

**An environmental DNA tool for monitoring the status of the Critically Endangered Smalltooth Sawfish,  
*Pristis pectinata*, in the Western Atlantic**

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## 1 **Abstract**

2 The Critically Endangered Smalltooth Sawfish, *Pristis pectinata*, was once widespread in the tropical and  
3 subtropical waters of the Atlantic Ocean, but following substantial declines over the past century, the core  
4 population is currently confined to southwest Florida in the U.S. and the Bahamas. Recent research and verified  
5 public encounter reports suggests that this core population may be stabilizing and, potentially, expanding into  
6 formerly occupied areas of their historic range in the Western Atlantic; however, the status of this species in non-  
7 core waters is not well understood. Environmental DNA (eDNA) methods provide a relatively cost effective and  
8 rapid assessment tool for monitoring species occurrence in aquatic habitats. Here, we have developed an eDNA tool:  
9 a species-specific Droplet Digital™ PCR (ddPCR™) assay targeting a 100-base pair portion of the mitochondrial  
10 NADH dehydrogenase subunit 2 gene in *P. pectinata*, with the ability to reliably detect as little as 0.25 pg of target  
11 DNA. The assay was validated by collecting and analyzing a water sample from known *P. pectinata* nursery habitat  
12 in Florida, which was found to contain an average of 11.54 copies of target DNA/μL (SE = 0.72) in the reaction.  
13 The assay was then further tested by placing a juvenile sawfish in an *ex situ* tank and analyzing water samples  
14 collected at time intervals. The implementation of this eDNA tool into field surveys will provide additional, reliable  
15 data to assess species recovery and aid in prioritizing localities beyond the core range in which to focus research and  
16 education initiatives.

17

## 18 **Introduction**

19 Sawfishes are among the most threatened families of marine fishes worldwide (Dulvy et al. 2014), with all five  
20 species listed as Critically Endangered or Endangered on the International Union for Conservation of Nature (IUCN)  
21 Red List of Threatened Species (Dulvy et al. 2016). All sawfishes have undergone global declines in range and  
22 abundance due to direct exploitation, bycatch in fisheries, and habitat loss (Dulvy et al. 2016). These threats are  
23 exacerbated by their life history traits (e.g., late maturity, low fecundity, and long life spans), which leave sawfishes  
24 susceptible to overexploitation, and makes population recovery a slow process (Stevens et al. 2000; Carlson and  
25 Simpfendorfer 2015).

26 The Critically Endangered Smalltooth Sawfish, *Pristis pectinata*, is thought to have experienced the largest  
27 global range contraction of all sawfishes and is currently found in less than 20% of its former range (Dulvy et al.  
28 2016). Once widespread in the tropical and subtropical waters of the Atlantic Ocean, remaining core population(s)

29 are thought to be limited to the U.S. and the Bahamas (Carlson et al. 2013), making these populations of global  
30 conservation significance (Dulvy et al. 2016). Within U.S. waters, *P. pectinata* were historically found from Texas  
31 to the Carolinas (Brame et al. 2019) but saw substantial losses in both range and abundance over the past century,  
32 with the current population restricted to southwest Florida (SWFL) by the 1980's (Norton et al. 2012).

33 Due to the dramatic declines in range and abundance, *P. pectinata* was listed as Endangered in 2003 under  
34 the U.S. Endangered Species Act of 1973 by the National Marine Fisheries Service (NMFS) (NMFS 2003), and a  
35 Species Recovery Plan (SRP) was developed to promote recovery and long-term viability of the species in U.S.  
36 waters (NMFS 2009; 2018). One characteristic of full species recovery is re-establishment in some or all of the  
37 former range (Akçakaya et al. 2018); therefore, the SRP for *P. pectinata* designated 15 recovery regions throughout  
38 their historic range in U.S. waters, wherein recovery efforts should occur if species presence is confirmed (NMFS  
39 2009; 2018). As a result of over 15 years of U.S. federal and state protections, scientific advances in the  
40 understanding of the biology and ecology of the species, and public education initiatives, the core population of *P.*  
41 *pectinata* in SWFL is believed to be stabilizing (NMFS 2018). One line of evidence for this potential stabilization is  
42 the emergence of relatively recent sawfish encounter reports within formerly occupied parts of their historic range in  
43 U.S. waters, including in designated recovery regions (NMFS 2018); however, the status of *P. pectinata* in these  
44 non-core areas is unknown.

