INTRODUCTION

Freshwater ecosystems rank among the most endangered habitats in the world and due to increasing human pressures conservation of these ecosystems remains a challenge (Chatterjee, 2017; Dudgeon et al., 2006; Reid et al., 2019; WWF, 2018). Among anthropogenic causes, habitat degradation, destruction or modification, unsustainable fisheries, pollution, and invasive species are persistent and significant drivers of population declines in freshwater ecosystems (Dudgeon et al., 2006; Reid et al., 2019). In North America, more than 80% of threats to fish, reptile, and amphibian populations are related to habitat degradation, exploitation, and invasive species (WWF, 2018). Reptilian and amphibian species face the highest proportion of decline among vertebrates (Böhm et al., 2013; IUCN, 2019). In Canada, wood turtle (Glyptemys insculpta) and the spiny softshell turtle (Apalone spinifera) are examples of species classified as...
threatened and endangered, respectively, by the Committee on the Status of Endangered Wildlife in Canada (COSEWIC, 2007, 2016). The major threats they face include habitat loss and fragmentation, road kills, pesticide exposure, and infectious diseases (Lesbarrères et al., 2014).

Habitat deterioration caused by pollution (i.e., toxic contaminants) organic pollution, and sediment loading, are also responsible for the important extinction rate of North American mollusks, especially for pollution-sensitive species such as freshwater mussels (Lopes-Lima et al., 2018; Ricciardi & Rasmussen, 1999). One example of a nationally imperiled mussel in Canada, the hickorynut, (Obovaria olivaria, Unionidea, COSEWIC, 2011), is currently suffering from the population decline of lake sturgeon (Acipenser fulvescens), the fish host needed to complete their life cycle. Another major cause for the hickorynut decline is the introduction of aquatic invasive species, such as the zebra mussel (Dreissena polymorpha) in the Laurentian Great Lakes and the St. Lawrence River (Hebert, Wilson, Murdoch, & Lazar, 1991; Schloesser, Metcalfe-Smith, Kovalak, Longton, & Smithee, 2006).

The introduction of invasive species, even if they are inconspicuous, can greatly modify freshwater habitats and jeopardize ecosystems integrity. For example, as a consequence of the introduction of the predatory water flea Bythotrophes longimanus in the mid-1980s, the crustacean zooplankton communities of the Laurentian Great Lakes have been drastically modified (Barbiero & Tuchman, 2004; Strecker, Arnot, Yan, & Girard, 2006). This predatory cladoceran also competes directly with larval fish for food resource (Branstrator, 1995).

Effective management of freshwater ecosystems also requires data on the distribution of exploited, rare, or invasive fish species. Expansion of invasive fish species is especially threatening for large interconnected freshwater ecosystems such as the Laurentian Great Lakes, which represent one of most important ecological natural resources as well as being of high socio-economic importance for recreational and commercial fishing industries. For example, the invasion of alewife (Alosa pseudoharengus) and sea lamprey (Petromyzon marinus) during the 1940s was linked to the decline in native fish abundance including the lake trout (Salvelinus namaycush), an important salmonid species for recreational fisheries as well as the lake whitefish Coregonus clupeaformis one of the most commercially important freshwater fishes in Canada (Madenjian et al., 2002; Wells & McClain, 1972). A salmonid stocking program was implemented to reduce alewife abundance by introducing a non-native salmonid species, that is, chinook salmon (Oncorhynchus tshawytscha), as well as creating interest for recreational fishing of this new species. More recently, the so-called “Asian carps,” including the grass carp (Ctenopharyngodon idella), bighead carp (Hypophthalmichthys nobilis), silver carp (Hypophthalmichthys molitrix), and black carp (Mylopharyngodon piceus) are being thoroughly monitored because of the threat they are representing for the socio-economic and ecological integrity of the Laurentian Great Lakes (Kolar et al., 2005).

For most freshwater species, assessment and monitoring are still mainly conducted using standard sampling methods such as gillnets for fish (Sandstrom, Rawson, & Lester, 2013; SFA, 2011), capture by traps, auditory surveys or visual observation for reptiles and amphibians (Hutchens & DePerno, 2009), and observation with an aqua-scope for mussels (OMNRF, 2018; Stoeckle, Kuehn, & Geist, 2016). However, in many cases, freshwater species may be very difficult to detect using these traditional methods due to their ecology and life-history traits as well as being a cause of habitat and population disturbance. Here, the analysis of environmental DNA (eDNA) may greatly contribute to improve the detection and monitoring of threatened, invasive, and exploited species without disturbing their habitat (Mauvisseau, Tönges, Andriantsaoo, Lyko, & Sweet, 2019; Mize et al., 2019). This approach allows tracing DNA from different sources, that is, epidermis, feces, mucus, collected in environmental samples such as water from lakes or rivers. Once filtered and DNA extracted, the presence of several or specific species is confirmed using different methods (e.g., qPCR or metagenomics), and more recently CRIPR-Cas (Williams et al., 2019) depending on the scope and goal of the study. In a metagenomics approach, all species of a targeted taxonomic community can be identified simultaneously while in qPCR or CRIPR-Cas the presence of a single targeted species is normally assessed. (Deiner et al., 2017; Rees, Maddison, Middleditch, Patmore, & Gough, 2014; Taberlet, Bonin, Zinger, & Coissac, 2018; Wilcox et al., 2013; Williams et al., 2019).

