Environmental DNA (eDNA) detects the invasive crayfishes *Orconectes rusticus* and *Pacifastacus leniusculus* in large lakes of North America

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Abstract We report results of a study that made reciprocal comparisons of environmental DNA (eDNA) assays for two major invasive crayfishes between their disparate invasive ranges in North America. Specifically, we tested for range expansions of the signal crayfish *Pacifastacus leniusculus* (Dana, 1852) into the Laurentian Great Lakes region known to be invaded by the rusty crayfish *Orconectes rusticus* (Girard, 1852), as well as for the invasion of *O. rusticus* into large lakes of California and Nevada, US known to be invaded by *P. leniusculus*. We compared eDNA detections to historic localities for *O. rusticus* within the Great Lakes, and to recent sampling for presence/absence and relative abundance of *P. leniusculus* in California and Nevada via overnight sets of baited traps. We successfully detected *O. rusticus* eDNA at six sites from the Great Lakes and *P. leniusculus* from six of seven lakes where it was known to occur in California and Nevada, but did not detect any range expansions by either species across the North American continent. eDNA appears suitable to detect benthic arthropods from exceptionally large lakes, and will likely be useful in applications for monitoring of new biological invasions into these and other freshwater and marine habitats.

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Introduction

Biological invasions profoundly affect aquatic ecosystems and their constituent organisms globally (Gallardo et al., 2016), and consequently demand ongoing development of tools for more effective prevention and control (Lodge et al., 2016). This includes surveillance approaches that are sensitive enough to detect new invasions early when eradication or containment efforts are likely to be most successful and cost-effective (Simberloff, 2003; Vander Zanden et al., 2010). Among the most promising advances in invasive species surveillance to emerge recently has been environmental DNA (eDNA), or DNA of macrobiota collected and identified from environmental samples (Ficetola et al., 2008; Lodge et al., 2012b). Research to date has consistently found eDNA to be highly sensitive to detection of potentially harmful species at the low population abundances associated with early stages of invasion (e.g., Egan et al., 2015; Smart et al., 2015; Dougherty et al., 2016; Matsuhashi et al., 2016). However, a number of questions related to the application and interpretation of eDNA methods persist and require ongoing research attention (Roussel et al., 2015; Barnes & Turner, 2016).

Among these concerns is the performance or suitability of eDNA for applications to more diverse taxa and habitats beyond the fish or amphibians and small ponds or mesocosms where this tool was initially pioneered (e.g., Ficetola et al., 2008; Goldberg et al., 2011; Barnes et al., 2014). As examples, the performance of eDNA for benthic arthropods like freshwater crayfish has been equivocal between initial studies (e.g., Tréguier et al., 2014; Figiel & Bohn, 2015; Dougherty et al., 2016), and tests of eDNA in the largest of freshwater habitats like the Laurentian Great Lakes of the United States (US) and Canada (hereafter Great Lakes) have been relatively rare (but see Tucker et al., 2016). Given that organisms like crayfish have invaded and negatively affected large freshwater habitats including the Great Lakes (Peters et al., 2014), we ask: is it feasible to provide early detection of invasive benthic arthropods in massive freshwater lakes using eDNA? We addressed this question by developing and applying eDNA assays for two major invasive crayfishes in large lakes of their reciprocal non-native ranges in North America. Our study allowed us to both test the feasibility of eDNA for detecting benthic arthropods in situ in large freshwater ecosystems while simultaneously screening for major range expansions of two problem species into new, potentially suitable regions.

The rusty crayfish *Orconectes rusticus* (Girard, 1852) and the signal crayfish *Pacifastacus leniusculus* (Dana, 1852) are two of the most widespread and impactful invasive crayfishes on the planet (Lodge et al., 2012a); their ecological effects include dramatically reducing populations of aquatic plants, other macroinvertebrates, native crayfishes, and some imperiled vertebrate species (Twardochleb et al., 2013). *O. rusticus* is native to the Ohio River drainage in the US, but was introduced northward through pathways including live bait use by anglers into the Great Lakes, as well as inland lakes and rivers of US states and Canadian provinces like Wisconsin and Ontario (Capelli & Magnuson, 1983; Edwards et al., 2009; Peters et al., 2014). More recently, *O. rusticus* has established its first populations in western North America (e.g., Olden et al., 2009), and species distribution modeling suggests that more of the western US is likely vulnerable to further invasion by *O. rusticus* (Morehouse & Tobler, 2013). Conversely, *P. leniusculus* is native to the Columbia River and adjacent Pacific drainages of western North America, but was introduced over the past century southward to the US states of California and Nevada, and then subsequently to Europe and Japan (Lodge et al., 2012a; Larson & Williams, 2015; Usio et al., 2016). Species distribution models predict that much of eastern North America—including the Great Lakes—is likely suitable for *P. leniusculus* to establish populations, although this invasive crayfish has not been previously observed from this half of the continent (Capinha et al., 2011; Larson & Olden, 2012).