45 Traditional survey methods for monitoring the status of rare species can be expensive and time-consuming  
46 (Lewison et al. 2004). Environmental DNA (eDNA) methods provide a relatively cost effective and rapid  
47 assessment tool for monitoring species occurrence in aquatic habitats (Rees et al. 2014; Evans et al. 2017). Water  
48 provides a medium for traces of DNA recently shed by organisms (e.g., cellular debris, skin cells, blood, feces,  
49 urine), which can be collected and analyzed via genetic assays (Jerde et al. 2011). EDNA has been shown to be as  
50 sensitive, and sometimes more sensitive in rare species detections compared to survey methods such as  
51 electrofishing (Evans et al. 2017), Baited Remote Underwater Video systems (BRUVs) and underwater visual  
52 censuses (UVCs) (Boussarie et al. 2018), and traditional net surveys (Thomsen et al. 2012). EDNA methods also  
53 negate the need to capture and handle the target species, making it an ideal tool to assess the presence or absence of  
54 a threatened species (Rees et al. 2014). EDNA has been used in targeted, single species detections for a growing  
55 number of threatened elasmobranchs, including the Endangered Maugean Skate, *Zearaja maugeana* (Weltz et al.  
56 2017), the Vulnerable Great White Shark, *Carcharodon carcharias* (Lafferty et al. 2018), the Vulnerable Chilean

57 Devil Ray, *Mobula tarapacana* (Gargan et al. 2017), and the Critically Endangered Largetooth Sawfish, *Pristis*  
58 *pristis* (Simpfendorfer et al. 2016).

59 Here, we develop and validate an eDNA assay to detect the presence of *P. pectinata* DNA in water  
60 samples, for use as a tool for monitoring their recovery in the Western Atlantic. This tool will allow scientists and  
61 managers to better understand the status of *P. pectinata* in non-core areas and provide quantitative baseline data  
62 from which to measure progress towards recovery. Such data can also aid in prioritizing recovery regions in which  
63 to focus research and education initiatives, playing an important role in adaptive management strategies as *P.*  
64 *pectinata* expands into its former range.

65

### 66 ***Pristis pectinata* eDNA assay development**

67 Field and laboratory controls

68 To reduce the risk of contamination by exogenous DNA or cross-contamination between samples, rigorous controls  
69 were used throughout all stages of this research (see Ficetola et al. 2016; Goldberg et al. 2016; Port et al. 2016;  
70 Schweiss et al. In press). All water collection bottles and filtering equipment were cleaned prior to each use using a  
71 combination of two methods of sterilization; cleaning with 10% bleach followed by either autoclaving at 120°C for  
72 20 min or exposure to UV light for 20 min, depending on the materials. To prevent contamination between the  
73 stages of sample processing, water filtration, DNA extractions, and PCR amplifications were conducted in  
74 physically isolated laboratories. Furthermore, water samples were filtered in laboratories where contemporary *P.*  
75 *pectinata* tissue had never been present (see Deiner et al. 2015). During water filtration and DNA extraction,  
76 designated sterile forceps for each sample were used to handle used filters and gloves were changed between  
77 samples to prevent cross-contamination between samples (see Pilliod et al. 2013; Goldberg et al. 2016). During  
78 DNA extractions and PCR, aerosol barrier pipette tips were used to prevent cross-contamination between samples  
79 (Schweiss et al. In Press). Additionally, no positive samples were included in any PCRs due to the risk of  
80 contamination from the positive itself, as per ancient DNA (aDNA) PCR protocols (see Mulligan 2005).

81 To test for the possibility of contamination, negative control samples were incorporated into water sample  
82 collection and each stage of laboratory processing and analyzed through to PCRs, which were conducted in  
83 replicates of five (Jerde et al. 2011; Bakker et al. 2017). To test for contamination in the field, 3 L of autoclaved DI  
84 water were brought onto the boat and stored in three sterile 1 L Nalgene® bottles on ice until filtration. To test for

85 contamination during filtration, 3 L of autoclaved DI water were filtered and processed through to PCR. Negative  
86 controls for DNA extractions contained no particulate matter or filters, and PCR negatives contained no DNA  
87 template. Analysis of all negative control samples, using the optimized protocols described below, found no  
88 evidence of target DNA across all PCR replicates.

89

90 Water collection, filtration, and DNA extraction

91 Three liter (L) water samples were collected for all aspects of this study using three sterile, 1 L high-density  
92 polyethylene Nalgene<sup>®</sup> bottles, which were kept on ice in pre-cleaned marine coolers until filtration, which occurred  
93 within 24 hours of collection. All water samples were vacuum-filtered using 47 millimeter (mm) 0.8 micron (µm)  
94 nylon filters, and used filters were rolled and preserved in 95% ethanol at room temperature. Total eDNA was  
95 extracted from filters using the QIAGEN<sup>®</sup> DNeasy<sup>®</sup> Blood & Tissue Kit following the Goldberg et al. (2011)  
96 protocol incorporating QIAshredder<sup>™</sup> spin columns. The qualities of DNA extracts were visualized using 2%  
97 agarose gels and the quantities of DNA were assessed using Thermo Fisher Scientific<sup>™</sup> NanoDrop<sup>™</sup> technology.

