Quantifying Impacts of an Environmental Intervention Using Environmental

2 **DNA**

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19 Abstract

Environmental laws around the world require some version of an environmental impact assessment surrounding construction projects and other discrete instances of human development. Information requirements for these assessments vary by jurisdiction, but nearly all require an analysis of the biological elements of affected ecosystems. Amplicon-sequencing - also called metabarcoding - of environmental DNA (eDNA) has made it possible to sample and amplify the genetic material of many species present in those environments, 24 providing a tractable, powerful, and increasingly common way of doing environmental impact analysis for 25 development projects. Here, we analyze a 12-month 18-month time-series of water samples taken before, during, and after a culvert removal project two culvert removals in a salmonid-bearing freshwater stream. We 27 use an asymmetrical Before-After-Control-Intervention (BACI) design with also sampled multiple control streams to develop a robust background expectation against which to evaluate the impact of this discrete 29 environmental intervention in the treatment stream. We generate calibrated, quantitative metabarcoding data from amplifying the 12s MiFish mtDNA locus and complementary species-specific quantitative PCR 31 data to yield multi-species estimates of absolute eDNA concentrations across time, creeks, and sampling stations. We then use a hierarchical Bayesian time-series-linear mixed-effects model to reveal patterns of 33 eDNA concentrations over time, and to estimate the effects of the culvert removal on salmonids in the treatment creek. We focus our analysis on four common salmonid species: cutthroat trout (Oncorhynchus clarkii), coho salmon (O. kisutch), rainbow trout (O. mykiss), and sockeye salmon (O. nerka). After accounting for temporal variability common to the sampled creeks, we find We find that one culvert in the 37 treatment creek seemed to have no impact while the second culvert had a large impact on fish passage. The construction itself seemed to have only transient effects on these salmonid species during the several months after construction two construction events. In the context of billions of dollars of court-mandated road culvert replacements taking place in Washington State, USA, our results suggest that culvert replacement can be conducted with only minimal impact of construction to key species of management concern. Furthermore, eDNA methods can be an effective and efficient approach for monitoring hundreds of culverts to prioritize culverts that are required to be replaced. More broadly, we demonstrate a rigorous, quantitative method for environmental impact reporting using eDNA that is widely applicable in environments worldwide.

46 Introduction

At present, it remains difficult to comprehensively measure the environmental impacts of discrete human activities, despite such assessment often being required by law. Within the United States, both state and federal laws often require a form of environmental-impact assessment for medium- to large-scale projects (i.e., those that might have a significant impact on the environment) (Morgan 2012). Outside the US, many nations have their own versions of these same laws. Specifically when measuring impacts on aquatic ecosystems, assessments generally are based on literature reviews or field measurements of key species selected beforehand (Rubin et al. 2017). These traditional methods are often expensive, rely on just a few species, and are limited in spatial and temporal coverage (Martin et al. 2012). Moreover, they often lack pre-project monitoring and any or sufficient post-project monitoring, given that the goals of a development project normally focus on construction itself and funding is often extremely limited. For example, a recent literature review of stream restoration projects cited that more than half of projects evaluated (62%) had no pre-project monitoring and only sampled once per year (for before, during, and post-project sampling) (Rubin et al. 2017). Thus, current assessment efforts relying on traditional survey methods often fall short in documenting and quantifying environmental impacts. A key difficulty in conducting ecosystem assessments is that there is no one way to survey the world and just 61 'see what is there." All methods of environmental sampling are biased, in the sense that as they capture a selective portion of the biodiversity present (Rubin et al. 2017). Net samples for fish, for example, fail to 63 capture species too small or too large to be caught in the net; bacterial cultures capture only those species that can be cultured on available media, and so forth. Environmental DNA (eDNA), however, comes as close to this goal as any method yet developed although not without bias (see below): a sample of water, soil, or even air, contains the genetic traces of many thousands of species, from microbes to whales. Sequencing eDNA is therefore a means of surveying many species in a consistent and scalable way (Taberlet et al. 2012, Thomsen and Willerslev 2015). Environmental assessments have begun to make use of eDNA for such work around the world (Muha et al. 2017, Duda et al. 2021, Klein et al. 2022, Massri et al. 2022, Moss et al. 2022), but are not yet common practice. Sampling water to collect eDNA before, during, and after a development project would be a new and powerful method of assessing that project's impacts on the local biological communities, and could conceivably become the standard approach to conducting such impact assessments (Hinz et al. 2022). 74 Surveying the natural world by amplifying and sequencing DNA from environmental sources Surveying the world by eDNA has long been commonplace in microbial ecology (Ogram et al. 1987, Rondon et al. 2000,

Turnbaugh et al. 2007) but has recently become popular for characterizing ecological communities of

eukarvotes eukarvotic communities (Taberlet et al. 2012, Kelly et al. 2014, De Vargas et al. 2015, Port et al. 2015, Valentini et al. 2016, Stat et al. 2017). Techniques that take advantage of such data may include non-PCR-based methods such as hybridization, but generally include an amplification step such as quantitative PCR, digital or digital-droplet PCR, or traditional PCR from mixed templates followed by high-throughput sequencing (Ruppert et al. 2019). This last technique is known as metabarcoding, eDNA amplicon-sequencing, or more generally, marker-gene analysiseDNA metabarcoding. In a metabarcoding approach, broad-spectrum PCR primers identify many-hundreds or thousands of taxa across a very wide diversity of the tree of the life (e.g., Leray et al. (2013)), but nevertheless. Nevertheless the absence of a taxon from a sequenced sample does not indicate the absence of that taxon from the environment but rather that the taxon failed to amplify (Shelton et al. 2016, Kelly et al. 2019, Buxton et al. 2021). Instead, the unsampled species simply may not have been susceptible to that set of PCR primers, and so failed to amplify. The result is often a dataset that represents hundreds or thousands of taxa, but these taxa are a fraction of a larger (and perhaps taxonomically broad) pool of species present, Gold et al. 2023). In virtually all comparisons, metabarcoding recovers far more taxa from an area than any other sampling method (Port et al. 2015, Kelly et al. 2017, Seymour et al. 2021). However, we expect results from metabarcoding to differ dramatically from non-PCR based sampling methods due to the fundamental differences in sampling genetic waste residual genetic material as opposed to whole organisms. Furthermore, eDNA analyses rely on several laboratory processes, including PCR amplification, all of which contribute to complicating the interpretation of results (see Shelton et al. (2016) and Kelly et al. (2019)). Specifically, PCR amplification is an exponential process for which the efficiency varies across 97 species and primer set (Gloor et al. 2016). By understanding these process-differences, we can correct for taxon-specific biases in amplification efficiency to yield quantitative estimates of the community composition prior to PCR (McLaren et al. 2019, Shelton et al. 2022). Other ways to conduct quantitative metabarcoding include using qSeq (Hoshino et al., 2021), a process in which a random tag is added to target sequences 101 before PCR. However, if different species amplify at different rates during PCR, these quantifications would 102 reflect not just the starting concentration but also the amplification efficiency. 103 After correcting for amplification biases, the resulting metabarcoding dataset is compositional, revealing the 104 proportions of each species' DNA present in each sample, but importantly, contains no information about the absolute abundance of DNA present (Gloor et al. 2016, McLaren et al. 2019, Silverman et al. 2021, Shelton 106 et al. 2022). We can tie these proportional estimates to absolute abundances using additional data such 107

as a quantitative PCR (qPCR) assay for one of the taxa present. Thus, a single qPCR assay and a single

metabarcoding assay can together provide quantitative estimates of many species as opposed to running as