The use of qPCR for species detection relies on the critical step of developing species-specific primers that only amplify the DNA of the target species, avoiding false-positive results caused by cross-amplification by DNA from sister species. To confirm the absence of cross-amplification, primers must be tested on all related species potentially present in the region of study thus validating that only the target species is amplified by the primers (Wilcox, Carim, McElvey, Young, & Schwartz, 2015; Wilcox et al., 2013).

Over the last years, we have developed qPCR primers and probes in order to monitor invasive, threatened, or exploited aquatic species for various eDNA projects in the province of Québec, Canada. Here, we describe 60 qPCR primer pairs and associated TaqMan probes designed to detect fish (45 species), amphibians (six species), reptiles (five species), mollusks (two species), and crustaceans (two species), as well as their PCR conditions and results of their tests for cross-amplification of related species. As the geographic distribution of essentially all of these species extends throughout northeastern North America and in even more widely in some cases, these qPCR assays should be broadly useful for the detection of these species.

### 2 | MATERIALS AND METHODS

#### 2.1 | Sequence data for primer development

Reference sequences from mitochondrial genes, either cytochrome oxidase subunit 1 gene (COI), NADH dehydrogenase subunits (NADH), and cytochrome b gene (CYTB) from the targeted and related species were downloaded from BOLD (Ratnasingham & Hebert, 2007; http://www.boldsystems.org) or GenBank (Bensen...
et al., 2013; https://www.ncbi.nlm.nih.gov/genbank/) and aligned in Geneious 9.0.5 (https://www.geneious.com/). Primers were designed from the COI sequence for most species; however, NADH or CYTB sequences were chosen when the COI sequences of the targeted species did not have enough mismatches with the related species. All primers and probes were designed in regions with low intraspecific divergence while maximizing mismatches among related species at the extreme 3’end (Wilcox et al., 2013). Sequences were downloaded for 45 targeted fish species from 17 families, for five reptile species from three families, for six amphibian species from two families, for two crustaceans and two mollusks as well as sequences of related species present in Québec (Table S1).

For the alewife floater (mollusk, Utterbackiana implicata) and related species, some sequences for the gene of interest were unavailable in the database. Thus, the NADH I sequence was generated by PCR amplification on extracted genomic DNA using primers developed by Serb, Buhay, and Lydeard (2003), Leu-uurF (5’-TGGCAGAAAAGTGCATCAGATTTAACGC-3’) paired with NJU-12073 (5’-TGGGAATCTCTCTGCGAAAGTC-3’) or LoGlyR (5’-CTGCTGGAGGCAAGTGTACT-3’) following these conditions: 34 cycles × [94–98°C, 40 s], 50–58°C for 1 min and 68–72°C (5′-CCTGCTTGGAAGGCAAGTGTACT-3′) following these conditions: 50°C for 2 min, 95°C for 10 min 50 cycles × [95°C for 1 min, 94°C for 20 s, 60°C for 30 s]. Finally, selected primers were tested with their probes in a TaqMan assay in a final volume of 20 µl including 1.8 µl of each primer (10 µM), 0.5 µl of probe (10 µM), 10 µl of TaqMan® Environmental Master Mix 2.0 (Life Technologies), 3.9 µl of dH₂O and 2 µl of DNA (10 ng) following these conditions: 50°C for 2 min, 95°C for 10 min 50 cycles × [95°C for 15 s, 60°C for 1 min].

2.2 | Primer development

Primers were designed to amplify fragments in a range of 101–250 bp to allow for Sanger sequencing in order to be able to validate eDNA detection when necessary. Annealing temperature was validated using Primer Express 3.0 (Life Technologies) and cross-amplification to unrelated species was verified using Primer Blast (Ye et al., 2012; https://www.ncbi.nlm.nih.gov/tools/primer-blast/).