We sought to evaluate whether eDNA could detect these two invasive crayfishes from among the largest freshwater habitats of their non-native ranges, using reciprocal surveillance efforts between the Great Lakes (*O. rusticus* established, *P. leniusculus* absent but at risk for invasion) and large lakes of California and Nevada (*P. leniusculus* established, *O. rusticus* absent but at risk for invasion).
absent but at risk for invasion). We used a quantitative PCR (qPCR) eDNA assay that was recently found effective at detecting *O. rusticus* in smaller inland lakes (Dougherty et al., 2016), and developed a new qPCR eDNA assay for *P. leniusculus* using a recently published, comprehensive molecular dataset on the phylogenetics of the *Pacifastacus* genus (Larson et al., 2016). We collected water samples for eDNA from locations across all five of the Great Lakes and several associated waters, including Lake Champlain (133,100 ha) and the St. Lawrence Seaway, as well as from the two largest natural lakes in California (Lake Tahoe at 49,000 ha, Clear Lake at 18,000 ha) and a number of nearby, smaller lakes and reservoirs. We also used citizen scientists in affiliation with public zoo and aquarium summer science programs to collect some water samples from the Great Lakes region. Finally, we compared eDNA detections for *O. rusticus* to recent distributional records for this species from a comprehensive review of crayfish across the Great Lakes (Peters et al., 2014), and compared eDNA detections for *P. leniusculus* to results of baited trapping for this species in a subset of the California and Nevada lakes. Cumulatively, our multi-region study tested whether eDNA can successfully detect invasive benthic arthropods from large lake habitats while screening for new potential crayfish invasions across the North American continent.

**Methods**

Our study included (a) design and testing of primers to amplify *P. leniusculus* DNA, (b) collecting water samples from the field in both the Great Lakes and California and Nevada study regions, (c) running single-species qPCR assays for both crayfish species on samples from both study regions, and finally (d) comparing eDNA detections to either known distributions (*O. rusticus*) or previous sampling for relative abundance (*P. leniusculus*) to evaluate the performance of eDNA for these crayfish.

**Primer design and testing**

We used a primer pair designed for the *O. rusticus* cytochrome *c* oxidase subunit I (COI) mitochondrial DNA gene previously published by Dougherty et al. (2016) that was tested for specificity against tissue samples from all other known Great Lakes region sympatric crayfishes (11 species), and that was also found effective at detecting this invasive crayfish from inland lakes of Wisconsin and Michigan, US. We refer readers to Dougherty et al. (2016) for more details on primer design, testing, and previous performance, but report here that the primer pair used was Orusticus_COI_5F (5′-CAGGGGCCGTCAGTAGGTTAGGTAT-3′) and Orusticus_COI_5R (5′-CATTCCGATC TATAGTCATTCCCGTAG-3′), which produces a 128 base pair (bp) amplicon.