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many qPCR assays as species of interest —(see also (Pont et al. 2022)). Together, we can use these data 110 to assess changes in eDNA concentrations of species over time, and due to environmental impacts, such as replacing a culvert under a road. 112 Because replacing culverts can require substantial intervention—for example, diverting the water from a creek 113 segment and rebuilding the road with a redesigned culvert—they require environmental impact assessments. 114 Furthermore, because these replacements occur serially according to a schedule, they present an attractive 115 experimental design to use eDNA to assess environmental impacts. As a result of a ruling in a federal court (Martinez 2013), Washington State is under a court-ordered mandate 117 to replace hundreds of culverts that allow water to pass under roads and highways, costing billions of dollars. 118 Improperly designed culverts can lead to many negative consequences for fish, especially anadromous salmon, including habitat fragmentation, loss of accessibility to spawning and rearing habitat, and genetic isolation 120 (Price et al. 2010, Frankiewicz et al. 2021). These impacts on salmonids furthermore violate the sovereign treaty rights of the region's indigenous tribes to manage their people, land, and resources (Schmidhauser 122 1976a (Martinez 2013). Salmonid species are of cultural and economic importance to the indigenous peoples 123 of the region, and without restoration of historic salmon-rearing habitat, the continued decline of salmonids 124 can lead to not only ecological destruction, but the loss of cultural and economic viability for many indigenous 125 tribes (Schmidhauser 1976b1976, Lackey 2003, Long and Lake 2018). Presently, the prioritization process for replacing culverts preventing fish passage conducted by the Washington 127 Department of Transportation is a protocol provided by the Washington Department of Fish and Wildlife, 128 which includes factors such as the amount of habitat blocked by the barrier, the types of species blocked by 129 the barrier, and estimated cost of repair, among other things (Washington Department of Fish and Wildlife 130 2019a 2019). However, data on fish presence upstream of barriers (i.e., culverts blocking fish passage) are rare 131 and often not included in these assessments. Using eDNA as a proxy for fish presence could provide another 132 important type of important data for project prioritization and increase efficiency in making prioritization decisions have the potential to be more cost effective. 134 Once a culvert has been designated as in need of repair, the intention is to improve conditions for biota, 135 including migrating fish, but it might be that the construction itself has might have a short-term negative effect on fish and other organisms before the longer-term improvements are realized. Specifically in the case 137 of culvert replacements, studies have cited the negative impacts of construction to include 138 sediment accumulation, removal of vegetation, and blocking flow and stranding fish (Wellman et al. 2000, 139 Washington Department of Fish and Wildlife 2019b 2019). However, it is unclear how long these effects might

last and if the long-term benefits of the culvert replacement justify the short-term costs of the construction.
These disruptions also underscore the importance of both properly assessing culverts to determine if they are
blocking fish passage and monitoring after construction to ensure the replacement actually improved fish
passage.

Many studies have attempted to quantify if-when culverts are barriers to fish passage and how effective culvert replacements are for fish passage, either by measuring physical parameters of the culvert and stream 146 after replacement (Price et al. 2010), or by measuring biological meansparameters, including electrofishing (Ogren and Huckins 2015) or utilizing genetic differentiation from fish tissues (Wood et al. 2018, Nathan 148 et al. 2018). In some cases, culverts deemed blockages did not prove to block fish passage (MacPherson et al. (2012) found in a study of over 200 culverts, that for certain species, including rainbow trout (O. mykiss), 150 culverts were not blocking fish passage despite being deemed blockages. As for how effective replacements 151 are, , while in others, blockages that were replaced were not found to improve fish passage (Price et al. 152 (2010) found in a study of ~75 culverts that, despite culvert replacement, about 30% of the new culverts 153 still remained blockages (by physical characterization), while or improve overal biotic integrity (Ogren and 154 Huckins (2015) found in a more in-depth study of just three culverts that after biological sampling (i. e., 155 electrofishing and macroinvertebrate surveys) 3-5 years after culvert replacements, the overall biotic integrity was not improved. Sampling water for eDNA analysis before, during, and post-restoration can provide 157 valuable information on if the restoration is needed, how the restoration negatively impacts communities during construction, and if the restoration efforts did in fact correct the blockage. 159

Here, we report the results of a year-long an approximately 18-month eDNA sampling effort before, during,
and after a small construction project in our experimental the replacement of two culverts (one small and one
large) in a creek, assessing the impact of that project these projects on the salmonid species present. We do
so using a combination of metabarcoding (12s mtDNA) and qPCR to yield estimates of the concentrations of
DNA present at each time point, and we use parallel samples from an additional four control creeks to develop
a causal analysis of changes in these concentrations. A clear opportunity for policy-relevant eDNA work is in
using its power to survey many species at a time to improve the way we assess the impacts of human activities.
Here, we demonstrate the utility of eDNA for such assessments policy-relevant environmental assessments by
surveying many species simultaneously and improving the way we assess the impacts of human activities.

Methods

170 Site and Species Selection

We used selected sampling locations based loosely on an asymmetrical BACI (Before-After-Control-Impact) 171 study design (Underwood 1992, 1994, Benedetti-Cecchi 2001) to measure the environmental impact of a 172 construction project replacing an under-road culvert culvert replacement using eDNA. We sampled four 173 control creeks in addition to the treatment creek where the culvert was replaced where construction was not 174 occurring, (Figure 1) at monthly intervals, both upstream and downstream of each creek's culvert. The eulvert 175 two culverts in the treatment creek (Padden) was were suspected to be partially impassible and thus was were 176 removed and replaced during the course of the study; one of the control creeks had a bridge, which allowed 177 . The four control creeks ranged from preventing fish passage (Portage), one control creek had a culvert 178 classified as having limited fish passability Barnes and Chuckanut), partially passable (Squalicum), and two 179 control creeks had culverts classified as preventing to allowing fish passage (Barnes and ChuckanutPortage; 180 see Supplemental Text 1) (Washington Department of Fish and Wildlife 2019a2019). These creeks were chosen due to their comparable size, flow, watersheds, and species presumed to be present to constrain as 182 many ecological variables as possible. The intervention (i.e., culvert replacement first culvert replacement (SR-11) in Padden Creek occurred over 184 about two months and included the "de-watering" of the creek, removal of the existing culvert, installation 185 of the new culvert, and then the "re-watering" of the creek from late August 2021 to early October 2021 186 (Supplemental Figure 3). We Text 1; Figure S1.4). The second culvert replacement (I-5) in Padden Creek 187 was a much larger construction project, including daylighting the creek and building a bridge under a large, five-lane interstate. In-water work for the I-5 culvert replacement began in late June 2022 and was completed 189 in September 2022. By sampling before, during, and after both construction events, we were then able to 190 quantify isolate the effect of the culvert replacement itself – controlling for temporal trends, background 191 environmental variability, and sampling variability – using a Bayesian time-series model to jointly model 192 salmon linear mixed effects model of eDNA abundances across creeks, time points, sampling stations, and 193 species. 194 Because salmonids are the primary species of management concern in these creeks, we focus the present 195 analysis on the four salmonid species most common in our data: cutthroat trout (Oncorhynchus clarkii), coho 196 salmon (O. kisutch), rainbow/steelhead trout (O. mykiss), and sockeye/kokanee salmon (O. nerka). As further 197 described below, we surveyed the salmonid DNA present in each creek via eDNA metabarcoding (targeting 198 a region of the 12s mtDNA gene) and complementary quantitative PCR (qPCR; targeting a region of the 199

CytB gene) for a reference species (cutthroat trout, O. clarkii), which in combination yielded quantitative
estimates for each fish species throughout the study area.

Not all four salmonids are expected to be found in all five of the creeks sampled. As documented by WA
Department of Fish and Wildlife SalmonScape (http://apps.wdfw.wa.gov/salmonscape/map.html), all creeks
contain cutthroat trout, steelhead trout, and coho salmon. Barnes Creek is the only creek documented to have
kokanee salmon , which are (a freshwater sub-type of sockeye salmonthat are not anadromous). However,
local spawner surveys conducted by the City of Bellingham from 2015-2020 in Padden Creek documented
kokanee salmon, as well as the other three species and importantly, several unknown species of live and dead
fish and redds (nests dug by fish in gravel to deposit eggs) ((City of Bellingham 2015).

The four salmonid species in this study have different life histories and behaviors that would impact when
fish (and therefore eDNA concentrations) occur in the creeks. For these four migratory salmonids, the run
timings vary for each species in the study area (Bellingham, WA). Adult coastal cutthroat (O. clarkii) are
documented to run throughout the entire year, whereas coho salmon (O. kisutch) run from September to
December, sockeye salmon (O. nerka) run from October to December, and steelhead trout (O. mykiss) run
from November to June. For migrating coho (O. kisutch) and steelhead trout (O. mykiss), juveniles may be
present in the creeks year-round (Supplemental Figure 3). Using eDNA methods, it cannot be determined
if the DNA found is sourced from adult or juvenile animals.

Furthermore, three of the four species in this study have both freshwater resident and saltwater migrating 217 behavior. Cutthroat trout (O. clarkii) encompasses both non-migrating, resident trout in the creeks and 218 coastal run cutthroat that migrate into Padden Creek from saltwater (Bellingham Bay). Similarly, O. nerka 219 includes both anadromous sockeye salmon and freshwater resident kokanee salmon and O. mykiss includes 220 both anadromous steelhead trout and non-migrating rainbow trout. Using eDNA, we cannot distinguish 221 between the migrating and non-migrating subspecies of O. clarkii, O. nerka, and O. mykiss. For the fish 222 exhibiting migratory behavior, the run timings vary for each species in the study area (see Discussion and 223 Figure S1.4). Therefore, our eDNA concentrations might reflect contributions from both migrating and 224 non-migrating individuals at any given time point in the dataset.