All designed primers and probes were validated for amplification of targeted species and for cross-amplification with related species (Table S1) using in-house extracted genomic DNA from various tissues for fish, amphibians, mollusks, and crustaceans using a salt DNA extraction protocol (Aljanabi & Martinez, 1997), and from blood for reptiles using the DNeasy blood and tissue kit (Qiagen). Preliminary primer screening was performed with FAST SYBR Green (Life Technologies). Amplifications were performed on a 7,500 Fast Real-Time PCR System (Applied Biosystems) in a final volume of 20 µl: 10 µl of Fast SYBR® Green Master Mix, 1 µl of each primer (10 µM), 2 µl of DNA (5–10 ng) and 6 µl of UltraPure Distilled Water (DNase, RNase, Free, Invtrogen™) following these conditions: 95°C for 20 s, 40 cycles × [95°C for 3 s, 60°C for 30 s]. Finally, selected primers were tested with their probes in a TaqMan assay in a final volume of 20 µl including 1.8 µl of each primer (10 µM), 0.5 µl of probe (10 µM), 10 µl of TaqMan® Environmental Master Mix 2.0 (Life Technologies), 3.9 µl of dH₂O and 2 µl of DNA (10 ng) following these conditions: 50°C for 2 min, 95°C for 10 min 50 cycles × [95°C for 15 s, 60°C for 1 min].

2.3 | Assay sensitivity

A standard curve experiment was performed following the same conditions as described above for the TaqMan assay. A synthetic DNA template of 500 base pairs (Integrated DNA Technologies Inc.) including the target amplicon sequence was designed from the COI, CYTB, or NADH gene sequence depending on the species. From the stock, diluted at 1.00E + 10 copies/µl, a nine-level dilution series (2,000, 1,000, 500, 100, 20, 8, 4, 2, and 1 copies per reaction) was prepared in a sterile yeast tRNA (10 µg/µl) solution. Ten replicates of each dilution were run to determine, for each primer/probe set, the amplification efficiency and the limit of detection defined as the lowest copies per reaction with >95% amplification success (Bustin et al., 2009).

3 | RESULTS

A final set of 60 assays were optimized and validated, one per targeted species which are presented in Tables 1–4. Species used for cross-amplification tests are presented in Table S1 and mismatches to primers with respect to related species are available on DRYAD. Only five tests showed a cross-amplification with the DNA of related species (primer set for S. namaycush, A. rostrata, E. lucius, M. thompsonii, and D. fuscus), thus confirming assay specificity for practically all primer-probe sets. In addition, 18 assays were tested for efficiency and limits of detection using a standard curve experiment with synthetic DNA, which revealed high amplification efficiency (Table 5). Most assays were developed for detection experiments, not for quantification, therefore no standard curve experiment with synthetic DNA was performed.