We designed and tested a new primer pair to amplify a subregion of the COI gene for *P. leniusculus*. We designed primers using Primer3 (Untergasser et al., 2012) and visual searches for nucleotide variants between in-group *P. leniusculus* sequences and out-group sequences of other crayfishes of the genus *Pacifastacus* and family Astacidae. The best primer pair we could identify, PacifastacusE_COI_F2 (5′-GGRGGATTTGGTAATTGGTTAATTC-3′) and PacifastacusE_COI_R2b (5′-CAATAGCCGCTGCTAGTAGAGGA-3′), produced a 184 bp amplicon. Specificity of the above primer pair was evaluated in the lab for successful amplification of tissue-derived *P. leniusculus* genomic DNA, as well as reduced amplification with tissue-derived genomic DNA for 2 other crayfish species of the genus *Pacifastacus* and family Astacidae. The best primer pair we could identify, PacifastacusE_COI_F2 (5′-GGRGGATTTGGTAATTGGTTAATTC-3′) and PacifastacusE_COI_R2b (5′-CAATAGCCGCTGCTAGTAGAGGA-3′), produced a 184 bp amplicon. Specificity of the above primer pair was evaluated in the lab for successful amplification of tissue-derived *P. leniusculus* genomic DNA, as well as reduced amplification with tissue-derived genomic DNA for 2 other crayfish species of the genus *Pacifastacus*, 4 of the European crayfish species of the family Astacidae, and 10 North American crayfish species of the family Cambaridae known to occur in the Great Lakes region, including *O. rusticus*. We evaluated performance of our primer pair using qPCR settings consistent with those reported in processing of our field samples (below). We required both a qPCR amplification curve and proper melting curve relative to that of the targeted in-group taxa to consider a sample positive, and also evaluated the timing of the quantification cycle (Cq) when fluorescence from amplification exceeds background fluorescence to further evaluate the strength of a non-target tissue-derived amplification as compared to amplification from target tissue-derived samples. Primer testing results confirmed our assay as specific enough to discriminate *P. leniusculus* DNA from non-target species, although we recommend post-qPCR Sanger sequencing confirmation. Additional details on *P. leniusculus* primer design and testing, particularly related to the complex taxonomy and phylogeny of the *Pacifastacus* genus (Larson et al., 2012, 2016), are available in Electronic Supplementary Material.
Field sample collection

We collected eDNA field samples from 14 sites within the Great Lakes region between 8 June 2015 and 22 October 2015, although a majority of samples were taken in the months of July and August (Electronic Supplementary Table S3). We collected eDNA samples from all five of the Great Lakes themselves, as well as from sites further downstream in the watershed in Lake Champlain and the St. Lawrence Seaway (Fig. 1). Specific site locations with sample dates and geographic coordinates and their abbreviations used throughout the manuscript (e.g., Fig. 1) are provided in Electronic Supplementary Table S3. These sites included a mix of shorelines of both densely populated urban centers (e.g., Chicago, Toronto) where releases of invasive crayfishes new to the Great Lakes by humans might be likely (Capinha et al., 2013), as well as more remote locations where the risk of crayfish introductions might be lower (e.g., Apostle Islands National Lakeshore of Lake Superior, Beaver Island of Lake Michigan). Further, at 9 of these 14 sites, eDNA samples were taken by citizen scientists, either lead author-supervised or unsupervised, who were participating in public zoo or aquarium summer programs, which included 24 organization staff or employees and 70 student or teacher volunteers. Participating organizations and citizen scientists are listed in Electronic Supplementary Table S3. In two cases where unsupervised citizen scientist groups took eDNA water samples, prior training on the field sampling protocol had been provided to organization staff by the authors.

We collected eDNA field samples from 11 lakes or reservoirs in California and Nevada between 26 and 31 August 2015 (Fig. 1). These sites included the two largest natural lakes in California (Lake Tahoe and Clear Lake), as well as a series of smaller lakes and reservoirs (Electronic Supplementary Table S3). We sampled from seven locations around the perimeter of Lake Tahoe and from both the southern and western

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**Fig. 1** Location of eDNA sample sites for the rusty crayfish *Orconectes rusticus* and signal crayfish *Pacifastacus leniusculus* in both the Great Lakes and California and Nevada regions of North America, with sample site abbreviations (Electronic Supplementary Table S3), whether or not *O. rusticus* or *P. leniusculus* eDNA was detected at a site, and historic localities for *O. rusticus* within the Great Lakes from Peters et al. (2014)
shoreline of Clear Lake, whereas most other study sites—regardless of region—had all replicated eDNA samples (below) taken from close proximity (within ~ 100 m) to a single location on their shoreline. Lake Tahoe in particular was chosen because of its history as a source for invasive *P. leniusculus* populations shipped to Europe (Larson & Williams, 2015), capacity to serve as a western North American surrogate for the Great Lakes in a cross-continental comparison, and finally for the availability of recent monitoring of *P. leniusculus* relative abundance throughout this lake (and some of its neighbors) for comparison to our eDNA results. Further details on recent *P. leniusculus* monitoring as related to eDNA results are addressed later in the methods.