226 Water Sampling

We collected water samples monthly between From March 2021 and to February 2022in each of five salmonid-bearing creeks in northwest Washington State, USA (Figure 1). We sampled each stream above and below under-road culverts., all five creeks were sampled monthly (n=12). Monthly sampling continued in Portage Creek, Padden Creek, and Squalicum Creek through August 2022, with one additional sampling

point in December 2022 (n=19). At each sampling station (N=2, upstream and downstream of a culvert) at 231 each creek(N = 5) in each month (N = 12), we collected three 2-liter water samples, for a total of 360 water samples. Samples were collected using Smith Root's an eDNA Backpack (Smith Root; Thomas et al. (2018)), 233 a portable pumping-and-filtering device set to filter at 1 L/min at 82.7 kPa (12 psi). In some months, less than 2 L of water was filtered due to clogging $\frac{\text{(min} = 1.02 \text{ L, mean} = 1.97 \text{ L, median} = 2.01 \text{ L; see Supplemental}}{\text{Supplemental}}$ 235 Figure 4). Water samples were filtered using single-use inlet tubes through $5\mu m$ self-preserving filters (Smith Root, Vancouver, WA), which were then dried and kept at room temperature until DNA extraction within 1 237 month of collection (Thomas et al. 2019). 238 Over the course of the year of sampling, water discharge varied from very low to no flow in summer months to high flow in winter months (Figure 2). Thus, when considering eDNA concentrations at a sampling site, 240 we need to account for the large difference in water volume over the course of the year-long time series. In 241 other words, given the same number of fish and a constant eDNA shedding rate, we would expect to see 242 higher concentrations of DNA in summer months and lower concentrations in winter months due to dilution 243 of eDNA in higher water volumes just from the difference in flow. Other eDNA time series datasets also 244 correct for discharge to present eDNA data as a mass flow rate mass/time(Tillotson et al. 2018, Thalinger 245 et al. 2019). Here, we convert eDNA). We account for this dilution by converting eDNA concentration $[copies/\mu L]$ to an eDNA mass flow rate [copies/s] by multiplying eDNA concentrations by discharge [L/s]. 247 (Tillotson et al. 2018, Thalinger et al. 2019). Flow gauges maintained by the United States Geological Survey 248 (USGS) were used for Padden Creek (USGS Gauge 12201905), Chuckanut Creek (USGS Gauge 12201700), 249 and Squalicum Creek (USGS Gauge 12204010; https://maps.waterdata.usgs.gov/mapper/index.html; 250 U. S. Geological Survey (1994); Supplemental Figure 1). During the year Figure S1.1). Over the course 251 of sampling, the flow gauges at Chuckanut Creek and Squalicum Creek became inoperable after a major 252 flooding event. To find discharge rates for Chuckanut and Squalicum Creeks, five years of historical data (2015-2020) were used to generate a daily-monthly averaged correction factor based on Padden Creek. 254 For the year of sampling (2021-2022), the discharge rates used at Chuckanut and Squalicum Creeks were estimated based on the correction factor from Padden Creek (Supplemental Figure 2(Supplemental Text 1, 256 Figure S1.3). No discharge data was available for Portage Creek or Barnes Creek. Based on field sampling 257 conditions, the discharge from Padden Creek was used as a proxy for both Portage and Barnes as they 258 are in similarly sized watershed areas and land-cover characteristics. Though in the year of sampling, the 259 discharge in Padden Creek ranged from no metered flow to 23 m³/s, the discharge on the dates of sampling only reached a maximum of 1.3 m³/s. 261

DNA Extraction, Amplification, Sequencing

All molecular work prior to sequencing was performed at the University of Washington. Bench-tops were 263 cleaned with 10% bleach for 10 minutes and then wiped with 70% ethanol. Molecular work was separated onto pre- and post-PCR benches; all DNA extractions and PCR preparation was conducted on a bench 265 where no PCR product was handled. DNA was extracted from half of each filter-Details of the molecular work can be found in Supplemental Text 1. Briefly, DNA was extracted off filters using a Qiashredder column 267 (Qiagen, USA) and the DNEasy DNeasy Blood and Tissue Kit (Qiagen, USA) with an overnight incubation 268 (Supplemental Text 1, Thomas et al. (2019)), such that the effective filtering effort was 1 L/sample; the 269 remaining half of each filter was archived at -20C. Extracts were eluted in 100 μ L of molecular grade water, 270 quantified via Qubit (Invitrogen, USA) and stored at -20°C until PCR amplification within 2 months of 271 extraction. 272 For the metabarcodingapproach metabarcoding, we targeted a ~186 bp hypervariable region of the mitochondrial DNA 12S rRNA gene for PCR amplification (MiFish; Miya et al. 2015), but using modified 274 primer sequences as given in Praebel and Wangensteen (unpublished; via personal communication) and 275 including the Illumina Nextera overhang sequences for subsequent indexing. The primers used were as 276 follows: F-5' TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGCCGGTAAAACTCGTGCCAGC 3', 277 R. 5' GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCATAGTGGGGGTATCTAATCCCAGTTTG 278 3' (italies indicate Nextera overhang). The. The primer sequences, final reaction recipe, and cycling 279 conditions can be found in Supplemental Text 1. Each month of samples was amplified on a single plate 280 with the addition of a no template control (NTC; molecular grade water in lieu of template) and a positive 281 control (genomic DNA from kangaroo). After PCR amplification, PCR, a species not present in the 282 environment). PCR products were visualized a 1-2% gel. If no band was present for a given sample, a 283 new amplification was attempted with extracts diluted 1:10 iteratively until a band was detected. PCR 284 products were, size-selected and cleaned using MagBind Beads (Omega Biotek, USA) at a sample: beads 285 ratio of 1.2. Bead-cleaned PCR products were clutted in 30 µL of molecular grade water and quantified via 286 Qubit (Invitrogen, USA). 287 An indexing PCR reaction added a unique index, and diluted iteratively if inhibited. After cleaning, a second 288 PCR amplification added unique indices to each sample using Nextera indices (Illumina, USA) to allow pooling 289 multiple samples onto the same sequencing run (See Supplemental Text 1 for details). Indexed PCR products 290 were also size-selected and purified using MagBind Beads (Omega Biotek, USA) at a sample: beads ratio of 291 0.8. Bead-cleaned PCR products were cluted in 30 µL of molecular grade water and quantified via Qubit. 292 Indexed and bead-cleaned products were normalized before pooling into libraries, which were subsequently 293

quantified via Qubit and visualized on a Bioanalyzer (Agilent, USA) before visualized before pooling for sequencing. Samples were randomized in 3-month blocks and each block split across 3 sequencing runs to avoid run effects, for a total of 12-14 sequencing runs. The loading concentration of each library was 4-8 pM and 5-20% PhiX was included depending on the composition of the run. Sequencing was conducted using an Illumina Miseq with v3 2x300 chemistry at the NOAA Northwest Fisheries Science Center and the University of Washington's Northwest Genomics Center.

Here, we used mock communities to determine the species-specific amplification efficiencies for each salmonid

in the study. Briefly, we-We constructed five communities with known proportions of starting DNA from 301 different species (total DNA as measured by Qubit). The communities ranged from having a total of 12 to 20 302 species, but six salmonid species were included in all five mock communities to have more information on the 303 amplification efficiencies of salmonids (Supplemental Table 32). We sequenced these communities using the 304 same metabarcoding primers and thermocycling conditions above and then determined the species-specific 305 amplification rates given the discrepancy between the known starting proportion and the proportion of 306 reads after sequencing. These The mock community data were then used to correct the sequencing reads 307 from the environmental samples to estimate the starting DNA proportions of each species in environmental 308 samples, which is the metric of interest (Figure 3, green boxes). This is the first application of the model to correct eDNA data from water samples with mock community data as described in Shelton et al. (2022) (see 310 Supplemental Text 2 for more information).

312 Bioinformatics

After sequencing, bioinformatic analyses were conducted in R (R Core Team 2017). A more detailed description of the bioinformatics pipeline is included in the supplement (Supplemental Text 1). Supplemental Text 1.

Briefly, primer sequences were removed using Cutadapt Cutadapt (Version 1.18) (Martin 2011) before dada2

dada2 (Callahan et al. 2016) trimmed, filtered, merged paired end reads, and generated amplicon sequence variants (ASVs). Taxonomic assignment was conducted via the insect insect package (Wilkinson et al. 2018) using a tree generated by the developers for the MiFish primers that was last updated in November 2018.

Only species level assignments from insect insect were retained and ASVs not annotated or not annotated to species level were then checked against the NCBI nucleotide database using BLAST+ (Camacho et al. 2009).

Query sequences that matched a single species at >95% identity were retained.

