from scientists and managers to determine if eDNA could be an alternative or complementary tool for monitoring spatio-temporal changes in populations. Considering the above, the aim of this study was to investigate the extent to which the relative abundance and/or biomass of golden perch estimated using electrofishing in three river systems over two years was correlated with eDNA concentrations in concurrent water samples. This information was then used to propose future sampling strategies.

Methods

A genetic assay was developed to amplify a fragment of the mitochondrial 12S ribosomal RNA (hereafter 12S rRNA) of golden perch (see [Supplementary material](#)). Water-sample collection for eDNA was done between 1 March and 14 May 2021 and between 28 February and 11 May 2022 and was followed by boat-based electrofishing at 15 pre-determined sites in the southern MDB. The sites were distributed throughout the Lachlan ($n = 5$), Murrumbidgee ($n = 5$), and Wakool rivers ($n = 5$) and have formed the basis of annual government electrofishing surveys since 2016 ([Figures 1A–E](#)). Each site was a 200-m-long stretch of river transect, comprising eight 50-m-long 'cells,' numbered with the first site farthest downstream ([Figure 1F](#)). Ninety seconds of electrofishing was carried

out in each cell using a 7.5 GPP Smith-Root electrofishing unit. The electrofishing units were adjusted depending on conductivity at each site with a direct current (DC) of 500 or 1,000 V, 120 Hz, a duty cycle of 10–40%, and 4.5–9.8 A applied (see [Supplementary material](#)). Key water quality parameters were recorded at each site using an Aqua TROLL 500 sonde.

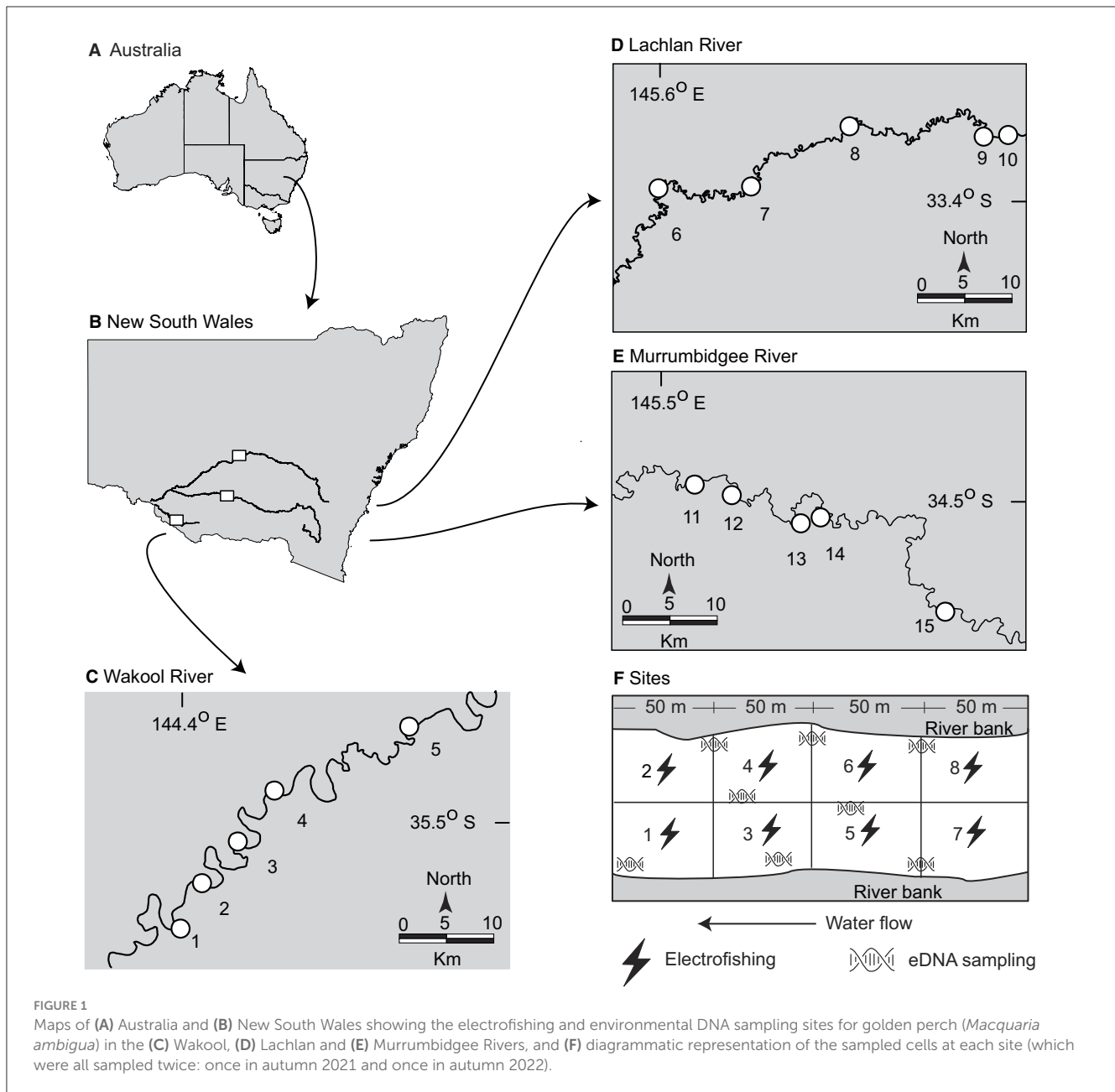
At each site, the boat was typically launched adjacent to the first cell or allowed to drift downstream to the first cell to avoid liberating DNA that may have accumulated in the sediment. Eight replicate surface-water samples (three from each bank and two moving in a transect upstream from the center of the river (i.e., evenly distributed throughout the 200-m transect; [Figure 1F](#))) were collected with a target volume of 2.0 l using a Smith Root eDNA Sampler (Smith-Root). The filter rate was set at 1.01 min^{-1} with a maximum pressure of 69 kpa using 5- μm polyethersulfone self-preserving filters (Smith-Root; (16)). An equipment control (EC) comprising 1.0 L of sterile water was filtered at each site before field sampling.