In both the Great Lakes and California and Nevada sampling regions, we took 10 replicated eDNA water samples at each sample site, with the exception of instead taking 5 replicated eDNA water samples at the spatially dispersed sites in Lake Tahoe (Electronic Supplementary Table S3). Surface water samples of 250 ml volume were taken in bottles that had previously been decontaminated by minimum 10-min soaks in 10% bleach solution prior to a rinse in deionized water (Goldberg et al., 2016). These water samples were then immediately filtered through funnels containing 1.2 μm cellulose nitrate filters using a hand vacuum pump (Actron CP7830; Bosch Automotive Service Solutions, Warren, Michigan, US) connected to a side-arm flask. Filters were then promptly placed in 2-ml microcentrifuge tubes (USA Scientific, Ocala, Florida, US) and completely submerged in 700 ml CI (24:1, Amresco) DNA extraction and isopropanol precipitation protocol outlined in Renshaw et al. (2015): (1) 2-ml microcentrifuge tubes were incubated in a 65°C water bath for a minimum of 10 min, (2) 700 μl of CI (24:1, Amresco) was added to each tube and samples were vortexed for 5 s, (3) tubes were centrifuged at 15,000xg for 5 min and 500 μl of the aqueous layer was transferred to a fresh set of 1.5 ml microcentrifuge tubes, (4) 500 μl of ice cold isopropyl alcohol and 250 μl of 5 M NaCl were added to the 500 μl removed from the aqueous layer and tubes were precipitated at -20°C overnight, (5) the precipitate was pelleted by centrifugation at 15,000xg at room temperature for 10 min and the liquid was decanted, (6) 150 μl of room temperature 70% ethanol was added to each tube to wash pellets, (7) tubes were centrifuged at 15,000xg at room temperature for 5 min and the liquid was decanted, (8) 150 μl of room temperature 70% ethanol was added to each tube to wash pellets a second time, (9) tubes were centrifuged at 15,000xg at room temperature for 5 min and the liquid was decanted, (10) pellets were dried in a vacufuge at 45°C for 15 min, followed by air drying until no visible liquid remained, and finally, (11) pellets were rehydrated with 100 μl of 1X TE buffer, low EDTA (USB).

Three qPCR replicates were run for each eDNA extract in the following 20 μl reactions: 4.85 μl of PCR-grade water, 4 μl of 5× colorless GoTaq® flexi buffer (Promega), 0.4 μl of 10 mM dNTPs, 1.6 μl of 25 mM MgCl2, 1 μl of each 10 μM primer (forward and reverse), 0.15 μl of GoTaq® flexi DNA polymerase (Promega), 1 μl of EvaGreen (20× in water; Biotium), 2 μl of 4 μg/μl bovine serum albumin (Amresco), and 4 μl of eDNA extract. Mastercycler® ep replex (Eppendorf) cycling conditions were as follows: an initial denaturation at 95°C for 3 min; 45 cycles of denaturation at 95°C for 30 s, annealing at 60°C for 45 s, and extension at 72°C for 1 min, followed by a melting curve analysis that transitioned from 60 to 95°C over a span of 20 min. For quantification of eDNA samples, we used a 500 bp gBlock gene fragment based on GenBank accession AY701249 for *O. rusticus*, and a 487 bp gBlock gene fragment based on GenBank accessions EU921148, JF437995, and JF437997 for *P. leniusculus*. All Laboratory eDNA sample processing

All eDNA extractions followed a modified chloroform–isooamyl alcohol (hereafter “CI”) DNA extraction and isopropanol precipitation protocol outlined in Renshaw et al. (2015): (1) 2-ml microcentrifuge tubes were incubated in a 65°C water bath for a minimum of 10 min, (2) 700 μl of CI (24:1, Amresco) was added to each tube and samples were vortexed for 5 s, (3) tubes were centrifuged at 15,000xg for 5 min and 500 μl of the aqueous layer was transferred to a fresh set of 1.5 ml microcentrifuge tubes, (4) 500 μl of ice cold isopropyl alcohol and 250 μl of 5 M NaCl were added to the 500 μl removed from the aqueous layer and tubes were precipitated at -20°C overnight, (5) the precipitate was pelleted by centrifugation at 15,000xg at room temperature for 10 min and the liquid was decanted, (6) 150 μl of room temperature 70% ethanol was added to each tube to wash pellets, (7) tubes were centrifuged at 15,000xg at room temperature for 5 min and the liquid was decanted, (8) 150 μl of room temperature 70% ethanol was added to each tube to wash pellets a second time, (9) tubes were centrifuged at 15,000xg at room temperature for 5 min and the liquid was decanted, (10) pellets were dried in a vacufuge at 45°C for 15 min, followed by air drying until no visible liquid remained, and finally, (11) pellets were rehydrated with 100 μl of 1X TE buffer, low EDTA (USB).
gBlock gene fragments were synthesized by IDT. Copy numbers for gBlock fragments were estimated by multiplying Avogadro’s number by the number of moles. A serial dilution of the gBlock fragment provided a range in copy numbers for the quantification of eDNA unknowns (Gunawardana et al., 2014; Renshaw et al., 2015; Svec et al., 2015). We note as well that Great Lakes samples were run using both primers for *O. rusticus* known to occur there and for *P. leniusculus* being screened for new invasions, as well as the inverse (California and Nevada samples run twice using primers for both species).