3.1 | Exploited fish and monitored fish species

Species-specific primers were designed for 22 key species for recreational fisheries and 20 of these were validated in eDNA studies (Table 1). Two species-specific assays were designed for monitored fish species, brown bullhead (Ameiurus nebulosus) and eastern silvery minnow (Hybognathus regius). Standard curve experiments were performed for 14 of these species, including six salmonids (Salmo salar, S. trutta, Coregonus clupeaformis, Prosopium cyundraeum, Oncorhynchus mykiss, Salvelinus alpinus), largemouth bass (Micropterus dolomieu), striped bass (Morone saxatilis), lake sturgeon (A. fulvescens), sand lance (Ammodytes sp.), Atlantic herring (Clupea harengus), capelin (Mallotus villosus), rainbow smelt (Osmerus mordax), and the redfish (Sebastes sp.). Based on the standard curve experiment, the assays for the salmonid species had an amplification efficiency varying between 93.5% and 108.9%, as expected for an efficiency considered as acceptable (Taylor et al., 2019) and a limit of detection varying between 20 mtDNA copies/rxn (S. trutta, M. dolomieu, O. mordax) and two mtDNA copies/rxn (S. salar) (Table 5).
| Scientific and Common name | Primer/Probe Gene | Sequence 5′ > 3′ bp eDNA | bp | eDNA |
|----------------------------|-----------------|--------------------------|----|------|
| Acipenser fulvescens       | ACFU_COIF       | GCTGGCGGGAAACCTG         | 179| v    |
| Lake sturgeon              | ACFU_COIR       | TGACTAATACAGATCACAAACAGAGT |    |
|                            | ACFU_COI Probe  | TACCATTATTAACATGAAACCC   |    |
| Ameiurus nebulosus (*)      | AMNE_CYTBF      | CCCCCCTGCTAGTTAATCTTCTGAGGA | 133| -    |
| Brown bullhead             | AMNE_CYTBR      | GGTTCATGAATAAAGGGCATGTAAG |    |
|                            | AMNE_CYTBR Probe| ACCCGATTCTCAGATTTT       |    |
| Ammodictidae sp             | AMSP_COIF       | GGTAGTTTTAACAATCTTCTCTGAGGA | 143| v    |
| Sandlance                   | AMSP_COIR       | ATTACCAAGCTCACAAAATACAGA |    |
|                            | AMSP_COI Probe  | AACTTCATCACACACATTA      |    |
| Clupea harengus             | CLHA_COIF       | ACATGGATATCCTCCTGGGATTG  | 193| v    |
| Atlantic herring            | CLHA_COIR       | GGTTCATGAATAAAGGGCATGTAAG |    |
|                            | CLHA_COI Probe  | AACTTCATCACACACATTA      |    |
| Coregonus clupeaformis      | COCL_CYTBF      | AAAACCTCTTCTCTGGGTG      | 198| v    |
| Lake whitefish              | COCL_CYTBR      | AGTGTGACCTCCTGGGATTG     |    |
|                            | COCL_CYTBR Probe| TTTGTCAGTAATCTGGA        |    |
| Cyprinus carpio             | CYCA_COIF       | CCAACTAATAATCAGGAGCTCAGGA | 173| v    |
| Common carp                 | CYCA_COIR       | GGTTCATGAATAAAGGGCATGTAAG |    |
|                            | CYCA_COI Probe  | AACTTCATCACACACATTA      |    |
| Esox lucius                 | ESLU_COIF       | CCATTATTGTTTGAAGCAGTCCTG | 152| v    |
| Northern pike               | ESLU_COIR       | GGTTCATGAATAAAGGGCATGTAAG |    |
|                            | ESLU_COI Probe  | AACTTCATCACACACATTA      |    |
| Esox masquinongy            | ESMA_COIF       | AGGGTTGGAAATCTGACTAATTTCTG | 189| v    |
| Muskellunge                 | ESMA_COIR       | GGTTCATGAATAAAGGGCATGTAAG |    |
|                            | ESMA_COI Probe  | TTTGTCAGTAATCTGGA        |    |
| Hybognathus regius (*)      | HYRE_COIF       | GCACTAGTGACCATATCTTCTCC | 204| -    |
| Eastern silvery minnow      | HYRE_COIR       | CATAGTGATCCGCGGAGCTAAA  |    |
|                            | HYRE_COI Probe  | CTGTGTCGCTGCTCTAT       |    |
| Mallotus villosus           | MAVI_COIF       | GCAATCTCGCTCAGCGG        | 185| v    |
| Capelin                     | MAVI_COIR       | AAGAAGAAACCAGTGTAATTAGCACA |    |
|                            | MAVI_COI Probe  | AACTTCATCACACACATTA      |    |
| Microgadus tomcod           | MITO_COIF       | CTTCTGACTTTTACCCCGTCA   | 166| -    |
| Atlantic tomcod             | MITO_COIR       | TGAAATTCCTCGCCAGATGAGGC |    |
|                            | MITO_COI Probe  | CCGGAGCGCTCGTCTGGA      |    |
| Micropterus dolomieu        | MIDO_COIF       | ACCATCTTCTCCTCATCTTGCG  | 173| v    |
| Smallmouth bass             | MIDO_COIR       | GCGAGGACTGGGAGGCATGATA   |    |
|                            | MIDO_COI Probe  | CCGGAGCGCTCGTCTGGA      |    |
| Morone saxatilis            | MOSA_COIF       | TGAAACTGGCTGAACGTTTAC    | 178| v    |
| Striped bass                | MOSA_COIR       | GTGTGATATTTGGGAGATGAGCA |    |
|                            | MOSA_COI Probe  | CATCTGAGACCTAACAATTT    |    |
| Moxostoma valenciennesi     | MOVA_CYTBF      | CTTGAGGATTATATATATGGATCCTACCTTAC | 251| -    |
| Greater redhorse            | MOVA_CYTBR      | GTGAAAGGCGGAAGAATCGTGGT |    |
|                            | MOVA_CYTBR Probe| CGCAGTACCTTATGTTG       |    |
Cross-amplification tests revealed co-amplification of *S. namaycush* primers with *S. alpinus*; however, these two species are rarely found in sympatry in North America. Testing for cross-amplification also revealed that *Esox lucius* primers amplified *E. americanus americanus*, with the Canadian distribution range of this latter species being limited in Québec, and hybridization being common throughout this genus (Crossman & Buss, 1965).