In total, sequencing runs generated ~42 million reads across all environmental samples (12 months x 2 stations x 5 creeks x 3 biological replicates = 360 filters) and 27 mock community samples (3 communities x 9 replicates 6 even, 3 skewed proportions) for calibration (see below). After quality-filtering and merging

samples, of which -81% of reads and -2% of ASVs were annotated to species level (per sample: mean = 78%, median = 88%, min = 0%, max = 99.99% of reads annotated). We only focus on the metabarcoding data 327 from four salmonids for the remainder of this paper. The four salmonids represent ~55\% of all environmental reads and ~68% of the annotated reads found in environmental samples. 329 In the mock community samples, 98.7% of the ~5 million reads after quality filtering were annotated to 330 species level. Importantly, the target salmonid ASVs in the mock communities were found in environmental samples, unambiguously linking the taxa in calibration samples with those in environmental samples. The 332 most common salmonid species found in the environmental samples was O. clarkii (cutthroat trout), which was found in ~90\% of samples, followed by O. kisutch found in ~60\% of samples, then O. mykiss found 334 in -40% of samples, and finally O. nerka found in -5% of samples. Not only was O. elarkii found in the 335 majority of environmental samples, but also ~63\% of samples across all times, creeks, and stations had at 336 least 50% of reads assigned to O. clarkii. 337

all runs, ~33 million reads remained from ~21,000 amplicon sequence variants (ASVs) in the environmental

338 Quantitative PCR and Inhibition Testing

We quantified cutthroat trout (O. clarkii) DNA in each sample, targeting a 114 bp fragment of the cytochrome b gene with a qPCR assay (Duda et al. 2021). The primer/probe sequences were: F 340 5' CCGCTACAGTCCTTCACCTTCTA 3', R 5' GATCTTTGTATGAGAAGTAAGGATGGAA 3', P 341 5' 6FAM-TGAGACAGGATCCAAC-MGB-NFQ 3'. The qPCR assay was multiplexed with TagMan 342 Exogenous Internal Positive Control Reagents (EXO-IPC) (Applied Biosystems, USA) to check for the 343 presence of PCR inhibitors (Duda et al. 2021). The EXO-IPC mix includes the primers and probe for the EXO-IPC DNA, with the probe having a VIC reporter, allowing it to be multiplexed with the O. clarkii 345 assay, which has a FAM reporter. Each DNA sample was run in triplicate; the final recipe, final recipe, and thermocycling conditions can be found in Supplemental Text 1. All qPCRs were conducted on an Applied 347 Biosystems StepOnePlus thermocycler. Each plate included a 8-point standard curve created using synthetic DNA (gBlocks) at the following 349 concentrations: $100,000 \text{ copies}/\mu\text{L}$, $10,000 \text{ copies}/\mu\text{L}$, $1,000 \text{ copies}/\mu\text{L}$, $100 \text{ copies}/\mu\text{L}$, $10 \text{ copies}/\mu\text{L}$, 10 copies/350 $copies/\mu L$, 3 $copies/\mu L$, 1 $copy/\mu L$ Additionally, six no template controls (NTCs) were included on each plate: 3 with the IPC DNA mix and 3 with molecular grade water instead of template or IPC DNA mix. 352 Plates were re-run if efficiency as determined by the standard curve was outside of the range of 90-110%. 353 To check for inhibition, the cycle threshold (Ct) value determined for the Each DNA sample was run in triplicate and was checked for inhibition using the EXO-IPC assay from the NTC was compared to the Ct

the mean Ct for the NTCs, the sample was deemed inhibited and diluted 1:10 and re-assayed until the Ct value fell within the accepted range. (Applied Biosystems). The majority of environmental samples (6560%) were inhibited and accordingly diluted for analysis. In 7580% of inhibited samples, a 1:10 dilution or less remedied the inhibition, but some samples (11%) required dilution by a factor of up to 1000 (Supplemental Figure 5)... 1000. Each plate included a 8-point standard curve created using synthetic DNA (gBlocks) ranging from 1 to 100,000 copies/μL and six no template controls (NTCs) were included on each plate with molecular grade water instead of template. All qPCRs were conducted on an Applied Biosystems StepOnePlus thermocycler.

All qPCR data was processed in R using Stan (Stan Development Team 2022), relating environmental samples to the standard curve via a linear model (Figure 3, blue boxes; Figure S2.1). We amended the standard linear regression model to more realistically capture the behavior of qPCR observations, accommodating non-detections as a function of underlying DNA concentration, and letting the standard deviation vary with the mean (lower-concentration samples had more uncertainty). See McCall et al. (2014) and Shelton et al. (2019) for similar models; see Supplemental Text 2 for full statistical details. Subsequent analysis corrected for sample-specific dilution if found inhibited and corrected for any variation in water-volume filtered during sample collection. Samples with standard deviations between technical replicates larger than 1.5 Ct values were removed from analyses.

374 Quantitative Metabarcoding

The intercalibration of the mock community samples demonstrated the rank order of amplification efficiencies for salmonids (Supplemental Figures 14 and 15). 10 and 11). Cutthroat trout (O. clarkiiand 376) and sockeye/kokanee salmon (O. nerka) had similar amplification efficiencies, both of which were higher than rainbow/steelhead trout (O. mykissand) and coho salmon (O. kisutch), which had the 378 lowest amplification efficiency. Calibrated metabarcoding analysis yielded quantitative estimates of the proportions of species' DNA in environmental samples prior to PCR. We then converted these 380 proportions into absolute abundances by expansion, using the qPCR results for our reference species, 381 We estimated the total amplifiable salmonid DNA in environmental cutthroat trout (O. clarkii). 382 sample i as $\frac{\text{qPCR}_{\text{reference}_i}}{\text{Proportion}_{\text{reference}_i}}$, $\frac{\text{Camplifiable}_i}{\text{Reportion}_{\text{reference}_i}}$, where C has units of [DNA] 383 copies/ull and then expanded species' proportions into absolute concentrations by multiplying these sample-specific total concentrations by individual species' proportions, such that for species i in sample 385 $i, \frac{\text{DNA}_{i,j} = \text{DNA}_{\text{salmonid}_i} * \text{Proportion}_{i,j}}{\text{C}_{i,j} = \text{C}_{\text{amplifiable}_i} * \text{Proportion}_{i,j}}$. Here, we combine the modeled output of the qPCR model for *O. elarkii* cutthroat trout (Figure 3 dashed blue box) and modeled proportions
of salmonid DNA from metabarcoding (Figure 3 dashed green box). Though Although in the future this
could be used as a joint model, here the precision of our modeled estimates were very high such that we used
the mean of the posterior estimates from each model to move forward as input to the time series model
(Figure 3 dashed purple box; see Supplemental Text 2 for more details).

Finally, due to the range of water discharge over the course of the year, we converted from DNA concentration [copies/L] to a mass flow rate [copies/s] after multiplying by the discharge of each creek [m³/s] (Figure 3, solid purple boxes).

₃₉₅ Estimating the Effects of Culvert Replacement and of Culverts Themselves

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Consistent with the asymmetrical BACI study design, we generated data from our-We sampled four control 396 creeks as context against which to compare the observations in Padden Creek, our treatment creek where 397 the two culverts were being replaced. At a given station in a given creek, some DNA concentration exists 398 for each species. For simplicity, we focus on a single species and a single station (downstream or upstream) for the moment. Our observations of the (log) DNA concentration in creek i at time t are distributed as 400 $Y_{i,t} \sim \mathcal{N}(\mu_{i,t}, \sigma^2)$. More complex versions of the model may let σ vary across creeks, time points, species, or with environmental covariates of interest. Recognizing that these observations are autocorrelated in time, we 402 use an AR(1) autocorrelation model, implemented in Stan via R, to capture the observed temporal trends. 403 We observe the log-DNA concentration are interested in how the DNA concentration changes over time, so we assert that the expected value of DNA in a creek at time t, Y, for a given species in a given sample as 405 a random variable drawn from a normal distribution with mean μ and observation variance $\sigma^2 \mu_{k,t}$, depends 406 upon its time point, in some way. We considered three ways of modeling the salmonid eDNA data, each in 407 a Bayesian framework, but each treating non-independence among time points somewhat differently: 408

- A linear auto-regressive (AR(1)) model, written in stan. For each species j in each creek, the expected log-DNA concentration μ at time t in creek i at station d concentration of eDNA of each month is a linear function of the DNA concentration for the same creek/station at t-1. expected value from the previous month. Within a species, the monthly autoregressive parameters are shared across creeks.
- A generalized additive model (GAM), written in brms (which itself writes a stan model). For each species in each creek, an independent set of spline (weighting) parameters describes the temporal trends in expected eDNA concentration; the number of spline knots is shared across species and creeks.
 - A linear mixed-effects (LME) model, written in rstanarm. For each species in each creek, sampling

month is treated as a random effect. Each species-creek-month effect is treated as an independent draw
from a common distribution.