Beyond our use of field controls to test for contamination, we also checked the eDNA extraction reagents and technique for contamination by the inclusion of a single extraction blank (one per each set of extracted eDNA samples) that involved just the reagents. On each qPCR assay, we checked the assay reagents and technique for contamination with two wells that included the same Mastermix as the rest of the plate with sterile water in place of the eDNA extract. The serial dilution of standards on each plate served as a qPCR-positive control. Finally, where amplification curves with the correct Cq and melting curves consistent with the target species (whether *O. rusticus* or *P. leniusculus*) were observed, a single qPCR replicate of these presumed positive eDNA detections was further confirmed through unidirectional Sanger sequencing with the reverse primer.

Comparison of eDNA to crayfish distributions or abundance

Unlike some previous studies including Dougherty et al. (2016) or Doi et al. (2016), we did not couple our eDNA field sampling with simultaneous conventional surveys for our target taxa. However, in both study regions, we had access to resources that could be used to relate eDNA results to either best available estimates of species distributions or recent measures of absence and relative abundance. First, Peters et al. (2014) recently summarized all known crayfish presence localities throughout the Great Lakes, using a combination of published academic and unpublished grey literature and government agency monitoring records (Fig. 1). While Peters et al. (2014) does not include reliable, widespread absence records for crayfish species such as *O. rusticus* or estimates of their population sizes or relative abundances in the Great Lakes, we still sought to relate our eDNA detections to distance to nearest observed *O. rusticus* presence locality. We did so with the expectation that some absences we observed for *O. rusticus* by eDNA might be the product of sampling sites that have not yet experienced *O. rusticus* invasion, and Peters et al. (2014) is the best available resource for this comparison. Similarly, we wanted to evaluate whether detections of *O. rusticus* eDNA tended to occur at sites either known to have been invaded by this crayfish or in close proximity to *O. rusticus* populations that may have spread to our sampling location recently. We estimated Euclidean distance in km from our eDNA sampling locations to the nearest *O. rusticus* records reported in Peters et al. (2014; Fig. 1), and tested for significant differences in distances between sites where *O. rusticus* eDNA was and was not detected. This analysis focused only on the 12 sample sites within the Great Lakes themselves and omitted 2 eDNA sample sites downstream in Lake Champlain and the St. Lawrence Seaway owing to the more limited geographic coverage of Peters et al. (2014).

In the California and Nevada region, 9 of our 11 study lakes had been recently sampled for *P. leniusculus* relative abundance as part of routine lake monitoring by the authors (Electronic Supplementary Table S4). This included Lake Tahoe and eight lakes in its immediate vicinity, but did not include the more remote Clear Lake or Camp Far West Reservoir (Fig. 1). Monitoring for invasive *P. leniusculus* in these lakes involves overnight sets of wire mesh crayfish traps (0.42 m long × 0.21 m diameter with two 60 mm diameter openings) over a depth gradient baited with dry dog food. Baited trapping results are reported as catch-per-unit effort (CPUE), the average number of crayfish collected per trap from the sampling event. Performance of baited trapping for detecting presence and reflecting relative abundance of crayfish in lakes is reviewed in Larson & Olden (2016). Monitoring of *P. leniusculus* populations within Lake Tahoe by baited trapping occurred at seven locations dispersed around the lake perimeter to which sample sites for eDNA were deliberately matched (Fig. 1). Seven of nine lakes previously sampled for crayfish by baited trapping were found to contain *P. leniusculus* of varying relative abundance, whereas two lakes have never had crayfish populations detected. Similarly, within Lake Tahoe, a gradient of
P. leniusculus relative abundance from scarce to extremely abundant was known a priori, in response to poor (sandy) versus good (rocky) benthic habitats for this crayfish (Larson & Olden, 2013). We used linear regression to relate average eDNA copy number from qPCR to recent average P. leniusculus CPUE for each of the nine lakes individually, as well as for the seven locations within Lake Tahoe. This analysis allowed us to evaluate whether eDNA results agreed with relative abundance estimated from a more conventional sampling method over gradients of P. leniusculus prevalence both between lakes, as well as within one large lake. Additional details on baited trapping data used in this analysis are reported in Electronic Supplementary Table S4.

