

1 **Environmental DNA analysis needs local reference data to**
2 **inform taxonomy-based conservation policy – A case study**
3 **from Aotearoa / New Zealand**

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26 **Abstract**

27 Effective management of biodiversity requires regular surveillance of multiple species.
28 Analysis of environmental DNA by metabarcoding (eDNA) holds promise to achieve this
29 relatively easily. However, taxonomic inquiries into eDNA data need suitable molecular
30 reference data, which are often lacking. We evaluate the impact of this reference data void in
31 a case study of fish diversity in the remote fiords of New Zealand. We compared eDNA-
32 derived species identifications against Baited Remote Underwater Video (BRUV) data
33 collected at the same time and locations as the eDNA data. Furthermore, we cross referenced
34 both eDNA and BRUV data against species lists for the same region obtained from literature
35 surveys and the Ocean Biodiversity Information System (OBIS). From all four data sources,
36 we obtained a total of 116 species records (106 ray-finned fishes, 10 cartilaginous fishes; 59
37 from literature, 44 from eDNA, 25 from BRUV, 25 from OBIS). Concordance of taxonomies
38 between the data sources dissolved with lowering taxonomic levels, most decisively so for
39 eDNA data. BRUV agreed with local biodiversity information much better and fared better in
40 detecting regional biodiversity dissimilarities. We provide evidence that eDNA
41 metabarcoding will remain a powerful but impaired tool for species-level biodiversity
42 management without locally generated reference data.

43 **Introduction**

44 Marine reserve (MR) networks conserve biodiversity by stabilizing communities and
45 maintaining food web structure (Wing & Jack, 2013). Effective management of MR
46 biodiversity requires regular surveillance, for example to avoid overexploitation by fishing
47 (Wing & Jack, 2013), or to avoid damage through influx of non-indigenous species
48 (Cunningham, 2019). Fish surveillance is of particular interest due to their sensitivity to most
49 forms of human disturbance, their usefulness at all levels of biological organization, and the
50 favourable benefit-to-cost ratio of fish assessment programmes (Harris, 1995).

51 Analysis of environmental DNA (eDNA) metabarcoding data is a well-established molecular
52 technique for multispecies surveys (Cristescu & Hebert, 2018). Environmental DNA
53 metabarcoding holds promise for biodiversity surveys intended to inform biodiversity
54 management – associated techniques are regarded more cost-efficient than traditional
55 methods (such as baited remote underwater video surveys – BRUV), less dependent on expert
56 taxonomic knowledge, can be standardized, and are able to inform on a broad range of taxa
57 (Sigsgaard et al., 2020).

58 Reliable low-level taxonomic annotation is a prerequisite for useful biodiversity management
59 and biological surveillance (e.g., Jack & Wing 2013). For example, in a southern New
60 Zealand context, *Parapercis colias* (blue cod) is of high commercial interest, but three other
61 of the 79 cod species are known from New Zealand (Roberts et al., 2019), so that genus
62 information alone is ambiguous for determining blue cod presence or absence. Accordingly,
63 higher-level taxonomic classifications (e.g., family, and order levels) are even less
64 informative for species level conservation management. This reality translates into the desire
65 for obtaining perfect 100 bp to 200 bp alignments (Huson et al. 2007) between an unknown
66 eDNA-derived query sequence and a well described reference sequence derived from a valid
67 species. In practice, absence of such reference data necessitates relaxation of taxonomy-
68 assigning alignment parameters to retain sufficient eDNA data for analysis, and in
69 consequence the data's informative quality suffers.

70 Availability of suitable reference data for metabarcoding is highly variable depending on
71 taxonomic groups and geographic locations, with fish considered relatively well covered in
72 Barcode of Life Data Systems (BOLD) and NCBI's GenBank (Benson et al. 2011) for some
73 regions such as Europe (Weigand et al., 2019). Arguably, fewer reference data are available
74 for fish of southern New Zealand. For example, for six commonly used 12S primer pairs,
75 recognized as well suitable for fish multispecies surveys (Weigand et al., 2019), an average
76 of 36% of all northern European fish species are available as reference data, but only 26% of
77 southern New Zealand species (GAPeDNA v1.0.1 web interface, 11-Sep-2021; Marques et
78 al. 2021; also see SI Table 1).

79 In this study, we evaluate the impact of taxonomic data limitations on multispecies surveys
80 using the example of fish in the UNESCO World Heritage Site Te Wahipounamu (Fiordland)
81 in southern New Zealand. We compare the results of concurrent eDNA and BRUV surveys
82 and cross-reference these data against species lists for the same region obtained from
83 literature and the Ocean Biodiversity Information System (OBIS). The fish diversity of Te
84 Wahipounamu has been described based on a diverse range of mostly visual methods. If our
85 BRUV and eDNA approaches work optimally, we should see a strong overlap between these
86 field data and previously described fish diversity records of the region.

87 **Methods**

88 For this study, we evaluated the presence of Actinopterygii (ray-finned fishes) and
89 Chondrichthyes (cartilaginous fishes) species in one MR, two commercial exclusion zones

90 (all “MR”), and corresponding control areas in southern Te Wahipounamu, New Zealand
91 (west coast, approximately from -44.3 to -46.25 Southern latitude; Fig. 1a). We obtained and
92 analysed eDNA and BRUV data as well as electronic records proximate to the field work
93 area from the Ocean Biodiversity Information System (OBIS; Ausubel 1999). Furthermore, a
94 reference list of ray-finned fishes and cartilaginous fishes that have been observed in
95 Fiordland was assembled from literature. All observations were formalized using NCBI
96 taxonomy (Federhen, 2012), including trivial names, and limited to classes Actinopterygii
97 and Chondrichthyes.

98 Literature data, itself obtained using various methods, were extracted from five sources,
99 including one meta-analysis (SI Table 2). OBIS data were downloaded for a 38 km radius
100 around all field work sites (centre point W 166.89°, S -45.80°), as well as for smaller areas
101 surrounding individual field work sites, (2.5 km radius; Fig. 1a).

102 For a detailed description of field and laboratory work please refer to SI. For eDNA
103 collection and BRUV filming we visited three locations in southern Te Wahipounamu
104 (Moana Uta / Wet Jacket Arm, Taumoana / Five Fingers, and Te Tapuwae a Hua / Long
105 Sound; henceforth WJ MR, FF MR, and LS MR), and accompanying control areas outside
106 those MRs (henceforth WJ CTRL, FF CTRL, and LS CTRL), from 12.–22. December 2019
107 (Fig. 1a.) Within each sampling location, at randomised sites, we collected eDNA (mean
108 depth 14.05 m, med.: 15, sd.: 1.4 m), and subsequently deployed BRUV assemblies (mean
109 depth 15.6 m, med.: 16, sd.: 2.6 m; SI). We considered data from 21 sites (FF: 2 FF MR and
110 3 FF CTRL, WJ: 4 WJ MR and 4 WJ CTRL, and LS: 4 LS MR and 4 LS CTRL). We
111 collected two 900 ml water samples with eDNA at each site, filtered them alongside negative
112 controls, then sealed and stored them until further processing. BRUV footage was obtained
113 for one hour and analysed by eye.