Immediately following water sample collection, electrofishing was conducted across the entire 200-m reach, positioned within each 50-m cell and beginning at cell 1 ([Figure 1F](#)). At each cell, any inert fish displaced to the surface were removed (defined as 'caught') and held in an aerated live well for up to 10 min before being measured (to the nearest 1 mm TL), weighed (to the nearest 1 g), and released well downstream of the next upstream cell. Any golden perch displaced to the surface but not retrieved were recorded as 'observed'. The weights of these latter fish were estimated as the averages caught at the site. The total caught and observed weights from cells 1–8 were used as the biomass estimate for each site. The abundance included the observed number of fish.

The DNA was extracted from water samples following conventional procedures, which are described in the [Supplementary material](#). To remove potential inhibitors, all samples were cleaned using a Zymo OneStep PCR Inhibitor Removal Kit (Integrated Sciences) following the manufacturer's instructions. The samples were evaluated for residual polymerase chain reaction (PCR) inhibition (see [Supplementary material](#)) and, if present, DNA concentrations were evaluated using 1:10 diluted DNA or (if no apparent inhibition) using neat DNA. Six replicate multiplex quantitative PCR (qPCR) reactions were performed sample⁻¹ using a standard curve generated from a dilution series of gBlockTM standards with seven concentrations between 10^{-2} and $10^{-8} \text{ ng } \mu\text{l}^{-1}$, as well as low copy number standards (10, 5, 2, and 1 copies μl^{-1} ; see the [Supplementary material](#)). The concentrations were then multiplied by 50 (for neat DNA) or 500 (for 1:10 DNA) to calculate the quantity (ng) of DNA filter⁻¹. The amount of DNA l⁻¹ of water was determined by dividing the quantity of DNA filter⁻¹ by the volume of water filtered. The concentration of DNA in copies reaction⁻¹ was calculated from the concentration in ng l⁻¹ (see [Supplementary material](#)).