98

99 Droplet Digital PCR assay

100 Primers were designed to amplify a 100-base pair (bp) fragment of the mitochondrial NADH dehydrogenase subunit  
101 2 (mtDNA ND2) gene in *P. pectinata*, but not in other elasmobranchs that could co-occur with this species in U.S.  
102 waters, or in other *Pristis* sawfishes. To design these primers, mtDNA ND2 sequences for *P. pectinata* (GenBank  
103 accession no. KP400584.1) and 17 genetically similar exclusion species were downloaded from GenBank (Online  
104 Resource 1) and aligned in CodonCode v. 6.0.2 (CodonCode Corporation, Dedham, MA, USA). Forward  
105 (PpecND21F: 5'-CTGGTTCACATTGACTCTTAATTTG-3') and reverse (PpecND21R: 5'-  
106 GCTACAGCTTCAGCTCTCCTTC-3') primers and an internal PrimeTime<sup>®</sup> double-quenched ZEN<sup>™</sup>/IOWA  
107 Black<sup>™</sup> FQ probe labeled with 6-FAM (PpecND2Probe1IBQF: 5'-TACCATAGCCATCATCCATTATTATTC-  
108 3') were designed to amplify DNA in only *P. pectinata* by including bp differences in the primers and the probe in  
109 all exclusion species (see Online Resource 1). To confirm that the combination of the primers and probe amplified  
110 the desired locus, PCRs were conducted using quantitative real-time PCR (qRT-PCR) and total genomic DNA  
111 (gDNA) from four *P. pectinata* individuals. Reaction mixtures contained 1.1 µL of extracted DNA (~25 ng/µL), 1X  
112 Bio-Rad<sup>®</sup> iTaq<sup>™</sup> universal probe supermix, 900 nanomolar (nM) of each primer, and 170 nM of probe, adjusted to

113 22  $\mu$ L using PCR-grade water. Cycling conditions consisted of enzyme activation at 95°C for 10 min, followed by  
114 40 cycles of: 94°C for 30 s and 64°C for 2 min, followed by enzyme deactivation at 98°C for 10 min, using a ramp  
115 rate of 1°C/s. The resulting amplicon from one *P. pectinata* individual was cleaned using a QIAGEN® QIAquick  
116 PCR Purification Kit using the manufacturer’s protocol, with the exception that all centrifugation steps were  
117 conducted at 12,000 rpm for 2 min. Forward and reverse sequences were generated using a BigDye™ Terminator  
118 v3.1 Cycle Sequencing Kit (Applied Biosystems™, Foster City, CA, USA) on an Applied Biosystems™ 3730XL  
119 DNA Analyzer. A consensus sequence was assembled in CodonCode v. 6.0.2 (CodonCode Corporation, Dedham,  
120 MA, USA) and its identity was verified as *P. pectinata* using the NCBI BLAST search function; the generated  
121 sequence was 99.3% similar to *P. pectinata* GenBank accession no. KP400584.1 (Chen et al. 2016).

122 The PCR reaction and cycling conditions were optimized for the Bio-Rad® QX200™ AutoDG™ Droplet  
123 Digital™ PCR System (Instrument no. 773BR1456) by systematically adjusting seven variables (i.e., primer and  
124 probe concentrations, cycle number, ramp rate, annealing temperature, denaturation time, and elongation time) to  
125 produce positive results with high relative fluorescence units (RFUs) and little to no “droplet rain” (i.e., droplets, or  
126 clusters of droplets, that lie between the positive and negative droplet bands on the ddPCR™ scatter plot) (see  
127 Online Resource 2). Optimized ddPCR™ reaction mixtures contained 1.1  $\mu$ L of extracted DNA, 1X Bio-Rad®  
128 ddPCR™ supermix for probes (no deoxyuridine triphosphate (dUTP)), 900 nanomolar (nM) of each primer, and 170  
129 nM of probe, adjusted to 22  $\mu$ L using PCR-grade water. Optimal ddPCR™ cycling conditions were enzyme  
130 activation at 95°C for 10 min, followed by 40 cycles of: 94°C for 30 s and 64°C for 2 min, with a final enzyme  
131 deactivation step at 98°C for 10 min, using a ramp rate of 1°C/s. To ensure the assay was species-specific for *P.*  
132 *pectinata* in U.S. waters, the optimized ddPCR™ reaction and cycling conditions were tested using 0.20 ng/ $\mu$ L  
133 gDNA derived from fin clips from four *P. pectinata* individuals and one individual for each of 12 representative  
134 exclusion species (Table 1). The target DNA fragment was amplified in all five ddPCR™ replicates for each *P.*  
135 *pectinata* individual but was not amplified in any of the ddPCR™ replicates for any representative species from five  
136 genetically similar ray genera and two shark genera that could co-occur with *P. pectinata*, or in other *Pristis*  
137 sawfishes.