114 Environmental DNA was isolated in a PCR-free facility alongside extraction and cross-
115 contamination controls (SI: four species of tropical freshwater fish). After *in silico* PCR (SI),
116 we amplified our extracts with the well-established and widely used 12S MiFish primers
117 (“MiFish-U”; Miya et al. 2015; see SI Table 1 for primer comparison), targeting
118 Actinopterygii. Chondrichthyes were targeted with slightly altered derivatives (“Elas02”,
119 Taberlet et al. 2018). Our single-step PCRs were cycled 45 times, with annealing
120 temperatures of 45 °C (MiFish-U) or 40 °C (Elas02). Amplified eDNA was then pooled,
121 visualised, purified, combined equimolarly, diluted to 4.5 pmol, and sequenced on an
122 Illumina MiSeq (Illumina, San Diego, US-CA; kit v2, 300 cycles, single-ended).

123 We defined Amplicon Sequence Variants (ASVs; Callahan et al. 2017) from eDNA after
124 demultiplexing with Cutadapt v3.0 (Martin, 2011), using Qiime2 2020-08 (Bolyen et al.,
125 2019) and DADA2 1.10.0 (Callahan et al., 2016). To yield high quality sequence data we did
126 not allow any mismatches, nor Expected Errors (Edgar & Flyvbjerg, 2015) during
127 demultiplexing. Taxonomic annotation of denoised data was obtained using Blast 2.10.0+
128 (Camacho et al., 2009) and a local full copy of the NCBI nucleotide collection (April 2020;
129 Benson et al., 2011) while excluding environmental samples. To yield a maximum of
130 taxonomically annotated ASVs, we chose relaxed taxonomic assignment parameters in
131 combination with an e-value to retain only the most significant alignments. We required a
132 minimum identity of 75% among all alignments and kept five high-scoring pairs for each
133 eDNA query, each of which needed a minimum coverage of 95% to be retained. The minimal
134 acceptable e-value was set to 10^{-10} . We retained the best high-scoring alignment of each
135 query-reference pair based on the highest bit score. We removed data contained in negative
136 controls, alongside ASVs covered by fewer than 15 reads (see SI).

137 To investigate how well the literature- and OBIS-derived biodiversity information were
138 resolved by eDNA and BRUV, we checked the concordance of all data sources on order,
139 family, genus and on species levels. To judge sampling effort and total species diversity
140 based on BRUV and eDNA observations, we inspected species accumulation curves and
141 calculated Good Turing estimators (giving the number of all species based on species already
142 seen in a small sample; Good, 1953), and then compared those values to combined Te
143 Wahipounamu literature and OBIS species records.

144 To verify the credibility of eDNA information, we checked all eDNA species lists against a
145 comprehensive list of all New Zealand fish (Roberts et al., 2019) and evaluated species
146 assignment and alignment qualities.

147 To investigate how useful BRUV, eDNA and OBIS records are in detecting regional
148 differences between fish biodiversity, we used Analysis of Similarity (ANOSIM; Clarke
149 1993). Thereby analysing Jaccard distances (Jaccard, 1912), we looked for significant
150 differences in taxon (species, genus, family, order) overlap depending on various factor
151 combinations, hence checking whether a particular observation method fared better in
152 detecting taxon composition differences either between different field work areas (WJ, FF,
153 LS), or according to protection status (MR or CNTRL).

154 Results

155 Each of our four data sources yielded different species counts. In total, we yielded 116
156 species (106 Actinopterygii, 10 Chondrichthyes), comprised of 59 species from previously
157 published Te Wahipounamu works, 44 unique species derived from eDNA, 25 from BRUV
158 and 25 from OBIS (large area; see Table 1, Fig. 2, SI. Table 4). While 21 field work sites
159 (Fig. 1a) yielded environmental DNA and BRUV data (Fig. 1b, c) matching local OBIS data
160 could only be obtained for nine field work sites (Fig. 1a, small circles, LS CNTRL, FF, WJ),
161 and hence those finer scaled OBIS data were later excluded from ANOSIM as incomplete
162 data (Fig. 1a, small circles, Fig. 1d). For further comparisons with Good-Turing estimates, we
163 posit the local “real” species count to 68 as the number of unique species observed across
164 literature and OBIS (Fig. 3).

165 Obtaining community composition comparable to literature and OBIS data within our works’
166 spatial constraints worked better with BRUV than with eDNA. Nineteen out of 25 species
167 detected with BRUV (76%) were contained in the literature or on OBIS, but only one out of
168 44 species detected with eDNA (2%) were contained in Te Wahipounamu-specific literature
169 or OBIS (Fig. 1a, large circle). Concordance of taxonomic information between the four data
170 sources dissolved with lowering taxonomic levels, and most decisively for eDNA data (Fig.
171 3). At species level, only two taxonomic assignments from eDNA matched other data
172 sources, namely *Notorynchus cepedianus* (broadnose sevengill shark), also found with
173 BRUV, and *Aldrichetta forsteri* (yellow-eye mullet) also listed in the literature (Fig. 2).
174 BRUV agreed better with available local biodiversity information, with 11 species mentioned
175 both in the literature and OBIS, and eight detected species mentioned in the literature only
176 (Fig. 3). On BRUV we identified six species (*Bodianus unimaculatus*, *Chelidonichthys kumu*,
177 *Galeorhinus galeus*, *Mustelus lenticulatus*, *Notorynchus cepedianus*, *Scorpaena cardinalis*)
178 not mentioned in literature, and not in OBIS, but occurring in New Zealand waters (Roberts
179 et al., 2019). While our plateauing species accumulation curves suggested exhaustive
180 sampling (SI Fig. 5), Good-Turing estimates of eDNA data inferred a presence of 60 species
181 in the study area (assuming 27% missed after 44 observations), and a presence of 26 species
182 using BRUV (assuming 7% missed after 25 observations).

183 Alignment qualities associated with taxonomic annotation of eDNA data were variable.
184 Forty-four species assigned among eDNA were defined by 92 ASVs (across 142
185 observations) of which only six yielded flawless alignments with reference data (i.e. 14%,
186 with full query coverage, no alignment gaps). Eighty-six ASVs had variable query coverage

187 (37 families, Tab. 1 and SI), while 32 ASVs had variable gap counts (15 families, Tab. 1 and
188 SI). Mean query coverage was 93.2% (min: 78.6%, med: 97%, sd 6.5%), mean gap count was
189 1 (max: 10, med: 0, sd 1.83; Tab. 1). Nineteen species assignments among eDNA (43%) had
190 not been observed in New Zealand, and none of these species were found using BRUV,
191 across literature, or OBIS data (apart from *Bovichtus variegatus* – thornfish, not in Roberts et
192 al., 2019, but in Roberts, 2005; Fig. 2). Importantly, 25 species observed with eDNA (56.8%
193 of eDNA-observed species) were known from somewhere New Zealand (Roberts et al., 2019)
194 but were not observed in BRUV or found in Te Wahipounamu literature. Interestingly, using
195 eDNA, we obtained perfect alignments between few ASVs and reference data for
196 *Arctocephalus forsteri* (New Zealand fur seal), *Balaenoptera musculus* (blue whale), and
197 *Tursiops truncatus* (bottlenose dolphin).

198 In ANOSIM, only BRUV data, and not eDNA data, exhibited location-specific differences
199 among species' presence overlaps among the 21 sites – on species, genus, family, and order
200 levels. Significant differences were calculated in overlaps between the six field work areas
201 but not between marine reserve nor control areas (SI Table 3).