Data analyses

The analyzed data comprised the concentration of golden perch DNA for each of the eight filters at each site (ng filter l⁻¹) and their



observed abundance (no. site⁻¹) and total biomass (kg site⁻¹) at the site level. Any null data for eDNA and fish abundance or biomass were removed but noted. All remaining data were log-transformed to act multiplicatively, and the relationships between eDNA concentrations (ng l⁻¹ as the response variable) and abundance and biomass (fixed effects) were initially plotted. Biomasses <1 kg were subsequently removed from analyses because these data (which were very few; see the ‘Results’ section) created an artificial change in the slope compared to those values ≥1 when log-transformed. The remaining log-transformed data were explored using linear mixed models (LMM). Random effects included ‘years,’ ‘rivers,’ and ‘sites’ with sites nested within rivers. Models were fitted within glmmTMB (v. 1.1.7) in R using penalized quasi-likelihood and with model checking (i.e., over-dispersion) and diagnostics (residual plots; (17, 18)). Because both regressions

appeared curvilinear, a quadratic term was included to improve model fit and compared against the linear model via the Akaike information criterion (AIC). Marginal and conditional r²s were estimated using the function r.squaredGLMM in the MuMIn package (v. 1.47.5) (19).

Results

The water-quality parameters varied considerably between and within rivers, with variability broadly increasing with water flow, which was always greatest in the Murrumbidgee River (Table 1). Furthermore, owing to excessive regional rainfall, the flow was greater across all rivers in 2022 than in 2021 (an average of 4,304 vs. 986 ml d⁻¹). Water sampling (0.44–1.901 of filtering) and

TABLE 1 The rivers and sites and their environmental conditions (mean \pm SE where indicated) were sampled for golden perch, *Macquaria ambigua*, eDNA prior to electrofishing.

Location	Latitude and longitude	Environmental conditions						Mean (\pm SE) eDNA (ng l^{-1})	^b Mean copies reaction ⁻¹ (\pm SE)	No. caught	No. observed	Mean (\pm SE) wt (g) caught	Mean (\pm SE) TL (mm)
		Mean (\pm SE) flow (ML d ⁻¹)	Temp (°C)	pH	DO (mg l ⁻¹)	Conductivity (mS cm ⁻¹)	Turbidity (NTU)						
<i>Wakool River</i>		258.1 (22.5)											