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141 Data analysis

142 Data were analyzed using three criteria for positive *P. pectinata* detections: 1) droplets fell above a manual threshold  
143 (MT) defined for this assay, 2) droplets above the MT fell within the prescribed range of the positive droplet  
144 population for this assay (5000–7000 RFUs; Fig. 1), and 3) the concentration of target DNA, determined using Bio-  
145 Rad® QuantaSoft™ software using the Rare Event Detection (RED) setting, was at or above the Limit of Detection  
146 (LoD) of the assay. Defining an assay-specific MT minimizes the likelihood of incorrectly calling artifact droplets  
147 (i.e., droplets that fall above the negative band population in the absence of target DNA; see Online Resource 3) as  
148 positive detections (e.g., Hunter et al. 2017). To define an appropriate MT for the *P. pectinata* eDNA assay, a No  
149 Template Control (NTC) plate with no target DNA was analyzed on the ddPCR™ platform, using the described  
150 reaction and cycling conditions. The highest amplitude of the artifact droplets was 2,700 RFUs; therefore, 3000  
151 RFUs was chosen to adopt a more conservative approach to minimize the risk of a calling a false positive.

152 To determine the LoD of the assay, ddPCR™ reactions were performed using gDNA from three *P.*  
153 *pectinata* individuals with a 6-fold series of 10X dilutions from starting concentrations of 20 ng/μL (i.e., 1:10 to  
154 1:1,000,000). Target DNA was reliably detected in all replicates for all individuals up to the 1:10,000 dilutions, but  
155 not in the 1:100,000 dilutions. The standard error of the 1:1,000,000 also overlapped with zero (Fig. 2a), making  
156 detection at this concentration unreliable. To further refine the LoD, ddPCR™ reactions were performed on  
157 subsequent 3-fold series of 2X dilutions from the 1:10,000 dilutions (Fig. 2b). The LoD of this assay was found to  
158 be the 1:80,000 dilutions, corresponding to 0.25 pg of target DNA in the reaction (Fig. 2c). The standard errors of  
159 the 1:80,000 dilutions did not include zero, or overlap with the 1:100,000 dilutions; so using the average number of  
160 copies of target DNA/μL in the 1:80,000 dilutions and applying the lower standard error as the relaxed detection  
161 threshold (see Baker et al. 2018), the LoD of the assay was determined to be 0.08 copies/μL.

162

### 163 **Validation of the *Pristis pectinata* eDNA assay**

164 To validate the ddPCR™ assay, positive *P. pectinata* eDNA samples were acquired via analysis of a water sample  
165 from known habitat and through a tank experiment. To collect these positive water samples, a pre-cleaned ~160 L  
166 tank was filled with ambient surface water from known *P. pectinata* nursery habitat in the Caloosahatchee River,  
167 Florida, approximately 330 m outside of the Harbour Isles Marina. A 3 L water sample was immediately collected  
168 from the tank to assess whether *P. pectinata* eDNA was present in the Caloosahatchee River water. One juvenile

169 female *P. pectinata*, measuring 786 mm stretched total length, was captured in a gill net inside the Harbour Isles  
170 Marina and placed into the tank. An aerator was added to the tank and dissolved oxygen and water temperature were  
171 monitored for the duration of the experiment. A 3 L water sample was collected from the tank immediately after the  
172 juvenile was added (time zero) and again after 30 min. All water samples were filtered, DNA extracted, run on  
173 ddPCR™, and analyzed using the methods developed in this study.

174 Applying all three criteria for a positive detection of target DNA, the ddPCR reactions containing DNA  
175 extracted from ambient water from the nursery contained an average of 11.54 copies/μL (SE = 0.72) of *P. pectinata*  
176 eDNA (Fig. 3). The amount of target eDNA increased to 739.4 copies/μL (SE = 38.31) immediately after the  
177 juvenile was added to the tank (time zero) and then increased to 3,175.8 copies/μL (SE = 589.3) after 30 min (Fig.  
178 3). At 30 min, the large quantity of target DNA isolated from the water sample oversaturated the PCR product,  
179 resulting in a high standard error.