202 Investigation of the strikingly homogenous structure of eDNA data by regression analysis of
203 the 142 non-unique eDNA observations (Tjur's R^2 0.027) suggested each additional
204 alignment gap to be associated with a 39% increased probability of observing a non-native
205 species (Odds Ratio 1.39, 95% CI from 1.19 to 1.66, $p < 0.01$). A 1% increase in alignment
206 concordance was associated with a 7% increased probability of non-native observation (OR
207 1.07, 95% CI 1.03–1.12, $p < 0.01$). Null deviance was 572.60 on 141 degrees of freedom,
208 residual deviance was 552.31 on 139 degrees of freedom (SI Figs 7 and 8).

209 **Discussion**

210 What is a realistic estimate of the fish biodiversity in Te Wahipounamu? Based on literature
211 and OBIS alone, we estimate the currently described combined ray-finned and cartilaginous
212 fish species count of Te Wahipounamu to be 68, minding that we constrained OBIS data to
213 surround field sites (Fig. 1, large circle), and that those data are predominantly based on
214 visual observations (SI Table 2). If species counts obtained from literature and OBIS were
215 close to a real value of 68, and the same was true for eDNA and BRUV data, both respective
216 Good-Turing estimates would be 68. Our BRUV-based Good-Turing estimate of 26 species
217 diverges strongly from this number. This may have several reasons. Firstly, we only
218 inspected an isolated area in Te Wahipounamu, while the literature describes a larger area.

219 Secondly, bait in BRUV does not attract all fish for the camera, particularly if deployed at
220 limited depth range, as done here. For eDNA, the Good-Turing estimate of 60 species is more
221 like the literature-inferred species count, but this could be coincidental.

222 How credible are eDNA derived species assignments with currently available reference data?
223 We believe lacking eDNA reference data to restrict accurate species annotation of ASVs.
224 There are several observations from our data that appear to support this hypothesis. First,
225 while there is a reasonably good concordance between species identified in our BRUV
226 analyses and species known from the area as combined from publications and OBIS, the
227 dissimilarity between eDNA data on one side, and BRUV, OBIS and publication data on the
228 other side, increases with decreasing taxonomic level, culminating in only two out of 44
229 eDNA species being either identified in our BRUV analyses or known from previous
230 publications (Fig. 3).

231 Secondly, every approach to identify species diversity in a marine ecosystem has its biases,
232 and published observations are mostly based on visual approaches. Thus, one could argue for
233 the existence of a bias favouring similarity between our visual BRUV observations and
234 published species occurrences to the detriment of eDNA data's similarity. However, we do
235 not believe this circumstance alone to be responsible for a bias favouring BRUV data to be
236 more similar with literature and OBIS observations in comparison to eDNA observations.
237 Literature and OBIS observation methodologies extend well beyond the specific biases of
238 BRUV, including a multitude of different observation techniques (poison stations, seine net
239 fishing, spear fishing, diver surveys and others, SI Table 1). Collectively, all observation
240 techniques should have provided an appropriately comprehensive overview of fish diversity
241 in Te Wahipounamu, lacking biases inherent to BRUV.

242 Thirdly, some divergence between eDNA data and the other data sources may be explained
243 by the known ability of eDNA to detect “cryptic” species that are not easily discovered by
244 any visual surveying. The most obvious candidates for this category would be the 25 eDNA
245 species that had previously been reported from New Zealand but not yet from Te
246 Wahipounamu (Fig. 2). However, such a bias should not prevent a broad overlap between
247 eDNA and visual approaches for species that can easily be detected visually. Clearly, we did
248 not find such an overlap.

249 Crucially, of the 25 species we detected by BRUV and that were therefore present at the time
250 of our concurrent water sampling for eDNA analyses, 24 species are not present in the NCBI

251 reference database (Fig. 2) and could therefore not be detected by our eDNA approach. This
252 highlights one of the main limitations of eDNA multispecies surveys today.

253 Nevertheless, and despite the lack of reference data, eDNA still identified a larger number of
254 species than our concurrent BRUV analyses. From where do these species assignments
255 come? In most cases during taxonomic assignment, where no perfect match can be found
256 between eDNA query and reference subject sequence, the algorithm assigning ASVs to
257 species information (BLAST) chose the next-closest matching species contained in the
258 reference data collection, as encouraged by our taxonomic assignment parameters. Our
259 taxonomic assignments correspond with this hypothesis, as binomial regression showed that
260 each additional gap in a sequences' reference alignment associated with a 39% increased
261 probability of observing of a non-native species.

262 Interestingly, a 1% increase in alignment concordance increased the likelihood of a non-
263 native observation as well, by 7%. At first sight this seems counter-intuitive, however the
264 latter observation is also consistent with our hypothesis: A poorly matching sequence would
265 not be assigned to a matching species but rather to a higher matching taxon such as genus or
266 family. A better fit increases the likelihood of a species level assignment, but without native
267 species contained among reference data, the likelihood increases that the query sequence is
268 assigned to a closely related species not occurring in New Zealand. Similar observations have
269 been made in other regions of the world (Stoeckle, Das Mishu, & Charlop-Powers, 2020).

270 The large number of species detected by our eDNA approach – although probably
271 misassigned in several instances – is a testament to the potential power of eDNA methods.
272 Arguably, any detected effect of lacking reference data could be less pronounced by using
273 another, or multiple primer pairs. For example, our primer evaluations with the recently
274 released software GAPeDNA (Marques et al., 2021) show that the “Fish 16S” primer set by
275 McInnes et al. (2017) would have covered 249 instead of the 119 New Zealand marine fish
276 species covered by our MiFish 12S dataset (SI Table 1). However, the overall conclusion
277 remains. Of the over 1294 known New Zealand marine fish species, molecular reference data
278 of any kind is available only for 489 species in southern New Zealand, and for no available
279 primer pairs sufficient reference data is available. Hence without substantial effort into
280 generating suitable reference data for a carefully selected range of similar primers, eDNA
281 analysis here and everywhere else will remain an impaired tool for biodiversity management.
282 While this insight holds true after almost two decades of eDNA research (Hebert, Cywinska,
283 Ball, & DeWaard, 2003) we note that a growing number of researchers are working hard on

284 closing reference data voids around the globe (reviewed in Marques et al., 2021; Weigand et
285 al., 2019).