Results

We detected O. rusticus eDNA at 6 of 14 locations in the Great Lakes, and did not detect P. leniusculus eDNA anywhere in this region (Figs. 1, 2). eDNA concentrations for O. rusticus were highest in the vicinity of Milwaukee and Chicago, where 10 out of 10 water samples contained O. rusticus eDNA, and were intermediate at Grand Traverse Bay of Lake Michigan and Toronto. Conversely, O. rusticus eDNA was most scarce in Cleveland and Detroit, where 2 and 1 of 10 water samples, respectively, were found to contain low copy numbers of O. rusticus eDNA. eDNA of O. rusticus was not detected from lower in the watershed in Lake Champlain and the St. Lawrence Seaway (Figs. 1, 2).

Pacifastacus leniusculus eDNA was detected at 6 of 11 lakes in California and Nevada, with high detections and copy numbers at most sites, excepting Prosser Reservoir where 2 of 10 water samples contained low eDNA copy numbers for this species (Figs. 1, 2). We did not detect O. rusticus eDNA from any California and Nevada lakes. We failed to detect P. leniusculus eDNA from one lake (Stampede Reservoir) known to previously support populations of this crayfish, but non-detections of P. leniusculus eDNA agreed with past sampling at two lakes believed to have no crayfish (Electronic Supplementary Table S4). We did not detect P. leniusculus eDNA from Clear Lake and Camp Far West Reservoir, which have not been sampled for crayfish by conventional (i.e., baited trapping) methods.

Finally, across qPCR runs for both species and regions, we found no evidence of contamination in either our field or laboratory controls. Across all sample runs, we observed average qPCR efficiencies of 99% (range 95–100%) and average R² values of 0.99 (range 0.92–1.00). All positive detections for O. rusticus and P. leniusculus were confirmed by Sanger sequencing and resulting match to available COI sequences for these species.

Proximity to Great Lakes O. rusticus localities

All O. rusticus eDNA detections in the Great Lakes were within 33.8 km of previously observed localities for this crayfish, whereas the majority of non-detections for O. rusticus were at sites where O. rusticus localities were more remote (52.1–100.0 km; Fig. 3). However, two localities (Duluth, Tawas Point on Lake Huron) where we did not detect O. rusticus eDNA were within 9.4 km or closer of recent historic collections of O. rusticus physical specimens (years 1999–2002). Accordingly, we did not observe any significant difference in distance to historically observed O. rusticus localities between sampling sites where we did and did not detect O. rusticus eDNA (Mann–Whitney rank sum U = 10,000, P = 0.240). We used a Mann–Whitney rank sum test for this comparison because of unequal variances between our two categories (Brown–Forsythe test P < 0.05).

eDNA copy number and P. leniusculus relative abundance

We found positive but weak relationships between eDNA copy number for P. leniusculus and relative abundance as CPUE for this crayfish estimated from recent baited trapping both in a nine lake subset of our California and Nevada study region, as well as for the seven locations within Lake Tahoe itself (Fig. 4). Again, we did not detect P. leniusculus eDNA from two lakes believed to have no crayfish populations, but also we did not detect P. leniusculus eDNA from one lake with notably high P. leniusculus CPUE values from recent sampling by baited trapping. Within Lake Tahoe, eDNA copy numbers were lowest at the one site within the lake known to have the lowest CPUE for this crayfish, but there was little relationship between eDNA copy number and crayfish relative abundance at the other six sites. In both cases, low R²
values (0.151–0.155) evidence little relationship between *P. leniusculus* CPUE and eDNA copy number.