$$\frac{Y_{i,t,d} \sim \mathcal{N}(\mu_{i,t,d}, \sigma^2)}{\mu_{i,t,d} = \alpha_{i,t} + \epsilon_{i,t,d} + \eta_{i,t,d}}$$

$$\epsilon_{i,t,d} \sim \mathcal{N}(\beta \mu_{i,t-1,d}, \phi^2)$$

- Ultimately, the three models yielded very similar results (Figure S2.2), and the LME model proved simplest and most flexible insofar as it could easily handle datasets with uneven sets of observations – for example,
- cases in which a species was detected downstream of a barrier, but not upstream.
- Intercept α varies by time, creek, and species, capturing creek-level deviations from the previous time-step.
- ϵ The autoregression term ϵ is itself a random variable drawn from a normal distribution with expected
- value $\beta \mu_{i,t-1,d}$ and process variance ϕ^2 , such that the species-specific slope term β estimates the degree of
- 425 autocorrelation in log-DNA concentration between one time-step and the next. The model shares information
- $_{426}$ across creeks and time-points via β . In R code using rstanarm, this model is coded as
- 427 Finally, η captures the difference in log-DNA concentration between upstream and downstream stations
- within a creek; we set $\eta_{d=1} = 0$ such that the value of $\eta_{d=2}$ explicitly captures the effect of the culvert within
- a given creek at a given time. The effect of construction in our focal Padden Creek, then, is the change in η
- 430 after construction versus prior to construction. We fit this model in a Bayesian framework using moderately
- informative priors on all parameters, and confirmed model convergence $(\hat{R} < 1.01)$ across 3 chains and 2500
- 432 model iterations. See statistical supplement (Supplemental Text 2) for prior values, diagnostics, and full
- 433 model details.
- 434 stan_glmer(log(observed) ~ (1 + time_idx|creek:species) + (1|station:creek:species:time_idx)
- See Supplemental Text 2 for more details on the linear mixed-effects model.

436 Results

- 437 Metabarcoding and Quantitative PCR
- In total, we generated ~ 52 million reads across all environmental samples and 27 mock community samples (3
- communities x 9 replicates [6 even, 3 skewed proportions]) for calibration (see below). After quality-filtering

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and merging all runs, ~45 million reads remained from ~24,000 amplicon sequence variants (ASVs) in the
440
    environmental samples, of which \sim 83\% of reads were annotated to species level (per sample: mean = 82%,
    median = 93\%, min = 0\%, max = 99.99\% of reads annotated). We only focus on the metabarcoding data
442
    from four salmonids for the remainder of this paper. The four salmonids represent ~54% of all reads and
    ~64% of the annotated reads found in environmental samples.
444
    In the mock community samples, 98.7% of the ~5 million reads after quality filtering were annotated to
445
    species level. Importantly, the target salmonid ASVs in the mock communities were found in environmental
    samples, unambiguously linking the taxa in calibration samples with those in environmental samples. The
447
    most common salmonid species found in the environmental samples was cutthroat trout (O. clarkii), which
    was found in ~85% of samples, followed by coho salmon (O. kisutch) found in ~62% of samples, then
449
    rainbow/steelhead trout (O. mykiss) found in \sim 40\% of samples, and finally sockeye/kokanee salmon (O.
450
    nerka) found in ~10% of samples. Over half of the samples across all times, creeks, and stations had at least
451
    50% of reads assigned to cutthroat trout.
452
    After calibrating metabarcoding data using mock communities (See Supplemental Texts 1 and 2), we estimated
    the salmonid composition across time points, creeks, and stations (Figure ?? Figures 4 and 5). The culvert in
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    one control creek (Barnes) appeared to be nearly a total barrier to salmonid passage, with salmonid eDNA
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    detected upstream of the culvert at only three time points, in contrast to being detected at every time point
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    in the downstream station of the same creek. The other four creeks had no such pattern associated with
457
    the culverts, suggesting that fish passage may have been possible through the culverts, or that there were
458
    resident populations upstream of the culverts.
459
    All environmental samples were quantified for absolute concentrations of cutthroat trout DNA across 30-32
460
    qPCR plates, resulting in \frac{280 \text{ samples}}{630 \text{ samples}} (\frac{60\%}{630 \text{ samples}}) with a positive detection in at least 1 of 3
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    technical replicates. The modeled output of cutthroat trout DNA concentrations, ranged from 10-50 copies/L
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    to 1.4 \times 10^6 copies/L, with a mean value of -58 - 47,000 copies/L (Figure 6).
    We combined compositional information from metabarcoding with absolute concentrations from qPCR for
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    our reference species, cutthroat trout (O. clarkii, from the qPCR), to estimate the total concentration of
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    DNA for each species (See Supplemental Text 2). The joint time-series model shared information across
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    stations and creeks; consequently, data from one of the control creeks (Barnes) could not be included because
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    of the nearly total absence of salmonids upstream of its culvert. However, data from the remaining creeks
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    characterized trends in the other. These quantitative data for all four target species well and could be modeled
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appropriately—were then used in the linear mixed-effects model to assess salmonid trends over time (Figure

471 7).

472 Effects of Culverts

Before considering the effect of construction, the difference in abundance trends between upstream and 473 downstream stations (Figure 7) demonstrates that the culverts themselves have some effect, but generally not 474 a large effect on the salmonid species surveyed. Therefore, these four creeks (which include 3 culverts and 475 one bridge) do not seem to be blocking salmonid passage, A noteable A notable exception was Barnes Creek, which was not included in the time series model, as the culvert was so clearly a barrier as most time points 477 had no salmonid DNA upstreamand therefore models including Barnes do not converge as a result of the large fraction of sampling points with no observations of salmonids. (Figure ??). Padden Creek upstream 479 of I-5 also was more clearly a barrier to fish passage, while the culvert across SR-11 seemed to be less of a barrier to fish passage. In other cases, salmonid DNA is found upstream but not downstream, indicating 481 that the culvert is likely not a barrier and there are resident individuals upstream of the culvert. 482 Summarizing over all species and the four creeks used in the time series model, the effect was largest during the dry periods of late summer / early fall (July to October), when flows were at a minimum (i.e., 484 September) and the connectivity culvert effect was minimal (Figure 8); the average log-fold change between 485 upstream and downstream was low (Figure 8). Salmonid species DNA concentrations were higher upstream 486 than downstream during this period, with mean upstream DNA concentrations only about 5% higher than 487 downstream DNA concentrations, sites was not significantly different from zero. Individual species' patterns 488 were similar, indicating that there is not a species-specific effect where culverts block the passage of 489 some salmon but not others (Supplemental Figure 17). A notable exception is O. kisutch in Chuckanut Creek, which was overall much more variable where salmonid DNA concentrations were up to 30% higher 491 upstream than downstream at certain time points and up to 44% higher downstream and upstream at others. Across all species Figure S1.12). The maximum positive log-fold change (i.e., upstream having a higher mass 493 flow rate) was 2.78 in Squalicum Creek for coho salmon (O. kisutch) in August 2021, while the maximum negative log-fold change (i.e., downstream having a higher mass flow rate) was -1.11 found in Squalicum 495 Creek for cutthroat trout (O. clarkii) in December 2021 (Figure S1.12). Of all species, creeks, and time points, Squalicum Creek had the lowest mean percent difference in upstream and downstreamsalmonid DNA 497 concentrations 23 of the 151 observations were within a log-fold change of -0.1 to 0.1, which corresponds with 498 eDNA mass flow rates upstream within 10% of mass flow rates downstream. 499 We also considered how the log-fold change in eDNA mass flow rates were impacted by the flow rate or 500

discharge of the creek itself (Figure 9. We found that at months of the lowest flow (summer months), the

log-fold changes between mass flow rates were the highest, while in winter months with highest discharge
 the log-fold changes were lower and downstream sites often had higher eDNA mass flow rates than upstream
 sites (Figure S1.12).