286 **Acknowledgments and Data**

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290 provided Good-Turing estimates, accumulation curves and advised P.C. M.H., M.L., P.C.
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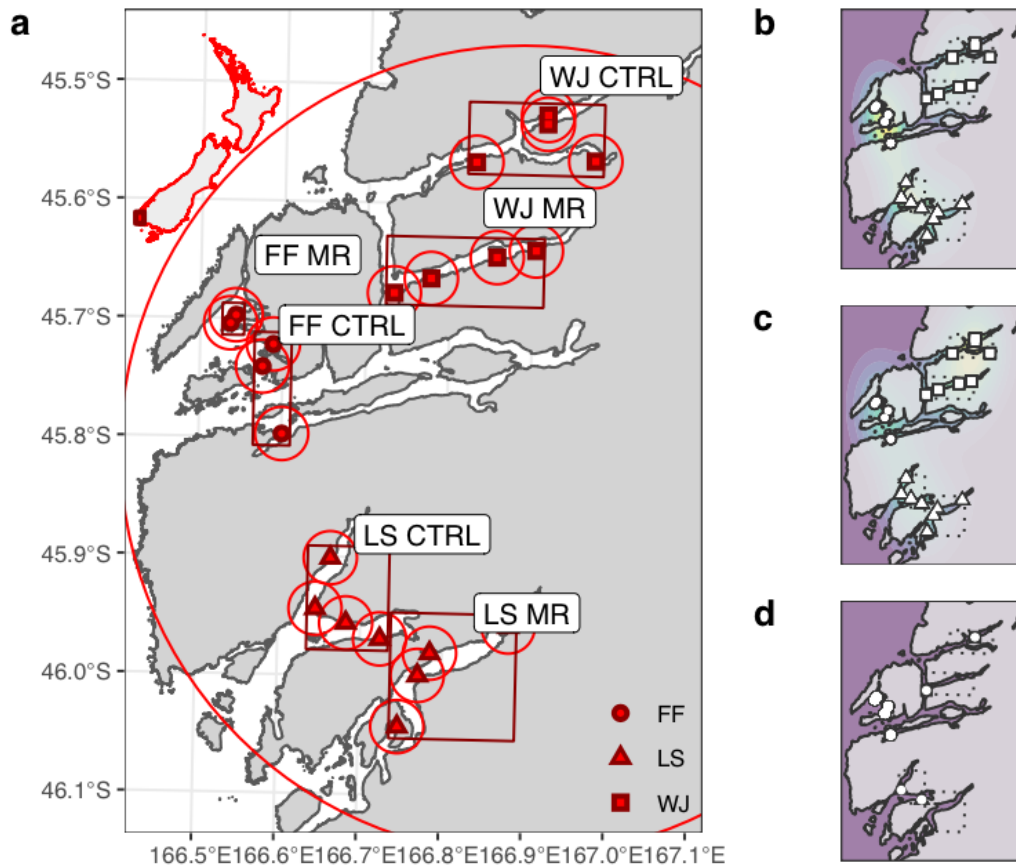
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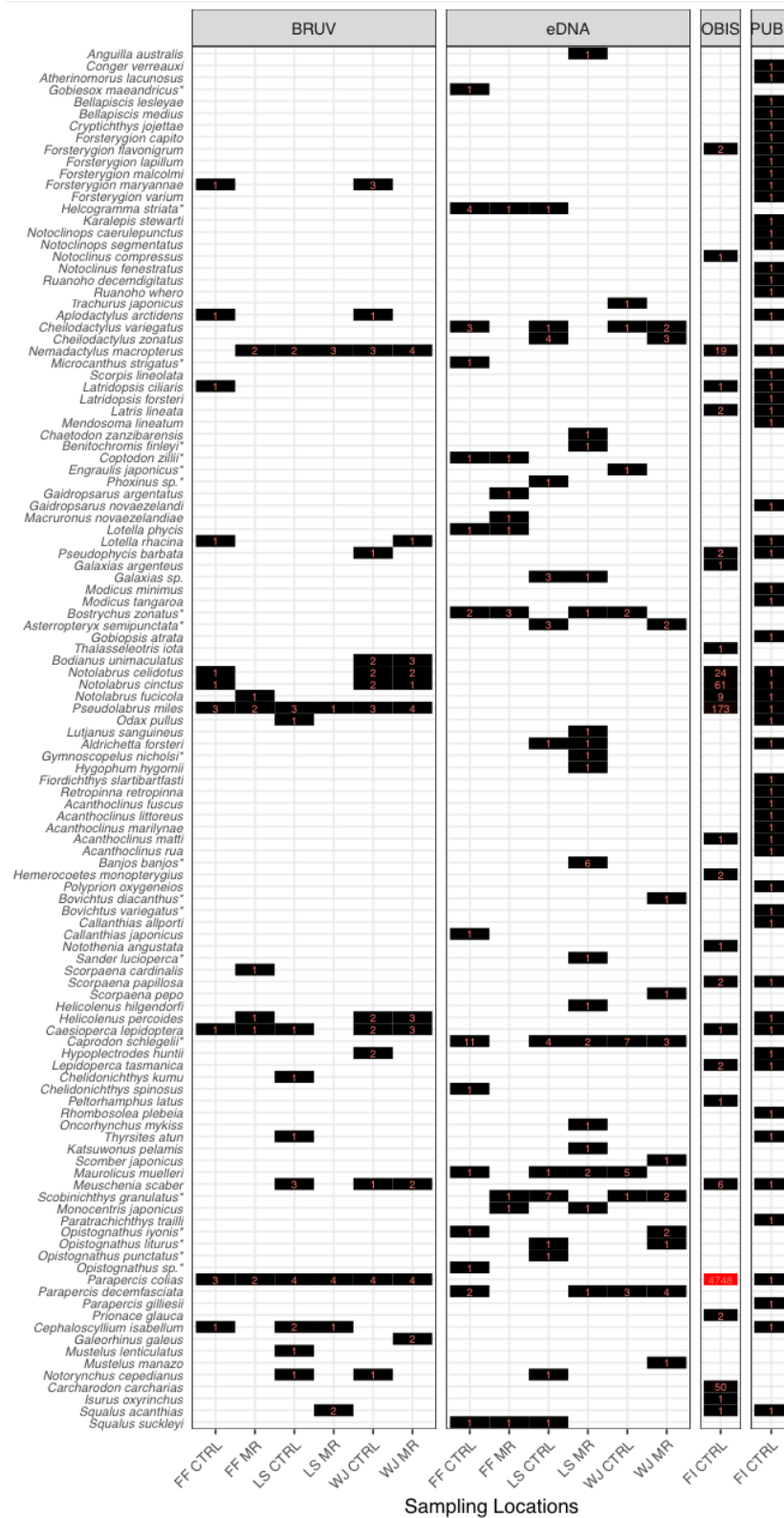
385 **Figures and Tables**



386

387 **Fig. 1:** Field work area, description, sites, and data coverage for eDNA, BRUV, and
388 OBIS data. **a:** We obtained biodiversity information from baited remote underwater video
389 (BRUV) footage and environmental DNA (eDNA) data from 21 field work sites across
390 three sampling regions (highlighted by rectangles) – Five Fingers (FF), Long Sound (LS)
391 and Wet Jacket (WJ). In each region we collected samples inside marine reserves /
392 commercial exclusion zones (MR) and outside in control areas (CTRL). To obtain
393 additional biodiversity information, we queried the Ocean Biodiversity Information
394 System (OBIS – <https://obis.org/>) for records within a 2.5 km radius of each field work
395 site (small circles) for the purpose of community structure analysis. Furthermore, we
396 obtained OBIS records for the entire sampling region (large circle) to extend our species
397 list alongside species mentioned across various literature sources (Table 1, SI Table 3). **b:**
398 Environmental DNA (eDNA), and **c:** BRUV data in a spatial context, lighter colour
399 indicates a higher density of distinct species observations (corresponding to numerical
400 values in Fig. 2). **d:** Species data for all field work sites could not be obtained from OBIS,