### 3.2 Threatened or invasive fish species

Specific primers were designed for 15 fish species listed as endangered, threatened, special concern, or susceptible to be special concern by the Species At Risk Act in Canada, by the Committee on the Status of Endangered Wildlife in Canada (COSEWIC) or by the Québec’s “Loi sur les espèces menacées ou vulnérables” (Table 2); eastern sand darter (*Ammodramys pellucida*), channel darter (*Percina copelandi*), copper redhorse (*Moxostoma hubbsi*), river redhorse (*Moxostoma carinatum*), American shad (*Alosa sapidissima*), Atlantic sturgeon (*Acipenser oxyrinchus*), American eel (*Anguilla rostrata*), brassy minnow (*Hybognathus hankinsoni*), chestnut lamprey (*Ichthyomyzon castaneus*), deepwater sculpin (*Myoxocephalus thompsonii*), grass pickerel (*Esox americanus vermiculatus*), marbled madtom (*Noturus insignis*), northern sunfish (*Lepomis punctatus*), yellow bullhead (*Ameiurus natalis*), rosyface shiner (*Notropis rubellus*); and for six invasive fish species, grass, silver and bighead carps (*Ctenopharyngodon idella, Hypophthalmichthys molitrix, Hypophthalmichthys nobilis*), goldfish (*Carassius auratus*), tench (*Tinca tinca*).
| Scientific and common name | Primer/Probe gene | Sequence 5′ > 3′ bp | eDNA |
|--------------------------|-----------------|----------------------|------|
| **(a) Invasive**          |                 |                      |      |
| Carassius auratus        | CAAU_COIF       | GGTGTGTCAGACACCTGCTAA | 165  |
| Goldfish                 | CAAU_COIR       | TTCTTCCTCATATCTCTGT  |      |
|                          | CAAU_COI_probe  | CTTTGGATGARACACCTGCTA |      | v |
| Ctenopharyngodon idella  | CTID_COIF       | TCAACACCGAGAGGCTAATAGTGG | 127  | v |
| Grass carp               | CTID_COIR       | GGTGGAAATGACTGATCCAT |      | v |
| Hypophthalmichthys       | HYMO_COIF       | TAGCAGGTGTGATCAATTGGA | 160  | v |
| molitrix                  | HYMO_COIR       | CACACGGCTAAACTGGAAGAT |      | v |
| Silver carp              | HYMO_COI probe  | CATCCGGTGCCAGCT      |      | v |
| Hypophthalmichthys       | HYNO_COIF F2    | TCAACACCAGAAGGCTAATAGTGG | 127  | v |
| nobilis                  | HYNO_COIR R2    | GGTTTGGAAATTGACTGATCC | 124  | v |
| Bighead carp             | HYNO_COI probe  | ACTCATGCTGCTCATCTCAG |      | v |
| Scardinius erythrophthalmus | SCER_COIF     | GAGTTTCTGACTTCTCCCTCG | 167  | v |
| Common rudd              | SCER_COIR       | ATACACCTGCCAGGTCAGAC |      | v |
|                          | SCER_COI probe  | ATGAAACTGATACCCACTCG |      | v |
| Tenca tinca              | TITI_CYTBF      | CAACCGCATTCGCTAGTAAA | 244  | v |
| Tench                    | TITI_CYTBR      | CAAAGGATATTTGCTCTCATG |      | v |
| **(b) Threatened or special concern** | | | |
| Acipenser oxyrinchus     | ACOX_COIF       | TGGTTGACTGACAGGCTATA | 171  | v |
| Atlantic sturgeon        | ACOX_COIR       | CCGAAGCCGCGCATC |      | v |
| Threatened COSEWIC       | ACOX_COI probe  | TGGCAGACAGCAGATT |      | v |
| Alosa sapidissima        | ALSA_COIF       | GCGGCTTTGGGAATTGACTG | 183  | v |
| American shad            | ALSA_COIR       | CAAGATTGCCTGCCAAAGGT |      | v |
| Special concern Quebec   | ALSA_COI probe  | CTCACCGATAGGGCTGATTT |      | v |
| Ameiurus natalis         | AMNA_COIF       | TAGTGGAGCCGGGGATATA | 205  | v |
| Yellow bullhead          | AMNA_COIR       | TGCAAGGTAAGTGAAGATAGT |      | v |
| Susceptible to be special concern Quebec | AMNA_COI probe | TCTCTTCTCTACTAGCCT |      | v |
| Ammocrypta pellucida     | AMPE_COIF       | GGGGATTCGGAAATCTCTCTGT | 162  | v |
| Eastern sand darter      | AMPE_COIR       | GGTACACGGTTTCATCGCGTG |      | v |
| Threatened SARA          | AMPE_COI probe  | AGACATGCGGTTTCTCT |      | v |
| Anguilla rostrata        | ANRO_COIF       | GGTCCATTAAATATGCGGCT | 131  | v |
| American eel             | ANRO_COIR       | CAGCCTGTACCCAGCCA |      | v |
| Threatened COSEWIC       | ANRO_COI probe  | TACGCTCTCTGAGTGAGTA |      | v |
| Esoc americanus vermiculatus | ESAMVE_CYTBF  | CTTGCCTAATTCATCTCATT | 227  | v |
| Grass pickerel           | ESAMVE_CYTBR    | GGGGTTAGGAGGAAAAATGAG |      | v |
| Special concern COSEWIC  | ESAMVE_CYTB probe | ATCTTTATTGACCTTCTAGT |      | v |

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3.3 | Threatened and invasive reptiles and amphibians