505 Effects of Culvert Replacement

Fish were excluded from Padden Creek on August 30th, 2021 in preparation for the stream to be diverted 506 on September 9th, 2021 and the diversion was removed October 7th, 2021. Water sampling occurred on September 10th, 2021, the day after the diversion, and on October 12, 2021, just 5 days after reconnecting 508 the stream (Supplemental Figure 3). By comparing the difference in upstream and downstream concentrations mass flow rates before and after construction in Padden Creek, we can assess how large of an impact the 510 replacement two culvert replacements had on salmonid species. 511 (Figure 10). The effects of the culvert replacement operation operations appeared to have been transient 512 and fairly minor for the four salmonid species surveyed. After the beginning of construction in September 513 2021 through the end of sampling in February 2022, we We saw very minor fluctuations in the difference 514 between upstream and downstream salmonid DNA concentrations mass flow rates, and did not see an increase 515 in this difference due to the culvert removal as the log-fold changes in Padden Creek were similar to those 516 in the control creeks at the same time points (Figure 10, grey shading points vs. no shading). Overall, O. 517 clarkii was the least impacted species of the construction while O. nerka was the most impacted species, 518 likely due to the very low concentrations in the creek and the migration timing of O. nerka being during and 519 post-restoration. The mean percent difference across all species prior to construction was 0.18% compared 520 to 1.6% during and post-construction (Supplemental Table 2 black points in areas of yellow shading).

2 Discussion

523 Environmental DNA can provide quantitative measurements of environmental impacts

Here, we used both eDNA metabarcoding and a single species-specific qPCR assay to rigorously quantify
both the effect of culverts and the impact of a culvert replacement on salmonids two culvert replacements
on salmonids in the same creek. We observed a clear seasonal pattern in the DNA concentrations of four
salmonid species detected in the study. The BACI sampling design and the time series model leveraged shared
information across linear mixed effects model leveraged information across treatment and control creeks to
integrate the change in eDNA concentrations due to time, whether a sample was collected below or above
a barrier (i.e., culvert), and whether or not there was construction occurring. Thus, we could isolate the
changes in eDNA concentrations as a result of the intervention (i.e., construction) while accounting for the

variance due to time and station (i.e., season and culvert).

A few other studies have used eDNA to measure environmental impacts in rivers and streams. Duda et al. 533 (2021) used 11 species-specific qPCR assays to document the distribution of resident and migratory fish after a large dam removal (Elwah Dam in project (Elwha River near Port Angeles, Washington). No eDNA 535 sampling was conducted before the dam removal, but the study provided a wealth of information about species returning after the dam removal, providing a very important dataset to use eDNA to monitor ecological 537 changes due to human intervention. Similarly, Muha et al. (2017) sampled three locations upstream and three locations downstream before and after the removal of a weir that was thought to be a barrier to salmonid 539 migrations. The authors only sampled once before and twice after the removal, spanning about a year, and used eDNA metabarcoding to look at the presence/absence of species detected. They found that in fact the 541 before sample demonstrated that the weir was not preventing fish passage (similar to the results found in this study) and furthermore documented a slight increase in alpha diversity in the first time point after the 543 barrier removal and then a return to a similar alpha diversity in the second time point after the removal (similar results found in this study using eDNA concentrations rather than diversity). Finally Yamanaka and Minamoto (2016) sampled along a river with three barriers, finding some fish able to cross barriers and 546 some not, suggesting that the eDNA can indicated habitat connectivity for fishes across barriers.

Importantly, our study demonstrates the value of combining a single qPCR assay with metabarcoding data to generate quantitative estimates of eDNA concentrations of many species without requiring n-n qPCR assays 549 for m-n species of interest. Here, we ultimately only quantified the impacts of four species, but importantly, we did not know a priori how many species of interest there might be and we reduced our efforts two fold by 551 only conducting two assays (one species-specific qPCR and one metabarcoding assay) as opposed to four assays (four species-specific qPCR assays). This can also be particularly helpful for taxa that don't have a 553 previously published qPCR assay, but are detected using universal metabarcoding assays. Metabarcoding data alone only gives compositional data, which cannot be used in a time series to quantify environmental impacts because there is no information about absolute eDNA concentrations. However, by anchoring or grounding proportions using a single qPCR assay, the proportional data can be turned into quantitative data. 557 The species for which to run the qPCR assay can be determined after the metabarcoding is completed; the most commonly found species with a robust qPCR assay should be used to glean the most information.

560 Fish life histories and expected patterns

The four salmonid species in this study have different life histories and behaviors that would impact when fish (and therefore eDNA concentrations) occur in the creeks. Three of the four species in this study have both freshwater and anadromous populations. Cutthroat trout (O. clarkii) encompasses both non-migrating, resident trout in the creeks and coastal run cutthroat that migrate into Padden Creek from saltwater (Bellingham Bay). Similarly, O. nerka includes both anadromous sockeye salmon and freshwater resident kokanee salmon and O. mykiss includes both anadromous steelhead trout and non-migrating rainbow trout.

Using eDNA, we cannot distinguish between the migrating and non-migrating subspecies of O. clarkii, O. nerka, and O. mykiss. Therefore, our eDNA concentrations might reflect contributions from both migrating and non-migrating individuals at any given time point in the dataset.

For these four anadromous salmonids, the run timings for the migrating populations vary for each species in the study area (Bellingham, WA). Adult coastal cutthroat (O. clarkii) are documented to run throughout the

the study area (Bellingham, WA). Adult coastal cutthroat (*O. clarkii*) are documented to run throughout the
entire year, whereas coho salmon (*O. kisutch*) run from September to December, sockeye salmon (*O. nerka*)
run from October to December, and steelhead trout (*O. mykiss*) run from November to June. For migrating
coho (*O. kisutch*) and steelhead trout (*O. mykiss*), juveniles may be present in the creeks year-round (Figure
S1.4). eDNA methods at present cannot distinguish adults versus juveniles from DNA found in a water
sample.

Despite the mix of migrating and non-migrating populations and various run timings, our metabarcoding 577 data demonstrate that in Padden Creek, there was a clear signal of sockeye/kokanee salmon (O. nerka) 578 both upstream and downstream only in November 2021 - February 2022 (and only upstream in March 579 2021).—and again in December 2022. This signal corresponds well with the documented run timing of 580 October to December and the presence of out-migrating juveniles in early spring. In contrast, cutthroat 581 trout (O. clarkiiand) and coho salmon (O. kisutch) were found nearly year-round in Padden Creek. The 582 persistent signal from O. clarkii could be explained by resident cutthroat trout. However, O. kisutch does not have a resident subspecies and the run timing is only documented from September to December. This 584 could potentially be due to juveniles maturing and residing in the creeks for 1-2 years after hatching while adults migrate into the creeks only during the run time to spawn. Visual surveys are conducted rarely and 586 even if they were conducted, it might be difficult to identify juveniles to species level (e.g., snorkel surveys, electrofishing, smolt traps) are conducted infrequently to determine adult and juvenile salmonid abundances. 588 Though O. kisutch eDNA was found year round, the highest concentrations were found near the expected 589 run timing as expected and the life history of O. kisutch includes rearing year-round in freshwater. Finally, 590 though the lowest concentrations on average, rainbow/steelhead trout (O. mykiss) was also found nearly 591 year-round in Padden Creek, which could be contributions from migrating steelhead (November to June), 592 juveniles maturing and migrating, or from resident rainbow trout. Though the O. mykiss signal is found 593 year-round, the highest concentrations do seem to correspond with the steelhead run timing.

Interpreting eDNA with respect to fish abundance and flow

By capturing residual eDNA from water samples, we are measuring a different signal than counting how
many fish are in the creek at each time of sampling. We should not expect the eDNA concentration for each
salmonid to directly correlate to the number of fish in the creek at the time of sampling, especially as we
often did not visually see any fish when we took water samples. Shelton et al. (2019) provides used a paired
eDNA sampling and seine netting analysis demonstrating to demonstrate that eDNA concentrations provide
a smoothed biological signal over space and time. We acknowledge this smoothing effect and emphasize that
in the context of using eDNA for environmental impact assessments, it is preferable to use a survey technique
such as eDNA that integrates signal across a larger spatial and temporal scale.

Many previous papers have commented on the "ecology" of eDNA and the various processes that contribute to eDNA concentrations in environmental samples (e.g., shedding rates, decay rates, transport) (Barnes and 605 Turner 2015). For example, higher concentrations of eDNA could be the result of a greater number (or biomass) of fish present, or increased shedding rates, or decreased decay. Many review papers document the nuances of 607 interpreting eDNA data and we recommend reviewing them for a deeper understanding (see Andruszkiewicz 608 Allan et al. (2020) for a review on shedding and decay rates and Harrison et al. (2019) for a review on transport). Other studies have also documented the relative importance of eDNA transport in streams. 610 Most notably, Tillotson et al. (2018) measured eDNA at four sites with similar discharge rates to the creeks in this study and specifically addressed spatial and temporal resolutions, finding that eDNA concentrations 612 reflect short time- (and therefore length-) scales by comparing peaks in eDNA concentrations to counts of salmon and accumulation by measuring both upstream and downstream sites. The authors found that the 614 sampling site furthest downstream did not accumulate eDNA and that two tributaries feeding into a main 615 channel were additive (Tillotson et al. 2018). For more general models and empirical data documenting 616 transport distances in streams, see Wilcox et al. (2016), Jane et al. (2014), Jerde et al. (2016), Shogren et 617 al. (2016), and Civade et al. (2016). Certainly eDNA concentrations can arise from different scenarios and 618 future work should continue to investigate how to tease apart the nuances of relating eDNA concentrations 619 to fish abundance.