Site 1 ^a	35.56° S, 144.35° E	NA	19.5 (1.80)	7.5 (0.20)	7.7 (0.39)	0.4 (0.12)	55.1 (10.63)	5.86×10^{-8} (1.94×10^{-8})	0.6 (0.1)	0	0	0	NA
Site 2	35.55° S, 144.35° S	NA	19.6 (2.25)	7.0 (0.23)	8.6 (0.55)	0.4 (0.14)	41.2 (6.20)	1.95×10^{-8} (1.30×10^{-8})	0.6 (0.2)	0	0	0	NA
Site 3	35.54° E, 144.37° S	NA	18.7 (1.91)	7.2 (0.18)	7.6 (0.19)	0.4 (0.12)	46.7 (8.78)	3.05×10^{-8} (8.74×10^{-9})	0.8 (0.1)	1	1	1,923.0	485.0
Site 4	35.53° E, 144.38° S	NA	18.1 (2.05)	7.5 (0.23)	8.3 (0.64)	0.4 (0.13)	51.3 (10.38)	3.40×10^{-8} (7.66×10^{-9})	1.0 (0.2)	6	0	1,741.0 (511.4)	426.7 (63.1)
Site 5	35.51° E, 144.42° S	NA	17.7 (1.52)	7.7 (0.22)	7.9 (0.32)	0.5 (0.13)	46.8 (9.42)	4.67×10^{-8} (9.21×10^{-9})	1.6 (0.3)	1	0	2,124.0	499.0
<i>Lachlan River</i>		2,469.8 (733.5)											
Site 6	33.37° E, 145.59° S	NA	14.6 (0.14)	7.2 (0.27)	8.7 (0.03)	0.3 (0.01)	56.9 (1.81)	1.74×10^{-7} (4.92×10^{-8})	1.3 (0.4)	1	0	1,575.0	450.0
Site 7	33.37° E, 145.66° S	NA	14.3 (0.04)	8.2 (0.01)	9.3 (0.04)	0.3 (0.0002)	67.5 (1.23)	2.64×10^{-7} (3.62×10^{-8})	1.4 (0.3)	4	0	1,413.8 (166.8)	450.0 (20.4)
Site 8	33.34° E, 145.72° S	NA	17.3 (1.34)	8.1 (0.10)	8.4 (0.61)	0.3 (0.02)	92.2 (14.18)	5.63×10^{-7} (9.70×10^{-8})	3.5 (1.1)	4	2	1,589.5 (158.4)	455.3 (14.6)
Site 9	33.34° E, 145.82° S	NA	17.4 (0.05)	8.1 (0.07)	8.7 (0.11)	0.3 (0.004)	90.6 (4.11)	2.50×10^{-7} (8.21×10^{-8})	2.4 (1.1)	5	0	1,386.2 (74.5)	445.2 (9.1)
Site 10	33.34° E, 145.84° S	NA	19.1 (0.64)	8.2 (0.12)	8.7 (0.35)	0.3 (0.01)	118.1 (7.70)	3.75×10^{-7} (5.40×10^{-8})	3.4 (1.0)	2	2	855.8 (533.2)	370.0 (58.0)
<i>Murrumbidgee River</i>		5,207.5 (1037.4)											
Site 11	34.45° E, 145.54° S	NA	14.8 (0.14)	7.7 (0.16)	10.3 (0.40)	0.2 (0.000)	41.2 (0.40)	1.17×10^{-7} (3.55×10^{-8})	1.2 (0.2)	2	0	1,150.5 (134.5)	420.0 (5.0)
Site 12	34.45° E, 145.56° S	NA	19.8 (1.60)	7.5 (0.07)	8.8 (0.81)	0.1 (0.01)	79.5 (7.65)	1.31×10^{-7} (3.15×10^{-8})	0.7 (0.2)	1	0	12.0	97.0
Site 13	34.47° E, 145.61° S	NA	19.8 (1.73)	7.5 (0.06)	8.4 (0.63)	0.1 (0.01)	70.8 (5.88)	1.51×10^{-7} (5.66×10^{-8})	1.4 (0.4)	1	0	1,062.0	410.0
Site 14	34.47° E, 145.62° S	NA	16.6 (0.94)	7.0 (0.22)	9.6 (0.39)	0.1 (0.01)	57.6 (11.25)	2.04×10^{-7} (4.96×10^{-8})	1.0 (0.2)	1	2	2,588.0	504.0
Site 15	34.52° E, 145.71° E	NA	19.3 (1.65)	7.2 (0.15)	8.5 (0.44)	0.1 (0.01)	72.4 (2.97)	2.92×10^{-7} (8.32×10^{-8})	1.6 (0.4)	1	0	122.0	200.0