180

## 181 Discussion

182 The developed eDNA assay provides a rapid-assessment tool to conduct targeted surveys to investigate the  
183 occurrence and infer the status of *P. pectinata* beyond their contemporary core range. This assay has been validated  
184 in the Caloosahatchee River, Florida, where *P. pectinata* is the sole species of sawfish; however, because the assay  
185 did not amplify DNA in other *Pristis* sawfishes, it can also be used in locations where other sawfishes have been  
186 known to co-occur, at least historically, such as Texas (Brame et al. 2019). It is promising to note that there are no  
187 bp differences in the primer or probe sequences designed in this study compared to a mtDNA ND2 sequence for a *P.*  
188 *pectinata* collected in Mexico (GenBank accession no. MF682494.1; Diaz-Jaimes et al. 2018), indicating the  
189 developed assay should amplify the target gene in this species from other, nearby waters in the Western Atlantic.  
190 Use of the assay outside these waters requires careful consideration and, potentially, further testing. MtDNA genes  
191 are often variable among populations within a species (Rubinoff et al. 2006); therefore, before using this assay to  
192 conduct eDNA surveys in other geographic regions (e.g., Eastern Atlantic), the primers and probe should be tested  
193 with *P. pectinata* tissue samples obtained from the local population. Where fresh *P. pectinata* tissue samples are not  
194 available for such testing due to the possibility of local extinctions, historic rostra can be used as an alternative  
195 source of DNA (Phillips et al. 2009). Finally, the primers and probe developed here were cross-tested with  
196 representative species from closely related genera found in U.S. waters; testing with additional exclusion species is



197 required to ensure that the assay remains species-specific in other geographic regions, highlighting the need for local  
198 fisheries knowledge (Poulakis and Grubbs 2019).

199 The use of ddPCR™ for single species detections is gaining popularity in eDNA research due to its unparalleled  
200 ability to detect minute quantities of target DNA amongst high concentrations of non-target DNA and in the  
201 presence of natural inhibitors found in water samples (Evans et al. 2017; Hunter et al. 2018). DdPCR™ assays  
202 developed for species such as the Bull Shark, *Carcharhinus leucas* (Schweiss et al. In Press) and Killer Whale,  
203 *Orcinus orca* (Baker et al. 2018), have found this platform to be capable of detecting less than 0.5 pg of target DNA  
204 in a reaction. Such highly sensitive assays are especially critical in eDNA surveys targeting Critically Endangered or  
205 Endangered species, where there can be substantial conservation outcomes based on the results of such surveys  
206 (Hunter et al. 2018; Poulakis and Grubbs 2019). For instance, the use of ddPCR™ could reduce the risk of false  
207 negatives (i.e., where target DNA is present but not detected) stemming from the use of a less sensitive PCR  
208 platform such as conventional or qRT-PCR, which are unlikely to detect such minute quantities of target DNA (Doi  
209 et al. 2015). Conservation and management strategies developed on the basis of such false negatives could lead to  
210 slower implementation and inadequate protections along with incomplete habitat designations for threatened species,  
211 ultimately hindering species recovery.

212 Using the three-criteria approach described here to define positive detections on the ddPCR™ platform  
213 provides a rigorous approach to interpret the results of eDNA surveys, reducing the risk of incorrectly calling PCR  
214 artifacts as positive species detections (e.g., false positives). For example, using only a MT, an artifact droplet just  
215 above the threshold could be incorrectly interpreted as a positive detection. Ensuring the quantity of target DNA is  
216 also within the detection capabilities of an assay allows for more robust and confident positive detections. False  
217 positives can also result from contamination between eDNA samples or from exogenous DNA. Given the detection  
218 capabilities of ddPCR™ assays, strict protocols to prevent contamination (see Goldberg et al. 2016; Schweiss et al.  
219 In press) coupled with testing for contamination at every stage in sample processing are critical in producing reliable  
220 data from eDNA surveys that may be used as a part of conservation planning. This is especially important when the  
221 results of eDNA surveys could be used to prioritize research and management initiatives as well as in the allocation  
222 of resources (Poulakis and Grubbs 2019).

223 With a well-designed water sampling regime, strict field and laboratory controls, and a highly sensitive  
224 ddPCR™ assay, targeted species eDNA surveys provide a powerful tool to improve our knowledge of the

225 occurrence of *P. pectinata*. The eDNA tool developed here can be used to provide quantitative baseline data in non-  
226 core areas from which to measure future progress towards species recovery. Recovery in *P. pectinata* populations is  
227 predicted to be a slow process due to their life history characteristics and will be dependent on the mitigation of  
228 anthropogenic activities (e.g., accidental fisheries mortalities; Carlson and Simpfendorfer 2015). Range re-expansion  
229 during recovery is predicted to begin in locations closest to the core population(s) as a result of spillover from  
230 adjacent areas, in a stepping-stone fashion (see Saura et al. 2014). There is, however, the possibility that because  
231 female *P. pectinata* have been shown to exhibit philopatry (Feldheim et al. 2017), occurrence and encounter reports  
232 of juvenile *P. pectinata* in non-core areas further away from SWFL (e.g. Texas, Mississippi) represent remnant *P.*  
233 *pectinata* populations scattered over portions of their former range; under such a scenario, patterns of recovery could  
234 be more complex and would ultimately depend on the availability of suitable habitat and the mitigation of threats  
235 from anthropogenic activities (Seitz and Poulakis 2006; Poulakis et al. 2011; Norton et al. 2012; Scharer et al.  
236 2017). Conducting targeted eDNA surveys for *P. pectinata* across all historically-occupied regions in U.S. waters  
237 could not only aid in conservation planning and prioritizing areas for research, but could also increase our  
238 understanding of patterns of recovery in a highly threatened marine species.