**Discussion**

Our study demonstrated that eDNA can detect benthic arthropods such as crayfishes in exceptionally large freshwater habitats, and also tested whether two major invasive crayfishes had expanded their ranges to climatically suitable regions on the North American continent. We were relieved to not find *O. rusticus* eDNA from California or Nevada, where it has never been directly observed, and also by our reciprocal failure to find *P. leniusculus* eDNA from the Great Lakes region. Further, within the known invasive range of each crayfish, eDNA generally performed well in comparison to either historic localities for *O. rusticus* within the Great Lakes or to lakes with and without known populations for *P. leniusculus* in California and Nevada. However, applications in each range had some apparent errors of omission or false negatives where other data sources indicated that the target crayfish species was present but it was not detected with eDNA. Finally, consistent with some past studies like Dougherty et al. (2016), we found weak relationships between eDNA copy number for the crayfish *P. leniusculus* relative to more conventional estimates of relative abundance for this species.
More work is needed to determine the conditions over which eDNA can faithfully reflect patterns of relative abundance for benthic arthropods such as crayfish and other taxa.

We were encouraged by the capacity of eDNA collected from surface water samples to detect invasive *O. rusticus* in the exceptionally large Great Lakes, and suggest that this result may have relevance to eDNA applications for benthic arthropods in marine environments, as well (e.g., Kelly et al., 2016). Our six positive detections of *O. rusticus* eDNA within the Great Lakes all appear reasonable in light of their relatively close proximity to historic localities for this species reported by Peters et al. (2014). Similarly, four of our non-detections of *O. rusticus* eDNA occurred at sites where the nearest known historic locality for this crayfish was relatively remote; specifically, at the undeveloped Apostle Islands National Lakeshore, geographically isolated Beaver Island, and shorelines of the cities of Muskegon and Rochester, which lacked *O. rusticus* records nearby in lakes Michigan and Ontario, respectively (Fig. 1). However, eDNA samples from Duluth and Tawas Point did not detect *O. rusticus* eDNA, despite being close to localities for this species reported by Peters et al. (2014). In these two cases, *O. rusticus* records were from the years 1999 and 2002, and it is possible that *O. rusticus* could have experienced population declines or collapses at these sampling sites over the intervening timespan (Simberloff & Gibbons, 2004). Conversely, it is also possible that if *O. rusticus* is present but rare at these locations that our level of replication of eDNA sampling was not adequate to detect this crayfish, although Dougherty et al. (2016) found eDNA to be sensitive to detection of *O. rusticus* down to very low relative abundances in smaller inland lakes with similar replication. Future tests of the ability of eDNA to detect crayfishes in large freshwater habitats like the Great Lakes would benefit from pairing of this emerging methodology to concurrent, conventional

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**Fig. 3** Distance to historic locality for the rusty crayfish *Orconectes rusticus* in the Great Lakes from Peters et al. (2014; Fig. 1) for sites where eDNA of this species was and was not detected. Sample site abbreviations are given in Electronic Supplementary Table S3 (see also Fig. 1).

**Fig. 4** Relationships between median eDNA copies per sample + 1 (from Fig. 2) for the signal crayfish *Pacifastacus leniusculus* relative to the most recent estimates of relative abundance for this crayfish as catch-per-unit effort (CPUE) from overnight baited trapping at the same lakes (a) or locations within Lake Tahoe (b). Sample site abbreviations are given in Electronic Supplementary Table S3 (see also Fig. 1) and CPUE data are available in Electronic Supplementary Table S4.
sampling approaches like baited trapping or visual searches by divers or snorkelers (Larson & Olden, 2016).

eDNA detected _P. leniusculus_ in six of seven lakes from which it was previously known to occur in our California and Nevada study region, and did not detect this crayfish from two lakes where it has never previously been observed. However, we did not detect _P. leniusculus_ eDNA from one lake with a high previous estimate of relative abundance for this crayfish by baited trapping. Similar to our two potential errors of omission or false negatives for _O. rusticus_ in the Great Lakes, this California lake had last been sampled for crayfish by baited trapping in a preceding year (2013), and a population decline or collapse could have occurred prior to our subsequent eDNA sampling. Yet a related issue is the apparently poor agreement between eDNA copy number and estimates of crayfish relative abundance by baited trapping observed both between and within our study lakes. In this case, timing of baited trapping is seemingly an inadequate explanation for disagreement between sampling methodologies, because at least within Lake Tahoe the majority of sites had been sampled for crayfish by baited trapping only a few days prior to our 2015 water collection for eDNA. Further, this result is consistent with Dougherty et al. (2016), who found little relationship between eDNA copy number and relative abundance of crayfish estimated by baited trapping that occurred immediately following water sample collection.