Primers were successfully designed for four salamanders including three species listed as threatened by the Species At Risk Act in Canada, Allegheny mountain dusky salamander (Desmognathus ochrophaeus), northern dusky salamander (D. fuscus), spring salamander (Gyrinophilus porphyriticus); as well as two frogs, spring peeper (Pseudacris crucifer), boreal chorus frog (P. maculata); four turtles species listed as endangered, threatened or special concern by the Species At Risk Act in Canada (endangered: Spiny softshell turtle—Apalone spinifera; threatened: Blanding’s turtle—Emydoidea blandingii, wood turtle—Glyptemys insculpta; special concern: northern map turtle—Graptemys geographica; and considered as invasive species: red-eared slider—Trachemys scripta) (Table 3). For all but one of these assays, cross-amplification tests returned negative results. The northern dusky salamander assay showed slight amplification of Allegheny mountain dusky salamander; however, these two species do not co-occur in North America.

Note: Primer name indicates gene amplified, fragment length (bp), and validation through eDNA studies (v: validated, –: not tested).
chorus frog, *P. maculata*. The assay had an amplification efficiency of 96.9% and a limit of detection of 2 copies/rxn (Table 5).

### 3.4 Invertebrate species

Primers for two invasive waterfleas, spiny waterflea (*Bythotrephes longimanus*), and fishhook waterflea (*Cercopagis pengoi*) and two freshwater mussels listed as threatened under the Species At Risk Act (*alewife floater-Utterbackiana implicata* and *Hickorynut-Obovaria olivaria*) were designed (Table 4). Standard curve experiments were performed for the two waterflea species. Assays for *B. longimanus* and *C. pengoi* had an amplification efficiency of 98.1% and 102.7%, respectively, and a limit of detection of 4 copies/rxn for both primer sets (Table 5).

### 4 DISCUSSION

The development of the 60 specific assays presented here was requested for specific needs and questions raised by government...
agencies, academics, or environmental consulting firms. These species are subject to ongoing monitoring either because they are exploited (e.g., Atlantic salmon, lake sturgeon), because of their invasive status (e.g., grass carp, spiny waterflea) or threatened status (e.g., Atlantic sturgeon, Blanding’s turtle, or alewife floater). All of our assays were developed using in silico tests by searching for nonspecific oligonucleotide hybridization using multiple alignments of the target species DNA sequences along with the sequences of related species.

### TABLE 4
Species-specific primers, probes for invertebrate species

| Scientific and Common name | Primer/Probe Gene | Sequence 5’ > 3’ | bp | eDNA |
|----------------------------|-------------------|------------------|----|------|
| **Mollusk**                |                   |                  |    |      |
| Utterbackiana implicata    | ANIM_NADHF        | TTTTATGTATTTCACCTGCTCACACT | 214 | –    |
| Alewife floater            | ANIM_NADHR        | ATGATGGCTCAAGTGAATGTTTATA |     |      |
|                           | ANIM_NADH_probe   | CAAATCTAAATACGCACTACT |     |      |
| **Obovaria olivaria**      | OBOL_COI_F2       | ATTCTGGGGCTCCGTTGGA | 200 | v    |
| Hickorynut                 | OBOL_COI_R2       | ACAGGCAATGCTGAACTGC |     |      |
|                           | OBOL_COI_probe    | CATCTCCTACTTGAAATA |     |      |
| **Crustacean**             |                   |                  |    |      |
| Bythotrephes longimanus    | BYLO_COIF         | GAGACTTATTGGGAGGACCAA | 214 | v    |
|                           | BYLO_COIR         | CCACTCAAATAGGGAATG |     |      |
|                           | BYLO_COI_probe    | TAATCGGAGGTGTTGAA  |     |      |
| Cercopagis pengoi          | CEPE_COIF         | GGAATAGCCTTGCCCTTCTGATG | 188 | v    |
|                           | CEPE_COIR         | GCTCCAGCGTGTCGCGATA |     |      |
|                           | CEPE_COI_probe    | ACTGGATGGACAGCTGAC |     |      |

**Note:** Primer name indicates gene amplified, fragment length (bp) and validation through eDNA studies (v: validated, –: not tested).

### TABLE 5
Percentage of amplification efficiency, limit of detection, intercept (y-inter), and the coefficient of the linear relation between cycle threshold and log DNA dilution ($r^2$) corresponding to for each standard curve developed with a synthetic DNA template