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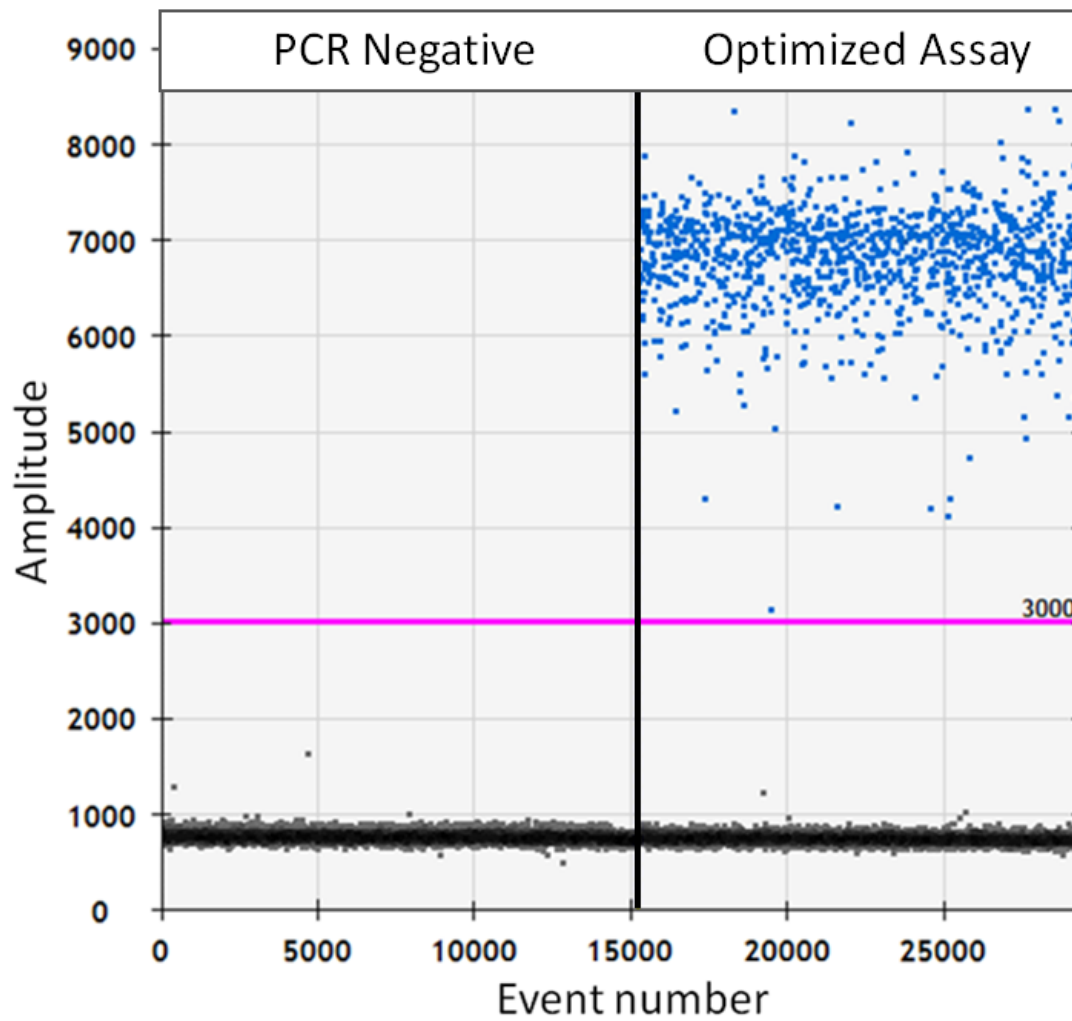
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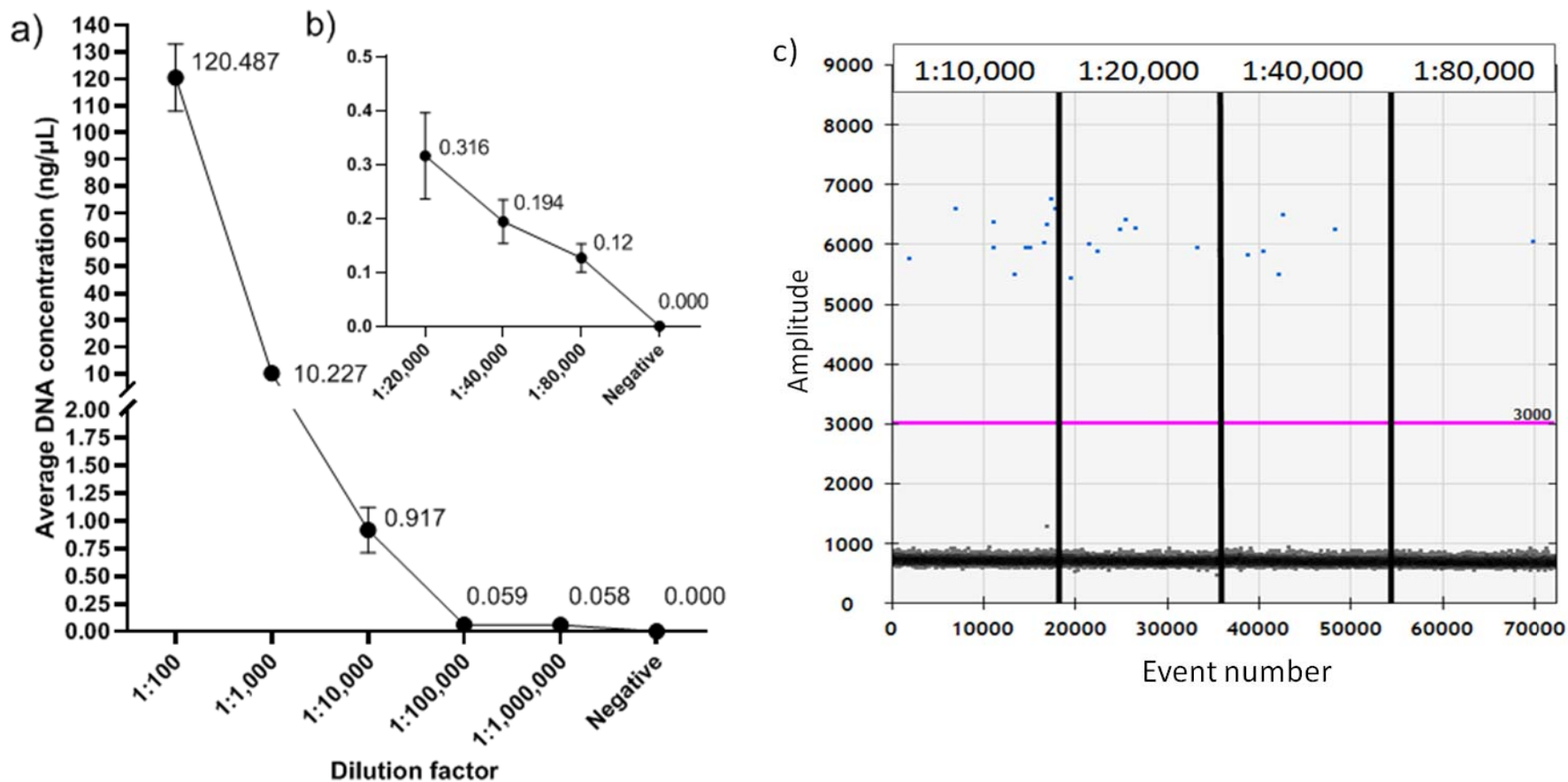
**Table 1** List of 12 exclusion species that were tested to ensure species-specificity of the primers and probe developed for the mitochondrial NADH dehydrogenase subunit 2 (mtDNA ND2) gene in *Pristis pectinata* on the Bio-Rad® QX200™ AutoDG™ Droplet Digital™ PCR System. The country of origin for each tissue sample is included