In this study, to assess the impact of a culvert on fish passage, we compare eDNA concentrations upstream and downstream at the same time point in a given creek. The distance between the upstream and downstream sampling was minimal (~60-300 m, average distance of ~150 m). Therefore, we assume that the small differences in spatial and temporal scale between sampling locations is minimal such that the impacts of these various processes will affect the downstream and upstream concentrations equally.

For assessing the impact of construction, we needed to account for differences within the same creek over 626 time (i.e., before and after construction). Because the sampling occurred over a whole year, transport and persistence times may have varied. However, the time series model uses information from the control creeks 628 to understand seasonal trends in eDNA concentrations without needing to link eDNA concentrations to fish abundance. The impact of construction in Padden Creek can be understood by comparing the measured 630 631 eDNA concentration during the time of construction to the expected eDNA concentration in the absence of construction by using information shared from the four other creeks that are not undergoing construction. 632 However, we did correct eDNA concentrations mass/volumeby discharge volume/timeand use a mass flow 633 rate mass/timefor the time series model (see below) given the wide range of discharge over the course of the 634 year. 635 Though eDNA can move downstream with water flow, here, we were measuring if culverts were barriers to 636 fish moving upstream, as we were focused on the impact of culverts on migratory salmon. In our case, we 637 were comparing if downstream stations had higher DNA concentrations than upstream stations as a result 638 of fish being unable to get upstream. This is of course complicated as a result of non-migratory fish, which 639 may be up or downstream and not attempting to pass through the culverts. However, the limited spatial 640 scale between upstream and downstream is such that we can assume the transport would affect upstream and downstream locations in the same way. That is, in the upstream station, some amount of eDNA is 642 coming from upstream of that location into the sampling station and leaving at the same time — in the same way that eDNA would be both entering and exiting the downstream station. Therefore, the relative change 644 between upstream and downstream stations should be the same in terms of eDNA transport. Additionally, at almost every single time point for all creeks and species, the upstream DNA concentration is higher than 646 the downstream DNA concentration. Based on that alone, we do not expect that downstream accumulation 647 of salmonid DNA is occurring to bias our results of whether fish can pass through these culverts. 648 Other studies have documented the relative importance of eDNA transport in streams. Most notably, 649 Tillotson et al. (2018) measured eDNA at four sites with similar discharge rates to the creeks in this 650 study and specifically addressed spatial and temporal resolutions, finding that eDNA concentrations reflect 651 short time- (and therefore length-) scales by comparing peaks in eDNA concentrations to counts of salmon 652 and accumulation by measuring both upstream and downstream sites. The authors found that the sampling 653 site furthest downstream did not accumulate eDNA and that two tributaries feeding into a main channel 654 were additive (Tillotson et al. 2018). For more general models and empirical data documenting transport 655 distances in streams, see Wilcox et al. (2016), Jane et al. (2014), Jerde et al. (2016), Shogren et al. (2016), 656 and Civade et al. (2016).

Finally, it should be noted that Lake Padden, about 1.5 km upstream from the sampling sites, was stocked with cutthroat trout in January 2021, rainbow trout in April and May 2021, and kokanee salmon in May 2021. Given that no sequencing reads in the metabarcoding data are found for O. nerka in May or June after 660 stocking in May, the potential transport of eDNA downstream from Lake Padden to the location of eDNA sampling is expected to be negligible. Given the transport distances documented in the literature and flow 662 rates in Lake Padden, we do not expect the stocking in Lake Padden to affect eDNA concentrations at the 663 sampling locations. 664

Not all culverts are barriers to salmonids 665

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By measuring DNA concentrations of salmonid species above and below culverts on a small spatial scale, 666 we were able to determine how much of a barrier each culvert was (or was not) to fish passage. We found by measuring eDNA concentrations that four of the five creeks sampled did not seem to be major barriers 668 to fish passage. The only creek that was determined to be a Barnes Creek was clearly a very large barrier 669 to fish passage was Barnes Creek, as we only found salmonid DNA in three months of the twelve months 670 of sampling, and those three months had very low concentration of salmonid DNA relative to the other 671 creeks. We note that our sampling occurred only over a single year and future work should monitor culverts 672 for longer time periods, different species, and different environmental conditions. 673 Of the four creeks where salmonid DNA was consistently found, Chuckanut Creekhad the largest discrepancies between DNA concentrations found below and above the barrier at each time point. The 675 culvert in Chuckanut Creek is suspected Within the treatment creek (Padden Creek), the SR-11 culvert 676 did not seem to be a barrier to fish passage and the State of Washington's Department of Transportation is planning to replace it in the near future. The bridge at Portage Creek and the culvert at Squalicum 678 Creek were more recently installed as compared to Padden, Chuckanut, and Barnes Creeks. They also were designated as only partially blocking fish passage, and here we find eDNA results suggest that they were in 680 fact not major barriers to fish passage. Squalicum Creek had the lowest difference between upstream and downstream concentrations across all the surveyed creeks, which corresponds well with the classification 682 that the culvert does not block fish passage. Also, Squalicum Creek is the only creek sampled that has 683 baffles inside the culvert, which should help fish passage. large barrier, while the I-5 culvert clearly was a 684 barrier, demonstrated by the difference in salmonid composition and eDNA mass flow rates over the course 685 of sampling.

Here, we find that instances where culverts designated as barriers were likely not blocking fish passage, while 687 others (Padden I-5 and Barnes Creek) were barriers to fish passage. Importantly, this demonstrates that collecting water samples for eDNA analysis might help to prioritize restoration of culverts suspected to be
barriers to salmonids and provide a new method for post-restoration monitoring to confirm that the barrier
has been corrected and allows for fish passage. Given the large amount of spending and effort required to
replace culverts, this finding is important and emphasizes the potential for new tools for environmental
impact assessments. We note that our sampling occurred only over a short temporal scale and future work
could monitor culverts for longer time periods, different species, and different environmental conditions.

695 Salmonids can quickly recover from a short-term intervention in a creek

The impact of the construction itself on salmonid species demonstrated construction had remarkably minimal effects on salmonid DNA concentrations. The disruption of disconnecting Padden Creek in late August the creek, demolition of the old culvert, installation of the new culvert, and the reconnecting of the creek in early October 2021 during both culvert replacement events showed almost no change in the difference in eDNA concentrations between downstream and upstream sampling sites. The differences in the control creeks between upstream and downstream were often higher than the treatment creek.

The construction timing did coincide with natural life history cycles for the salmon species. In the fall an 702 influx of DNA would be expected not only from adults returning to spawn as they move through the system, 703 but also from the presence of spawning material in the creek and decaying adults that die post reproduction. 704 This may explain a portion of the changes in DNA concentrations found here as the construction timing 705 coincided with run timings of the salmonids, however our time series model accounts for changes in season 706 in attempt to isolate the effects of the culvert and construction. Regardless, the changes between upstream 707 and downstream concentrations were very minor across time points and before and after construction. The post-construction sampling point of the I-5 culvert replacement (only one time point), does show that the 709 composition of salmonid DNA after replacement is now very similar to the two downstream stations, whereas before construction compositions were very different (because the culvert was a barrier). 711

This pattern of minimal disruption and quick recovery was consistent for all four species of salmonids, but
the more abundant species seemed to have a dampened effect (i.e., less overall change)compared to the rarer
species (i. e., O. clarkii was the least impacted and O. nerka was the most impacted). This also corresponds
to species with different life histories and behaviors, and it might be that our most commonly and abundant
species, O. clarkii, was more robust to the intervention because it displays both freshwater resident and
saltwater migrating behaviors.

Our findings here demonstrate that in addition to the value of using eDNA to select culverts to prioritize for replacement, sampling during and after construction can provide important information about the impacts

(or lack of impacts) on salmonids However, we lack the quantitative analysis as the site upstream of SR-11 and downstream of I-5 had no quantifiable cutthroat DNA. More time points would help demonstrate the effect of the culvert replacement. Here we found very minimal effects of both culverts in general and construction, but that one culvert had very minimal effect on salmonid passage while the other culvert had a large effect on salmonid passage. We note that these findings are likely not universal and certainly projects need to monitor comprehensively and quantitatively in order to assess the passability of culverts and impacts of construction.