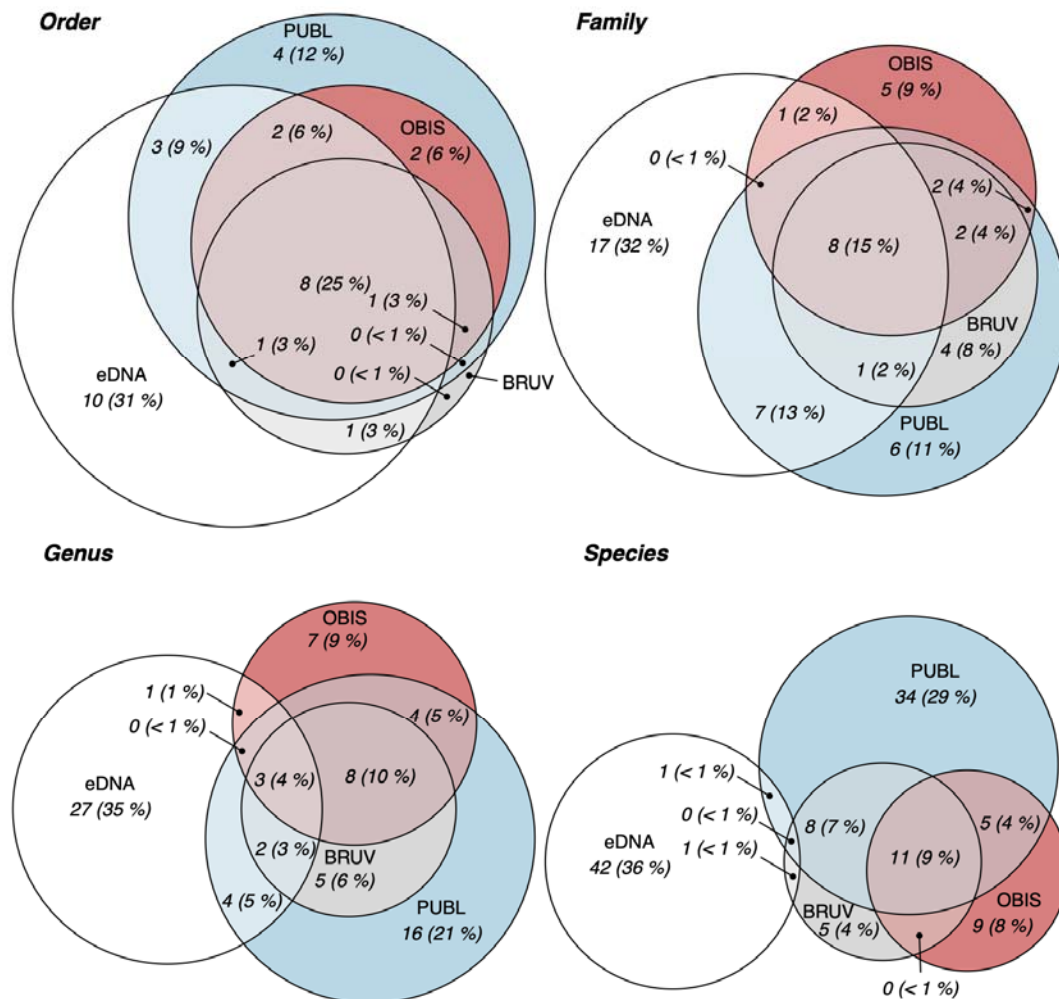
401 necessitating the exclusion of this data in the statistical analyses of regional biodiversity
 402 data. Graph created using R package *ggplot2* (3.3.5).



403

Sampling Locations

404 **Fig. 2:** Distinct species observations across data sources and field work locations.
 405 Observation types: **BRUV** – Observations from baited remote underwater surveys; **eDNA**
 406 – environmental DNA observations; **OBIS** – data retrieved from the Ocean Biodiversity
 407 Information System (<https://obis.org/>) for the area surrounding field work sites (large
 408 circle in Fig.1); **PUBL** – Fiordland fish species collated from multiple literature records
 409 as summarized by Inglis (2008). Sampling Locations: FF – Five Fingers area; LS – Long
 410 Sound area; WJ – Wet Jacket area; MR – marine reserve or commercial exclusion zone;
 411 CTRL – neither marine reserve nor commercial exclusion zone. Species list: Order
 412 follows Table 1, species not listed as New Zealand Species in (CD Roberts et al., 2019)
 413 are marked with an asterisk (*). Graph created using R package *ggplot2* (3.3.5).



414
 415 **Fig. 3:** Concordance of taxonomic information across four data sources of Fiordland fish
 416 biodiversity. Biodiversity data (Table 1, SI Table 2) is summarized at four different

417 taxonomic levels, shown are unique observation counts at each level, as well as the
418 corresponding percentage of those counts in comparison to all data. Circle sizes
419 proportional to observation count. Observation types: BRUV (grey) – Observations from
420 baited remote underwater surveys; eDNA (white) – environmental DNA observations;
421 OBIS (red) – data retrieved from the Ocean Biodiversity Information System
422 (<https://obis.org/>) for the area surrounding field work sites (large circle in Fig.1); PUBL
423 (blue) – Fiordland fish species collated from multiple literature records as summarized by
424 (Inglis et al., 2008). Graph created using R package *eulerr* (6.1.0).

425 **Table 1:** Details on taxonomic observations across data sources. Taxonomic hierarchies conform with NCBI taxonomy where available, thus
426 allow analysis in relation to environmental DNA (eDNA) data and are sorted alphabetically – the resulting species order is identical to Fig. 2.
427 Taxa not listed as New Zealand species by Roberts et al., (2019) are highlighted with asterisk (*). Trivial names are indicated where
428 available from NCBI. For all taxonomic assignments also yield from eDNA we provide the alignment coverage and alignment gaps. Since
429 identical species were assigned to multiple Amplicon Sequence Variants (ASV's; Callahan et al., 2017) in some instances, ranges are
430 provided for alignment coverages and gap counts for species-specific alignments.

431

Phylum	Class	Order	Family	Genus	Species	Common name	Algn. covrg.	Algn. gaps
Chordata	Actinopteri	Anguilliformes	Anguillidae	<i>Anguilla</i>	<i>Anguilla australis</i>	Australian shortfin eel	100%	0
			Congridae	<i>Conger</i>	<i>Conger verreauxi</i>	conger eel		
		Atheriniformes	Atherinidae	<i>Atherinomorus</i>	<i>Atherinomorus lacunosus</i>	hardyhead silverside		
		Blenniiformes	Gobiesocidae	<i>Gobiesox</i> *	<i>Gobiesox maeandricus</i> *	northern clingfish	79.3%	2
				Tripterygiidae	<i>Bellapiscis</i>	<i>Bellapiscis lesleyae</i>	mottled twister	
			<i>Bellapiscis medius</i>			twister		
			<i>Cryptichthys</i>		<i>Cryptichthys jojettae</i>			
			<i>Forsterygion</i>		<i>Forsterygion capito</i>		spotted robust triplefin	
				<i>Forsterygion flavonigrum</i>		yellow-and-black triplefin		
		<i>Forsterygion lapillum</i>			common triplefin			
			<i>Forsterygion malcolmi</i>					

			<i>Forsterygion maryannae</i>			
			<i>Forsterygion varium</i>	striped triplefin		
		<i>Helcogramma*</i>	<i>Helcogramma striata*</i>		88.2- 92.3%	1
		<i>Karalepis</i>	<i>Karalepis stewarti</i>			
		<i>Notoclinops</i>	<i>Notoclinops caerulepunctus</i>			
			<i>Notoclinops segmentatus</i>			
		<i>Notoclinus</i>	<i>Notoclinus compressus</i>			
			<i>Notoclinus fenestratus</i>			
		<i>Ruanoho</i>	<i>Ruanoho decemdigitatus</i>			
			<i>Ruanoho whero</i>	spectacled triplefin		
Carangiformes	Carangidae	<i>Trachurus</i>	<i>Trachurus japonicus</i>	Japanese jack mackerel	99.4%	0
Centrarchiformes	Aplodactylidae	<i>Aplodactylus</i>	<i>Aplodactylus arctidens</i>			