<b>Species</b>	<b>Tissue origin</b>
Green Sawfish, <i>Pristis zijsron</i>	Australia
Dwarf Sawfish, <i>Pristis clavata</i>	Australia
Largetooth Sawfish, <i>Pristis pristis</i>	Australia
Atlantic Guitarfish, <i>Rhinobatos lentiginosus</i>	United States
Atlantic Stingray, <i>Hypanus sabinus</i>	United States
Bluntnose Stingray, <i>Hypanus say</i>	United States
Cownose Ray, <i>Rhinoptera bonasus</i>	United States
Spotted Eagle Ray, <i>Aetobatus narinari</i>	United States
Clearnose Skate, <i>Raja eglanteria</i>	United States
Roundel Skate, <i>Raja texana</i>	United States
Bigeye Thresher Shark, <i>Alopias superciliosus</i>	United States
Atlantic Sharpnose Shark, <i>Rhizoprionodon terraenovae</i>	United States



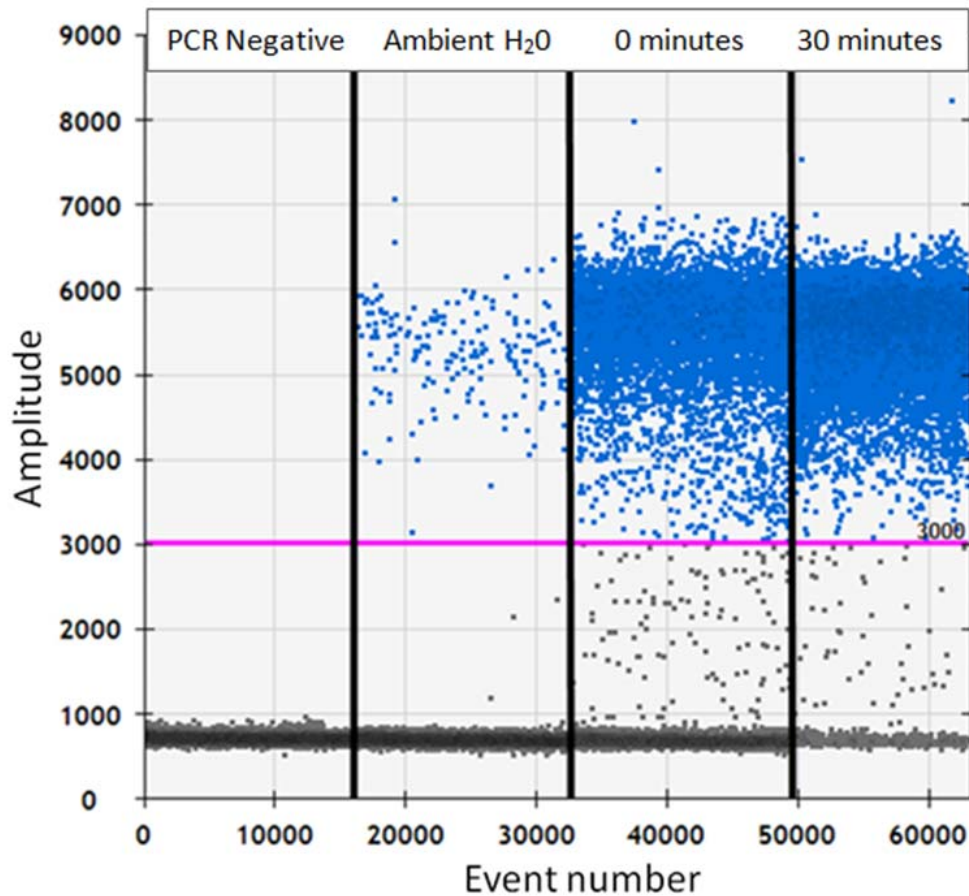


**Fig. 1** Raw droplet scatter plot of ddPCR™ products using genomic DNA from one Smalltooth Sawfish, *Pristis pectinata*, individual with the optimized assay conditions with a corresponding negative control. Each droplet in each well was classified as either positive (above 3000) or negative (below 3000) for target DNA based on a manual threshold amplitude of 3000 relative fluorescence units (RFUs); detected using a Bio-Rad® QX200™ Droplet Reader, QuantaSoft™ software and RED analysis setting. Each well is separated by vertical lines, and is labeled to correspond with the sample it represents



**Fig. 2** Average target DNA concentrations (copy number/μL) of the Limit of Detection (LoD) dilution series, using genomic DNA from three Smalltooth Sawfish, *Pristis pectinata*, individuals with five replicates each in a) a 6-fold series of 10X dilutions from a starting concentration of 20 ng/μL, b) a 3-fold series of 2X dilutions from the 1:10,000 dilution, and c) a corresponding raw ddPCR™ scatter plot of serial dilution reactions from one replicate of one Smalltooth Sawfish, *Pristis pectinata*, individual. The Bio-Rad® QX200™ Droplet Reader and QuantaSoft™ software using the RED analysis setting was used across all samples, and each droplet in each well was classified as either positive (above 3000) or negative (below 3000) for target DNA based on a manual threshold amplitude of 3000 relative fluorescence units (RFUs)





**Fig. 3** Raw droplet scatter plot of ddPCR<sup>TM</sup> serial dilution products from one reaction for each of the Smalltooth Sawfish, *Pristis pectinata*, positive eDNA samples. Ambient water refers to water samples collected from the Caloosahatchee River, known *P. pectinata* nursery habitat. 0 and 30 min reactions correspond to the positive water samples collected from the *ex situ* tank containing a live *P. pectinata*. Each droplet in each well was classified as either positive (above 3000) or negative (below 3000) for target DNA based on a manual threshold amplitude of 3000 relative fluorescence units (RFUs) detected using a Bio-Rad<sup>®</sup> QX200<sup>TM</sup> Droplet Reader, and QuantaSoft<sup>TM</sup> software using the RED analysis setting. Each well is separated by vertical lines and is labeled to correspond with the sample or time stage it represents. Note: “Droplet rain” (i.e., droplets, or clusters of droplets, that lie between the positive and negative droplet bands on the ddPCR<sup>TM</sup> scatter plot) is seen at 30 min due to an oversaturation of target DNA