Flow is the mean flow of each sampling day in each river system. The mean (\pm SE) concentrations of eDNA (ng l^{-1} and copies reaction⁻¹) and the total numbers and mean (\pm SE) weights (wt) and total length (TL) of 'captured' electrocuted fish and the numbers observed (but not captured) for each site ($n = 8$ cells site⁻¹ combined across two years: 2021 and 2022). NA, not applicable; DO, dissolved oxygen; NTU, Nephelometric Turbidity Units. ^aIncludes data from 2022 only, due to evidence of contamination for the 2021 samples. ^bCopies reaction⁻¹ were calculated using successfully amplified samples only.

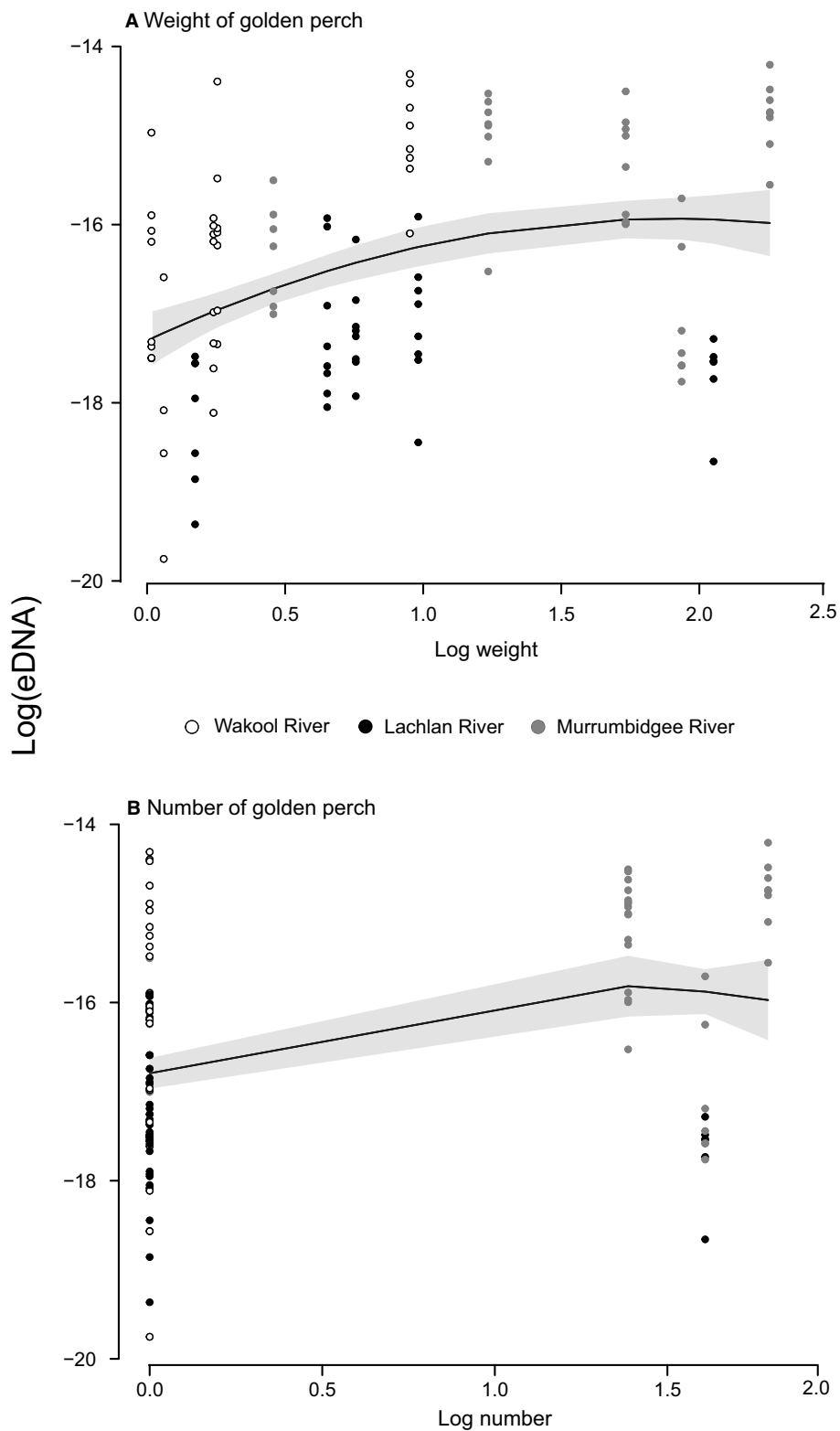


FIGURE 2
 Relationships between the log eDNA concentrations (ng l^{-1}) and log **(A)** biomass (kg) and **(B)** numbers of golden perch (*Macquaria ambigua*) at 15 sites across three river systems, during two years in the southern Murray–Darling Basin, Australia. Some weights were estimated (for fish observed but not caught), and abundances included fish caught and observed. Both axes are on the log scale.

electrofishing were successfully completed for all sites during both years, with a total of 30 golden perch (200–504 mm TL) caught and weighed (12–2588 g) and 7 observed (Table 1). At least one

golden perch was captured or observed at each site, except for sites 1 and 2 (in the Wakool River), and all but five were > 1 kg (Table 1, Figure 2).

TABLE 2 Akaike's information criterion (AIC), marginal r -squared (r^2m), and conditional r -squared (r^2c) values for models with differing random structures (years, rivers, and sites).

Model	AIC	Δ AIC	r^2m	r^2c	Δr^2c
Log (eDNA)~log(biomass) + year + river (site)	281.4	0.0	0.23	0.77	0.64
Log (eDNA)~log (biomass) + year + river	311.8	30.4	0.11	0.60	0.47
Log (eDNA)~log (biomass) + year	356.4	75.0	0.13	0.13	0.00

Sites were nested in the rivers.