| Scientific name            | Amplification efficiency (%) | Limit of detection (mtDNA copies by rxn) | y-inter | $r^2$ |
|---------------------------|------------------------------|------------------------------------------|---------|-------|
| **Exploited fish species**|                              |                                          |         |       |
| Acipenser fulvescens      | 100.1                        | 8                                        | 38.4    | .975  |
| Ammodytes sp.             | 102.7                        | 4                                        | 39.8    | .985  |
| Clupea harengus           | 103.9                        | 8                                        | 39.7    | .970  |
| Coregonus clupeaformis    | 98.4                         | 8                                        | 40.0    | .971  |
| Mallotus villosus         | 100.0                        | 8                                        | 39.6    | .970  |
| Micropterus dolomieu      | 102.8                        | 20                                       | 40.9    | .949  |
| Morone saxatilis          | 101.7                        | 4                                        | 40.4    | .963  |
| Oncorhynchus mykiss       | 94.6                         | 8                                        | 38.8    | .974  |
| Osmerus mordax            | 103.8                        | 20                                       | 42.1    | .969  |
| Prosopium cylindraceum    | 94.3                         | 4                                        | 40.3    | .969  |
| Salmo salar               | 98.7                         | 2                                        | 38.7    | .969  |
| Salmo trutta              | 108.9                        | 20                                       | 43.1    | .958  |
| Salvelinus alpinus        | 98.4                         | 4                                        | 39.4    | .970  |
| Sebastes spp.             | 95.5                         | 8                                        | 39.9    | .981  |
| **Invasive fish species** |                              |                                          |         |       |
| Ctenopharyngodon idella   | 96.5                         | 4                                        | 40.8    | .949  |
| **Amphibian**             |                              |                                          |         |       |
| Pseudacris maculata       | 96.9                         | 2                                        | 37.1    | .975  |
| **Crustacean**            |                              |                                          |         |       |
| Bythotrephes longimanus   | 98.1                         | 4                                        | 37.8    | .983  |
| Cercopagis pengoi         | 102.7                        | 4                                        | 40.1    | .978  |
that were available in online DNA databases and then predicting probe performance. They were also tested in vitro by amplifying tissue-extracted DNA from both targeted and related species. None of our assays resulted in cross-amplification of DNA for species from the same family, with five exceptions (see Table S1). Since assay development and tests should be specific to a defined geographic area and perhaps population (Goldberg et al., 2016; Wilcox et al., 2015), the cross-amplification tests were done for related species that are present in the same area of the targeted species in Québec. Consequently, before using our assays in other regions, it would be preferable to (a) verify the presence of all related species in the area of interest, (b) verify that cross-amplification tests were done with all related species present in the area of interest and, if not, (c) perform the necessary cross-amplification tests.

The development of eDNA studies is relatively recent and various protocols for eDNA collection, extraction, detection, and analysis have been developed depending on the taxa being studied (Tsuij, Takahara, Doi, Shibata, & Yamanaka, 2019). To the best of our knowledge, qPCR assays targeting the same gene of interest have already been published for 20 of the species addressed in the present study (See Table 6). For ten of them (A. fulvescens, E. lucius, G. insculpta, H. molitrix, M. saxatilis, O. mordax, P. crucifer, S. namaycush, S. salar, and S. trutta), the amplicon was less than 100 bp. In addition, for Trachemys scripta, only the TaqMan probe was designed by us, and we used the primers developed by Davy, Kidd, and Wilson (2015). Here, all of our assays produce amplicons of at least 101 bp which allows the authentication of the positive amplifications by Sanger sequencing in order to avoid false-positive detections. This is particularly crucial for projects where the objective is to detect threatened or invasive species. In addition, we chose to use a probe-based qPCR to allow for more specific detection and quantification of eDNA (Farrington et al., 2015; Mauvisseau, Burian, et al., 2019; Mauvisseau, Tönges, et al., 2019; Wilcox et al., 2013). The amplification efficiency and detection limit tests are usually performed using purified target molecules such as synthetic DNA or reference DNA from biological samples (Bustin et al., 2009). However, to standardize the analysis, the choice of reference DNA from biological samples requires an important amount of DNA and does not allow estimating the number of DNA copies in qPCRs. For these reasons, we used synthetic DNA to standardize our method for our assay development. The results