	Cheilodactylidae	<i>Cheilodactylus</i>	<i>Cheilodactylus variegatus</i>		97.6-98.8%	0
			<i>Cheilodactylus zonatus</i>	blackbarred morwong	97-97.6%	0
		<i>Nemadactylus</i>	<i>Nemadactylus macropterus</i>	tarakihi		
	Kyphosidae	<i>Microcanthus*</i>	<i>Microcanthus strigatus*</i>	stripey	85.3%	3
		<i>Scorpis</i>	<i>Scorpis lineolata</i>	silver sweep		
	Latridae	<i>Latridopsis</i>	<i>Latridopsis ciliaris</i>	blue moki		
			<i>Latridopsis forsteri</i>	bastard trumpeter		
		<i>Latris</i>	<i>Latris lineata</i>	striped trumpeter		
		<i>Mendosoma</i>	<i>Mendosoma lineatum</i>			
Chaetodontiformes	Chaetodontidae	<i>Chaetodon</i>	<i>Chaetodon zanzibarensis</i>		81.5%	4
Cichliformes	Cichlidae	<i>Benitochromis*</i>	<i>Benitochromis finleyi*</i>		79.4%	0
		<i>Coptodon*</i>	<i>Coptodon zillii*</i>	redbelly tilapia	90.6%	3

Clupeiformes	Engraulidae	<i>Engraulis*</i>	<i>Engraulis japonicus*</i>	Japanese anchovy	98.8%	0
Cypriniformes	Cyprinidae	<i>Phoxinus*</i>	<i>Phoxinus sp.*</i>		97.1%	0
Gadiformes	Gaidropsaridae	<i>Gaidropsarus</i>	<i>Gaidropsarus argentatus</i>	Arctic rockling	90.1%	1
			<i>Gaidropsarus novaezealandi</i>			
	Merlucciidae	<i>Macruronus</i>	<i>Macruronus novaezealandiae</i>	blue grenadier	100%	0
Moridae		<i>Lotella</i>	<i>Lotella phycis</i>		95.9%	0
			<i>Lotella rhacina</i>	rock cod		
			<i>Pseudophycis</i>	<i>Pseudophycis barbata</i>	southern bastard codling	
Galaxiiformes	Galaxiidae	<i>Galaxias</i>	<i>Galaxias argenteus</i>			
			<i>Galaxias sp.</i>		96.5%	0
Gobiesociformes	Gobiesocidae	<i>Modicus</i>	<i>Modicus minimus</i>			
Gobiiiformes	Eleotridae	<i>Bostrychus*</i>	<i>Modicus tangaroa</i>			
			<i>Bostrychus zonatus*</i>	barred gudgeon	85.4-86%	5

	Gobiidae	<i>Asterropteryx*</i>	<i>Asterropteryx semipunctata*</i>	starry goby	81.9-82.5%	6
		<i>Gobiopsis</i>	<i>Gobiopsis atrata</i>			
	Thalasseleotrididae	<i>Thalasseleotris</i>	<i>Thalasseleotris iota</i>			
Labriformes	Labridae	<i>Bodianus</i>	<i>Bodianus unimaculatus</i>	red pigfish		
		<i>Notolabrus</i>	<i>Notolabrus celidotus</i>	New Zealand spotty		
			<i>Notolabrus cinctus</i>			
			<i>Notolabrus fucicola</i>	yellow-saddled wrasse		
		<i>Pseudolabrus</i>	<i>Pseudolabrus miles</i>			
	Odacidae	<i>Odax</i>	<i>Odax pullus</i>	greenbone		
Lutjaniformes	Lutjanidae	<i>Lutjanus</i>	<i>Lutjanus sanguineus</i>	humphead snapper	80.8%	6
Mugiliformes	Mugilidae	<i>Aldrichetta</i>	<i>Aldrichetta forsteri</i>	yellow-eye mullet	96-100%	0
Myctophiformes	Myctophidae	<i>Gymnoscopelus*</i>	<i>Gymnoscopelus nicholsi*</i>		79.8%	1

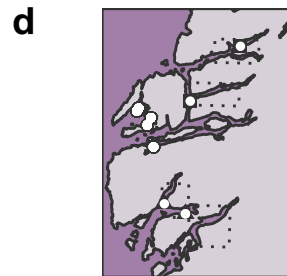
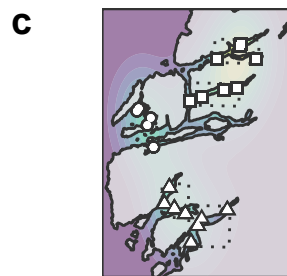
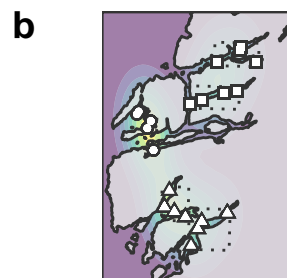
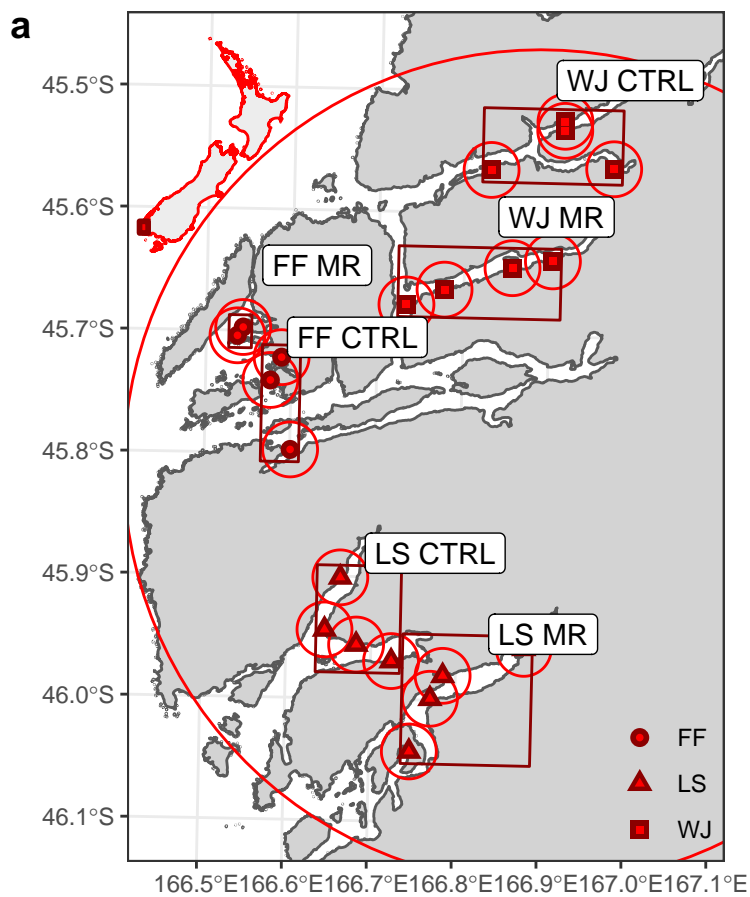
		<i>Hygophum</i>	<i>Hygophum hygomii</i>	78.7%	3
Ophidiiformes	Bythitidae	<i>Fiordichthys</i>	<i>Fiordichthys slartibartfasti</i>		
Osmeriformes	Retropinnidae	<i>Retropinna</i>	<i>Retropinna retropinna</i>	cucumberfish	
Ovalentaria	Plesiopidae	<i>Acanthoclinus</i>	<i>Acanthoclinus fuscus</i>		
			<i>Acanthoclinus littoreus</i>		
			<i>Acanthoclinus marilynae</i>		
			<i>Acanthoclinus matti</i>		
			<i>Acanthoclinus rua</i>		
Pempheriformes	Banjosidae	<i>Banjos*</i>	<i>Banjos banjos*</i>	84.7- 85.2%	0
	Percophidae	<i>Hemerocoetes</i>	<i>Hemerocoetes monopterygius</i>		
	Polyprionidae	<i>Polyprion</i>	<i>Polyprion oxygeneios</i>		
Perciformes	Bovichtidae	<i>Bovichtus*</i>	<i>Bovichtus diacanthus*</i>	94.1%	0