Species-specific primers and a hydrolysis probe (TaqMan[®]) were successfully designed to amplify 200 base pairs (bp) of the 12S rRNA gene (see [Supplementary material](#)). A synthetic oligonucleotide (gBlock[™]) based on the golden perch 12S rRNA gene was designed with an 8-bp reverse complement sequence for use as the positive control on every qPCR plate (see [Supplementary material](#)). The 8-bp reverse complement sequence facilitated detecting of false positives due to contaminating gBlock[™]. Testing of the assay on genomic DNA from co-distributed species in the same family failed to amplify, indicating the assay was specific to golden perch (see [Supplementary material](#)). The assay efficiency was 96.7% ($R^2 = 0.99$, the limit of detection (LoD) (lowest standard with at least 95% positive detection) was 4 copies reaction⁻¹ and the limit of quantification (LoQ) (with coefficient of variation (CV) threshold set at 35%) was modeled at 5 copies reaction⁻¹ (see [Supplementary material](#)).

Of 240 samples (DNA extracts), 171 were partially inhibited following purification (71%). These samples were diluted to 1:10 for subsequent qPCR. A further four samples (all from different sites) exhibited complete inhibition and were not included in subsequent qPCRs (1.7%). Of the remaining 236 samples, golden perch DNA was detected in 220 samples (i.e., in at least 1 of 6 qPCR replicates sample⁻¹) and at every site during both years (Table 1). The eDNA data from site 1 in 2021 were discarded due to endogenous DNA and golden perch DNA in the EC, indicating contamination had occurred during sampling. All other positive and negative controls exhibited strong or null amplification, respectively, indicating no issues with the laboratory methods. There was an average of 0.6–3.5 copies reaction⁻¹ across the 15 sites (Table 1). These concentrations were below the LoQ and resulted from diluting the DNA extracts to overcome inhibition, which may affect the precision of concentration estimates. Despite this uncertainty, we are confident that the number of technical replicates increased the accuracy of the concentration estimates.

Overall, the agreement between eDNA and electrofishing, with respect to mutual detections across sites, was 42%. However, the agreement in 2021 was considerably higher (56%) than the agreement in 2022 (28%). Given that golden perch eDNA was detected at all sites in all years, mismatches were driven by zero values obtained from electrofishing, with more non-detections during 2022, particularly in the Lachlan River.

The LMMs investigating the relationships between DNA concentrations and the abundance and biomass of electrofished golden perch across the three rivers were improved when a

quadratic term was fitted (AIC abundance: linear = 293.97, quadratic = 288.03; AIC biomass: linear = 143.30, quadratic = 134.34). There was a significant relationship between eDNA and estimated biomass, which was positive between approximately 1.1 and 5.5 kg, after which there were consistent eDNA concentrations to 10.3 kg (LMM $p < 0.01$; Figure 2A). Model comparison using AIC supported all random effects in the model, which explained 77% of the variance (Table 2). The fixed effect of biomass in this model explained 23% of the variance in the eDNA concentration. A comparison of marginal and conditional r^2 s within and between models indicated that for random effects, rivers explained the greatest variance, followed by sites and then years (Table 2). The relationship between eDNA and abundance was not significant (LMM $p > 0.05$; Figure 2B).

Discussion

The significant relationship detected between the eDNA concentration of golden perch in water samples and their electrofished biomass supports a trend in the literature describing the utility of eDNA for non-lethal aquatic monitoring—although the lack of significance for abundance also reiterates the need for species-specific verification of the approach—which might be best done under controlled conditions, quantifying individual variability in DNA shedding followed by degradation (7, 20). Nevertheless, the data support using eDNA to assess golden perch biomass, albeit with some caveats that warrant consideration.

It is important to reiterate that in this study neither eDNA nor electrofishing provided estimates of absolute abundance. Rather, our focus was a comparative analysis of these two approaches for providing indices of relative quantities and was similar in theme to studies comparing traditional active (trawls and seines) or passive (hooks, traps, and gillnets) fishing gears (e.g., (21, 22)). Ultimately, such comparisons can help ascertain whether alternative methods offer statistically equivalent abundance/biomass indices and might be preferred because of not only lower operational costs but also logistical benefits. Implicit within the latter is an adequate understanding of the limitations of each method.