At present, eDNA copy number does not appear to be a good surrogate for crayfish relative abundance as estimated by baited trapping. This could be the result of biases or limitations associated with baited trapping for crayfish, including dependency of this passive sampling method on behavior (rather than exclusively abundance) of target organisms, which can be influenced by factors including the presence or abundance of crayfish predators (Collins et al., 1983; Larson & Olden, 2016). Alternatively, it could be because these comparisons to date have primarily used surface water samples for eDNA in comparison to trapping data for a benthic organism (but see Tréguier et al., 2014), and this might have been especially relevant in a large lake like Tahoe where _P. leniusculus_ is known to use exceptionally deep water habitats (Abrahamsson & Goldman, 1970; Electronic Supplementary Table S4). Future tests of eDNA performance for crayfish or other benthic arthropods could benefit from taking water samples from deeper habitats over depth gradients to determine if the match between eDNA copy number and other estimates of relative abundance can be improved.

Our study contributes to a developing literature that increasingly suggests eDNA may be a viable monitoring tool for occupancy, if perhaps not abundance, of crayfish. Although Tréguier et al. (2014) found some disagreements between eDNA detections of the invasive red swamp crayfish _Procambarus clarkii_ (Girard, 1852) in French ponds as related to results of baited trapping for this species, Dougherty et al. (2016) instead found generally high concordance between baited trapping estimates of presence or absence and eDNA results for _O. rusticus_ from inland lakes of the northern US. Beyond applications to invasive crayfishes, Ikeda et al. (2016) tested an eDNA assay for the endangered crayfish _Cambaroides japonicus_ (De Haan, 1841) in streams of Japan, and detected eDNA for this crayfish from all sites where it was manually collected. Improvements in performance of eDNA for detecting crayfish occupancy, or better reflecting relative abundance for these species, might also be achieved from more mechanistic laboratory studies that can address habitat or environmental factors affecting eDNA persistence times or transport distances (Barnes & Turner, 2016; Goldberg et al., 2016; Shogren et al., 2016). For example, Figiel & Bohn (2015) used laboratory studies to evaluate where crayfish eDNA was most prevalent, finding most frequent detections for the species _Procambarus zonangulus_ (Hobbs & Hobbs, 1990) in sediments rather than surface waters, similar to results for some fish species (Turner et al., 2015). Ultimately, crayfish are both highly imperiled globally (Richman et al., 2015) and have also produced several major invasive species (Lodge et al., 2012a), and eDNA offers promising applications for better monitoring and management of both rare and introduced crayfishes.

eDNA is rapidly moving from proof of concept to actual implementation for purposes like surveillance for the early arrival of new or spreading invasive species (e.g., Jerde et al., 2013; Egan et al., 2015) or documenting distributional or population trends for imperiled species (e.g., de Souza et al., 2016). Here we report the first application of eDNA to test for major range expansions of invasive crayfishes into new regions where they were not previously known to
occur, but where climatic conditions are anticipated to be suitable for their population establishment and spread (Larson & Olden, 2012; Morehouse & Tobler, 2013). Although we did not detect O. rusticus from California and Nevada or P. leniusculus from the Great Lakes, our ability to detect each crayfish from its reciprocal, known invasive range supports that eDNA can likely be used in surveillance for new invasions by benthic arthropods into even large aquatic habitats like the Great Lakes. In addition, our study involved successful collaboration with citizen scientists to collect some of our eDNA water samples, making it among the first efforts to combine citizen science with eDNA sampling for monitoring of biodiversity (Biggs et al., 2015; Miralles et al., 2016). For example, unsupervised citizen scientists collected water samples that detected O. rusticus eDNA from Cleveland on Lake Erie, yet no citizen scientist water samples were affected by contamination or false positives that might be a concern for merging these two sampling methodologies. Together, our study provides an optimistic demonstration for eDNA to be applied in biodiversity monitoring for increasingly diverse taxa and habitats by increasingly diverse communities of researchers.

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