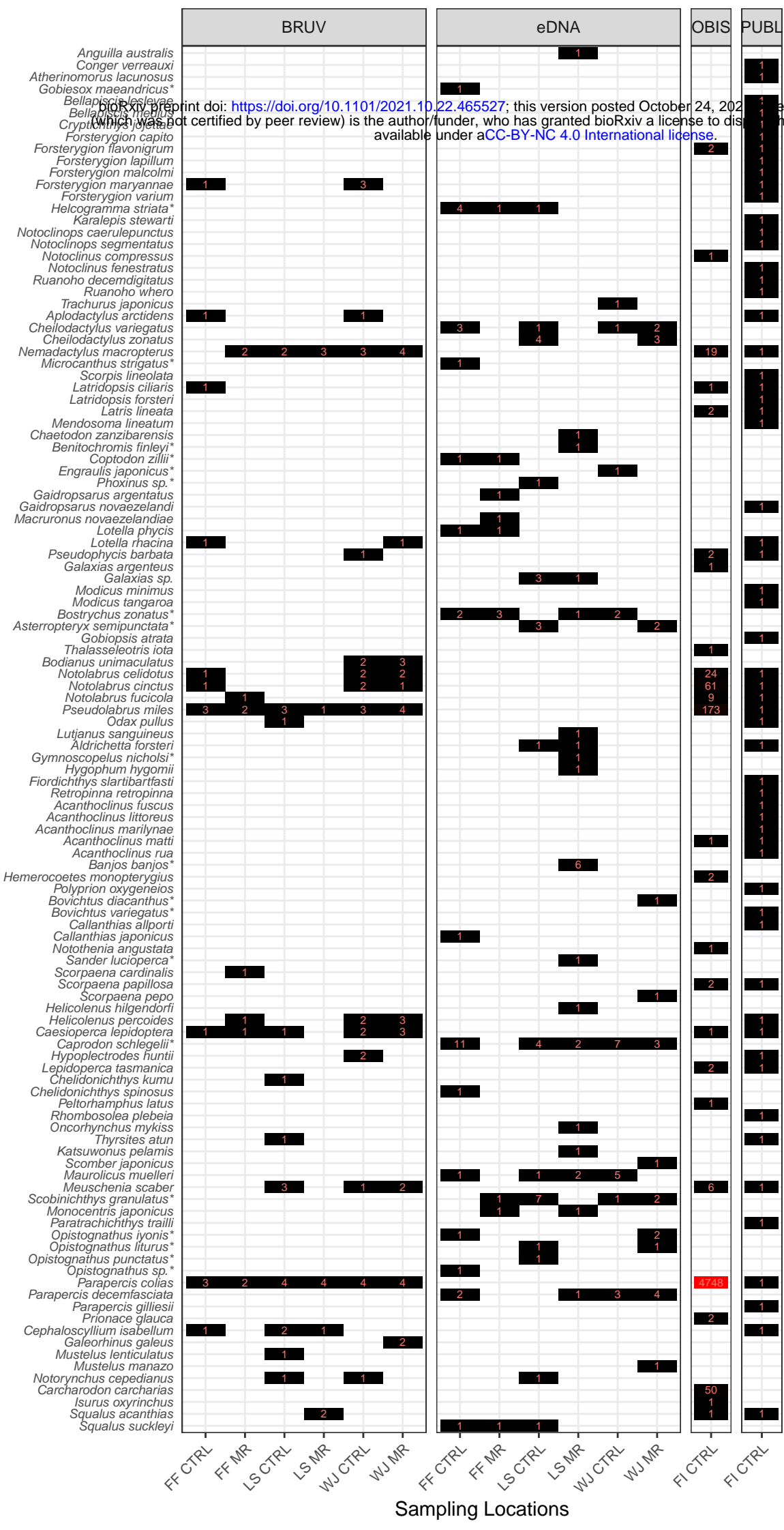
		<i>Bovichtus variegatus*</i>	thornfish		
Callanthiidae	<i>Callanthias</i>	<i>Callanthias allporti</i>			
		<i>Callanthias japonicus</i>		95.2%	1
Nototheniidae	<i>Notothenia</i>	<i>Notothenia angustata</i>	Maori chief		
Percidae	<i>Sander*</i>	<i>Sander lucioperca*</i>	pikeperch	81.4%	3
Scorpaenidae	<i>Scorpaena</i>	<i>Scorpaena cardinalis</i>	red rock cod		
		<i>Scorpaena papillosa</i>			
		<i>Scorpaena pepo</i>	pumpkin scorpionfish	85.7%	0
Sebastidae	<i>Helicolenus</i>	<i>Helicolenus hilgendorfi</i>		95.4%	0
		<i>Helicolenus percoides</i>			
Serranidae	<i>Caesioperca</i>	<i>Caesioperca lepidoptera</i>			
	<i>Caprodon*</i>	<i>Caprodon schlegelii*</i>	sunrise perch	90.6-98.2%	0-2

		<i>Hypoplectrodes</i>	<i>Hypoplectrodes huntii</i>			
		<i>Lepidoperca</i>	<i>Lepidoperca tasmanica</i>			
	Triglidae	<i>Chelidonichthys</i>	<i>Chelidonichthys kumu</i>	bluefin gurnard		
			<i>Chelidonichthys spinosus</i>	red gurnard	99.4%	0
Pleuronectiformes	Rhombosoleidae	<i>Peltorhamphus</i>	<i>Peltorhamphus latus</i>	speckled sole		
		<i>Rhombosolea</i>	<i>Rhombosolea plebeia</i>	New Zealand flounder		
Salmoniformes	Salmonidae	<i>Oncorhynchus</i>	<i>Oncorhynchus mykiss</i>	rainbow trout	100%	0
Scombriformes	Gempylidae	<i>Thyrsites</i>	<i>Thyrsites atun</i>	snoek		
	Scombridae	<i>Katsuwonus</i>	<i>Katsuwonus pelamis</i>	skipjack tuna	95.9%	0
		<i>Scomber</i>	<i>Scomber japonicus</i>	chub mackerel	100%	0
Stomiiformes	Sternoptychidae	<i>Maurolicus</i>	<i>Maurolicus muelleri</i>	pearlsides	86.3-99.4%	0-10
Tetraodontiformes	Monacanthidae	<i>Meuschenia</i>	<i>Meuschenia scaber</i>	velvet leatherjacket		