Both eDNA and electrofishing are affected by various biotic and abiotic factors, including river flow and temperature, which have been discussed in detail elsewhere (e.g., (7, 23, 24)). Here, river flow was not only variable among the three rivers but was also much greater across all rivers in 2022 than in 2021. Golden perch have lower electrofishing capture probabilities when daily flow increases and among individuals approximately <400 mm TL (23). Possibly, electrofishing capture probability was reduced during 2022, especially in the Murrumbidgee River, due to higher flow, deeper water, and increased turbidity following flooding and/or was affected by the relatively large proportion of small fish. Certainly, there was a lower rate of detecting golden perch by electrofishing than via eDNA in 2022. Nevertheless, water flow can also reduce eDNA concentrations and thus detectability (25), which could impact defining relationships with abundance and/or biomass.

Environmental DNA sampling and laboratory processing also have inherent limitations. For example, the number of samples site⁻¹ required to accurately estimate mean concentrations in this system was unknown *a priori*. Based on findings from other eDNA studies focused on the species-specific detection of fish in the

MDB and to maximize precision, we collected eight samples site⁻¹ (26, 27). Given golden perch DNA was noted in approximately 93% of our filters, it is likely our sampling strategy was adequate to collect sufficient DNA. Furthermore, we assessed every sample for partial inhibition following inhibitor removal and subsequently used either neat or diluted DNA to compensate for the remaining partial inhibition, providing confidence the eDNA concentration estimates were accurate.

Notably, the partial inhibition did not preclude amplification of the endogenous control and its presence would have gone undetected without a comprehensive assessment of all samples. The potential ramifications of partial inhibition on the outcomes of future studies are considerable, potentially leading to a delay in the amplification of the target DNA and uncertain quantification (28). Therefore, it remains crucial for studies assessing eDNA concentrations to address inhibition concerns to ensure the accuracy of results. Additionally, other extraction methods or inhibitor-resistant PCR reagents could be trialed (29, 30), but these are not a panacea. Nonetheless, we suggest checking for low levels of inhibition before commencing qPCR analysis, particularly in embedded riverine ecosystems.

Regardless of the specific influences of any intrinsic and extrinsic factors, collectively, these were encompassed within the observed quadratic relationship between eDNA and electrofishing. This asymptotic relationship deviates from patterns observed in many similar studies comparing eDNA concentration to biomass, where simple linear relationships have been reported, although competing model fits are often not described. Nevertheless, the quadratic relationship reported here is similar to that reported for silver carp, *Hypophthalmichthys molitrix*, in a large river, assessed using eDNA and an acoustic survey (31), and may be influenced by various factors, including, but not limited to, water temperature or flow (discussed earlier), eDNA production, or degradation (7). Deciphering the importance of potentially influencing abiotic or biotic factors warrants future investigation.

Despite the challenges posed by variability within biotic and abiotic factors and processing considerations, and although the data are few, we nevertheless identified a statistically useful relationship between eDNA concentration and biomasses ranging between 1.1 and approximately 5.5 kg per site⁻¹. This result justifies ongoing research to better describe the observed relationship and any important factors affecting variability. Such work might involve additional eDNA sampling concurrent to ongoing electrofishing assessments and would ultimately help inform appropriate future monitoring strategies for golden perch.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

The animal study was approved by the Fisheries Animal Care and Ethics Committee, Department of Regional New South Wales.

The study was conducted in accordance with the local legislation and institutional requirements.

Author contributions

MR: Writing – original draft, Writing – review & editing. MB: Writing – original draft, Writing – review & editing. AF: Writing – original draft, Writing – review & editing. JH: Writing – review & editing. JD: Writing – review & editing. EF: Writing – review & editing.

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Conflict of interest

This study received funding from the Fisheries Research and Development Corporation. The funder was not involved in the study design, collection, analyses, interpretation of data, the writing of this article or the decision to submit it for publication. All authors declare no other competing interests.

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Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/frish.2024.1358572/full#supplementary-material>

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