| Species                | Gene | Amplicon length | Reference                                                                 |
|------------------------|------|-----------------|---------------------------------------------------------------------------|
| Acipenser fulvescens   | COI  | 57              | Yusishen, Eichorn, Anderson, and Docker (2020)                            |
| Carassius auratus      | COI  | 110             | Roy, Belliveau, Mandrak, and Gagné (2018)                                 |
| Ctenopharyngodon idella| COI  | 141             | Roy et al. (2018)                                                          |
| Desmognathus fuscus    | COI  | 170             | Beauclerc, Wozney, Smith, and Wilson (2019)                                |
| Desmognathus ochrophaeus| COI  | 170             | Beauclerc et al. (2019)                                                   |
| Esox lucius            | COI  | 94              | Olsen, Lewis, Massengill, Dunker, and Wenburg (2015)                       |
| Glyptemys insculpta    | COI  | 71              | Lacoursière-Roussel, Dubois, Normandeau & Bernatchez (2016)                |
| Hypophthalmichthys molitrix| COI | 81 | Roy et al. (2018) |
| Hypophthalmichthys nobilis| COI | 117 | Roy et al. (2018) |
| Micropterus dolomieu   | COI  | 147             | Hulley, Tharmalingam, Zarnke, and Boreham (2019)                           |
| Morone saxatilis       | COI  | 63              | Brandl et al. (2015)                                                      |
| Myoxocephalus thompsonii| COI  | 148             | Hulley et al. (2019)                                                      |
| Oncorhynchus mykiss    | CytB | 153             | Minamoto, Hayami, Sakata, and Imamura (2019)                               |
| Osmerus mordax         | COI  | 76              | Hulley et al. (2019)                                                      |
| Perca flavescens       | COI  | 146             | Hulley et al. (2019)                                                      |
| Pseudacris crucifer    | COI  | 99              | Beauclerc et al. (2019)                                                   |
| Salvelinus namaycush   | COI  | 101             | Lacoursière-Roussel, Côté, Leclerc & Bernatchez (2016)                     |
| Salmo salar            | COI  | 74              | Atkinson et al. (2018)                                                    |
| Salmo trutta           | COI  | 61              | Gustavson et al. (2015)                                                   |
| Trachemys scripta      | COI  | 179             | Davy et al. (2015)                                                        |
obtained for each of the 18 assays that were tested (between 2 and 20 mtDNA copies per reaction) were comparable to previous studies on eDNA fish detection with limit of detection between 2 and 50 mtDNA copies per reaction (e.g., Carim et al., 2019; Farrington et al., 2015; Wilcox et al., 2015).

In situ tests were done on 36 of the 60 specific qPCR assays on eDNA studies, which confirmed the assay performance on eDNA samples. Most of these eDNA studies were done at the request of the Province of Québec’s government in order to monitor species with a threatened or invasive status. The results required by these studies were either presence/absence detection or relative quantification. For instance, since the first confirmed capture of a female of the invasive grass carp in 2015 in the St. Lawrence River, our qPCR assay has been thoroughly tested on eDNA to monitor the evolving distribution of this species in this river system (https://mffp.gouv.qc.ca/wp-content/uploads/avis-scientifique-carpes-asiatiques-quebec-confirmation-presence.pdf). Validation of sites with positive amplifications was performed by Sanger sequencing and confirmed the assay performance. Another governmental study required the development of a S. trutta qPCR assay in order to follow the patterns of eDNA diffusion in the St. Lawrence River (Laporte et al., 2020). This assay has been thoroughly tested and showed the efficiency of these primers to detect eDNA of confined S. trutta down to 5 km from the emission point (Laporte et al., 2020). Moreover, some assays developed for exploited fish species such as S. salar and M. dolomieu were also thoroughly tested on eDNA samples to assess their spatio-temporal distributions and habitat use (O’Sullivan et al., 2020). The performance of these assays was also validated by Sanger sequencing. In addition, qPCR assays developed for other clades showed good performance for detecting the presence or absence of specific species found in Québec. The spiny and fishhook waterfleas are of big concern since their introduction, probably through ballast water or recreational boats. These invasive species are already being monitored in the Laurentian Great Lakes using nets, sediment, or eDNA analysis (Walsh, Spear, Shannon, Krysan, & Vander Zanden, 2019). Here, our qPCR assay allowed the detection of B. longimanus in water samples from diverse regions of the Province of Quebec (Hernandez, Bougas, Perrault-Payette, Normandeau, & Bernatchez, 2018). These results were validated by Sanger sequencing as well as actual specimen collections done in the field in 2018.

5 | CONCLUSION

The use of eDNA analysis is booming and already modifying the design and implementation of biodiversity monitoring programs. The greatest advantage of this tool probably lies in the capacity to monitor threatened and invasive freshwater species without disturbing individuals at risk or their environment. Thus, the costs in terms of both technical resources and ecological impacts in the field are considerably reduced when compared to, for example, methods using gillnets to monitor fish species. eDNA analysis by qPCR is now widely and successfully used to detect a wide range of target species (Tsuji et al., 2019). Despite the challenge to design optimal specific primers throughout a species’ geographic range due to differences in co-occurring sister species, rare mitochondrial introgression, or local haplotypic variation, we hope that our 60 qPCR assays will be of broad usefulness not only for monitoring studies in Québec but also wherever these species are present in North America or have been introduced on other continents.

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AUTHOR CONTRIBUTION

C. H and B. B. should be considered joint first author. L. B., G. C. and A. S designed the project. G. C and A. S. shared tissue samples. B. B. and C. H. drafted the manuscript and all authors contributed to the writing and approved the final draft of the manuscript. C. H., B. B., A. P. P realized the primers design and the experiments.

DATA AVAILABILITY STATEMENT

Data has been upload to Dryad: https://doi.org/10.5061/dryad.12jm63xtw.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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