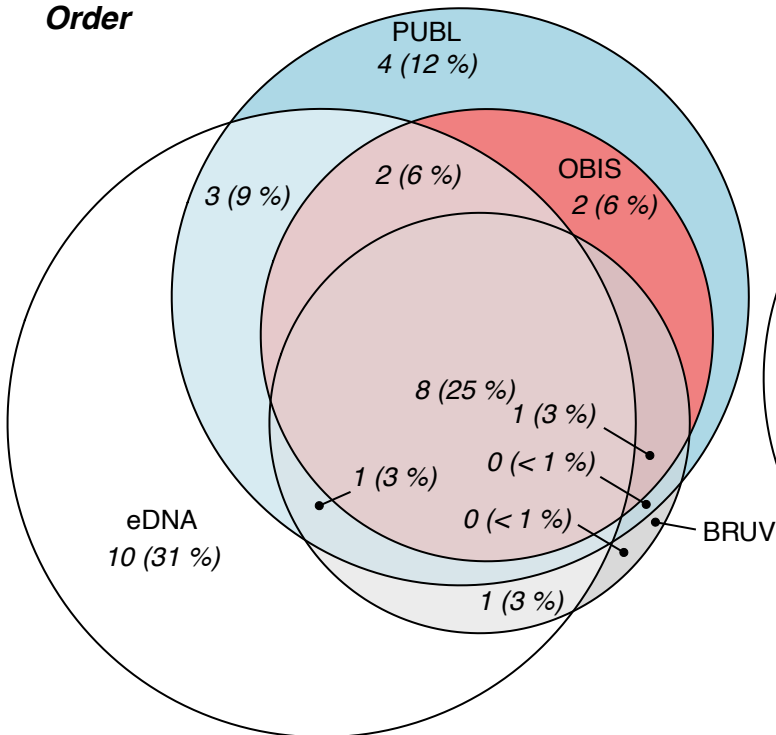
			<i>Scobinichthys</i> *	<i>Scobinichthys granulatus</i> *	rough leatherjacket	98.8-99.4%	0
	Trachichthyiformes	Monocentridae	<i>Monocentris</i>	<i>Monocentris japonicus</i>		94.4-97.6%	2-3
		Trachichthyidae	<i>Paratrachichthys</i>	<i>Paratrachichthys trailli</i>	sandpaper fish		
	undefined	Opistognathidae	<i>Opistognathus</i> *	<i>Opistognathus iyonis</i> *		89.9-90.5%	2-3
				<i>Opistognathus liturus</i> *	seto-amadai	89.3-90.5%	2
				<i>Opistognathus punctatus</i> *	finespotted jawfish	81.7%	5
				<i>Opistognathus sp.*</i>		85.7%	2
	Uranoscopiformes	Pinguipedidae	<i>Parapercis</i>	<i>Parapercis colias</i>	New Zealand blue cod		
				<i>Parapercis decemfasciata</i>		80.9%	1
				<i>Parapercis gilliesii</i>	yellow weaver		
Chondrichthyes	Carcharhiniformes	Carcharhinidae	<i>Prionace</i>	<i>Prionace glauca</i>	blue shark		
		Scyliorhinidae	<i>Cephaloscyllium</i>	<i>Cephaloscyllium isabellum</i>			

	Triakidae	<i>Galeorhinus</i>	<i>Galeorhinus galeus</i>	tope shark		
		<i>Mustelus</i>	<i>Mustelus lenticulatus</i>	spotted estuary smooth-hound		
			<i>Mustelus manazo</i>	starspotted smooth-hound	98.9%	0
Hexanchiformes	Hexanchidae	<i>Notorynchus</i>	<i>Notorynchus cepedianus</i>	broadnose sevengill shark	98.4%	0
Lamniformes	Alopiidae	<i>Carcharodon</i>	<i>Carcharodon carcharias</i>	great white shark		
		<i>Isurus</i>	<i>Isurus oxyrinchus</i>	shortfin mako shark		
Squaliformes	Squalidae	<i>Squalus</i>	<i>Squalus acanthias</i>	spiny dogfish		
			<i>Squalus suckleyi</i>	Puget Sound dogfish	100%	0

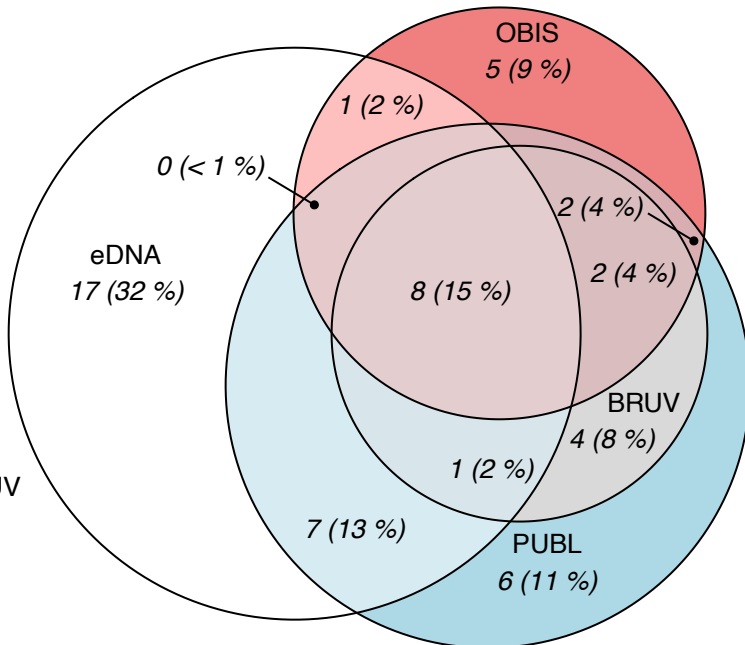




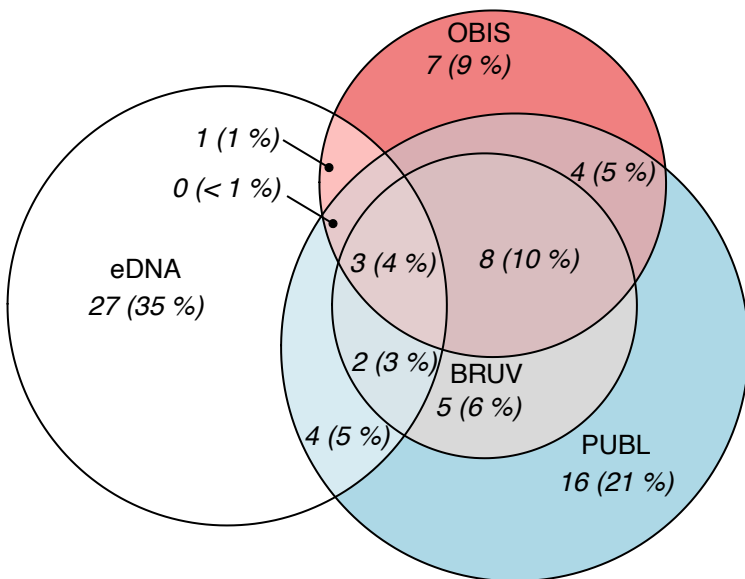
Order



Family



Genus



Species