727 Conclusion

It is notoriously difficult to quantify the environmental impact of discrete human impacts on ecosystems and species. Surveying species and communities by eDNA provides an opportunity for monitoring before, during, 729 and after impacts in a scaleable and cost-effective way. Here, we demonstrate that monthly eDNA sampling before, during, and after an intervention alongside control sites for one year can quantify the environmental 731 impact of replacing a road-culvert. We found that in our treatment creek and control sites, four of the five six barriers did not prohibit salmonid passage and that the culvert replacement. We found that of the 733 two culvert replacements in the treatment creek, one was a barrier and one was not, but both had minimal impacts on the four salmonid species monitored over the course of construction. We also provide a framework 735 in which compositional metabarcoding data can be linked with qPCR data to obtain quantitative estimates of eDNA concentrations of many species. This provides a practical way to utilize the large amount of information 737 from metabarcoding data without needing a unique qPCR assay for every species of interest. Environmental 738 DNA is moving into practice and this study demonstrates how eDNA can be broadly used for environmental impact assessments for a wide range of species and environments. 740

741 Conflict of Interest Statement

The authors declare there are no conflicts of interest.

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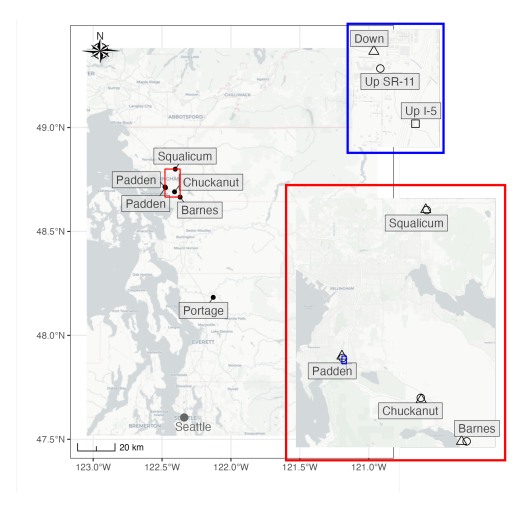
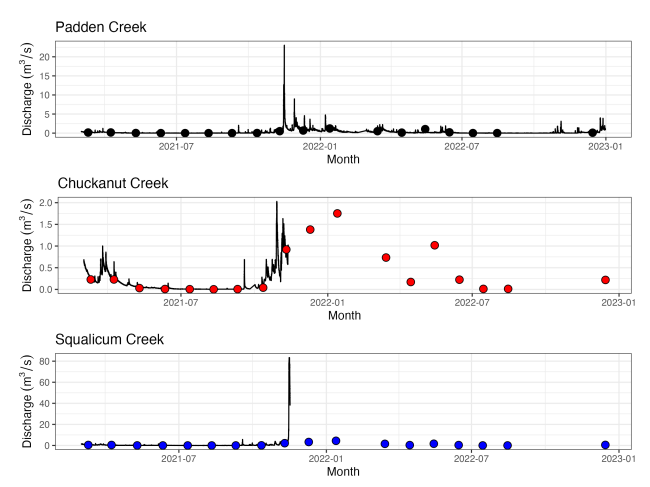


Figure 1: Map of sampling locations near Bellingham, Washington. In the red subset, triangles designate the downstream sampling location and circles designate the upstream sampling location. Padden Creek (the treatment creek where the two culverts were replaced) is shown in the blue subset where downstream is a triangle, upstream of the first culvert (SR-11) is a circle, and upstream of the second culvert (I-5) is a square.



Additionally, in the creek of interest, Padden Creek, rainbow trout (O. mykiss) were stocked in Lake Padden, approximately 1.5 km upstream of the sampling sites. Occasionally, cutthroat trout (O. clarkii) and kokanee salmon (O. nerka) have been stocked in the past as well. During the course of the study, a total of 10Circles designate the day of sampling. For Padden Creek, 000 rainbow trout were stocked in April and May 2021 the nearest 15 minute interval of flow was used. For Chuckanut and 30Squalicum Creeks, 000 kokanee salmon were stocked in May 2021 (Supplemental Figure the correction factor from five years of historical data from Padden Creek was used (see methods section and Supplemental Figures 2 and 3). Despite the stocking of 30,000 kokanee salmon in May in Lake Padden, O. nerka was only detected by metabarcoding in March 2021, August 2021, and then November 2021 through February 2022 (see results below). Importantly, this suggests that Lake Padden is far enough upstream that the eDNA signal at the sampling sites by the culvert is not a result of stocking the lake 1.5 km upstream (see discussion for more information).

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Figure 2: Discharge (m³/s) in Padden (USGS Gauge 12201905), Chuckanut (USGS Gauge 12201700), and Squalicum (USGS Gauge 12204010) Creeks over the course of sampling. Open circles show the days when sampling occurred. Gauges at Chuckanut and Squali84m Creek went offline in November 2021 after a major storm event. Portage Creek and Barnes Creek did not have stream gauges.

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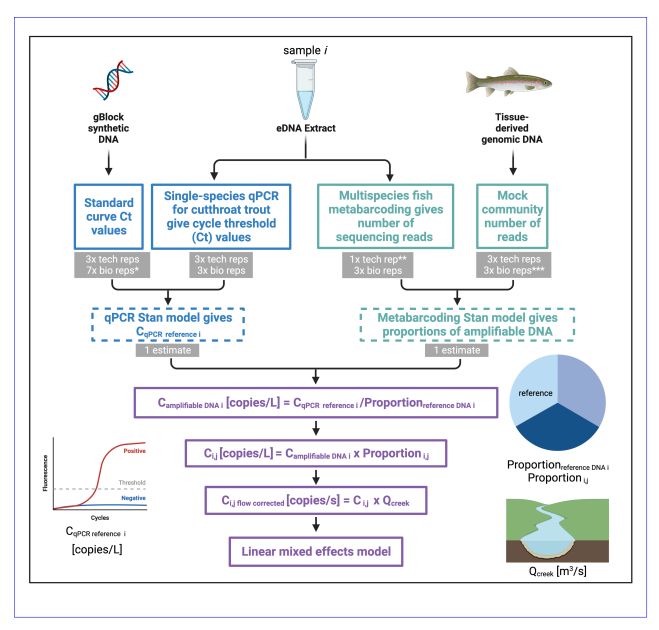


Figure 3: Conceptual figure of different datasets and models used for analyses. * indicates that here, biological replicates are different dilutions of the synthetic gBlock. ** indicates that for most samples, only one technical replicate was sequenced but for one sample per sampling month, three technical replicates were sequenced to check for consistency across replicates. *** indicates that here, the three biological replicates indicate three different mock communities with varying species compositions, but all containing the four salmonids of interest. Created with BioRender.com.

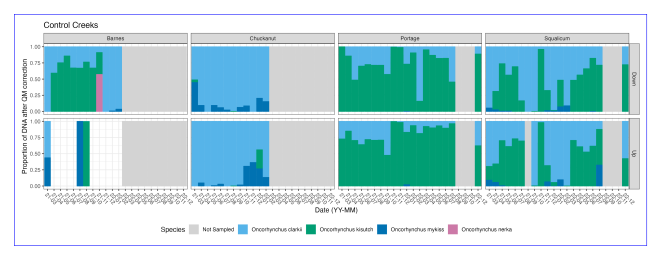


Figure 4: Compositions of salmonid DNA in control creeks as determined by metabarcoding after correction for amplification bias. Note Grey shading denotes time points that no sampling occurred were not sampled (Barnes and Chuckanut Creeks after March 2023 and Squalicum Creek in September 2021 at Squalicum Creek because the creek which was dry). The empty bars in the Barnes upstream sites indicate that no salmonid DNA was found at those time points.

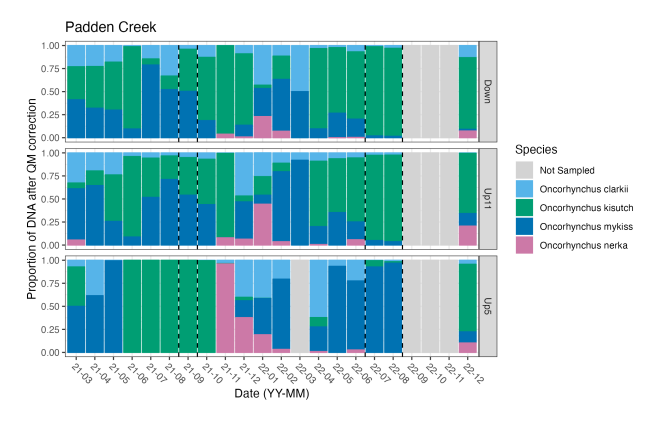


Figure 5: Compositions of salmonid DNA in Padden Creek as determined by metabarcoding after correction for amplification bias. The empty bar in the March 2023 in Up5 indicates that no salmonid DNA was found. The vertical dashed lines indicates the time period in which the culverts were replaced (SR-11 and I-5, respectively).

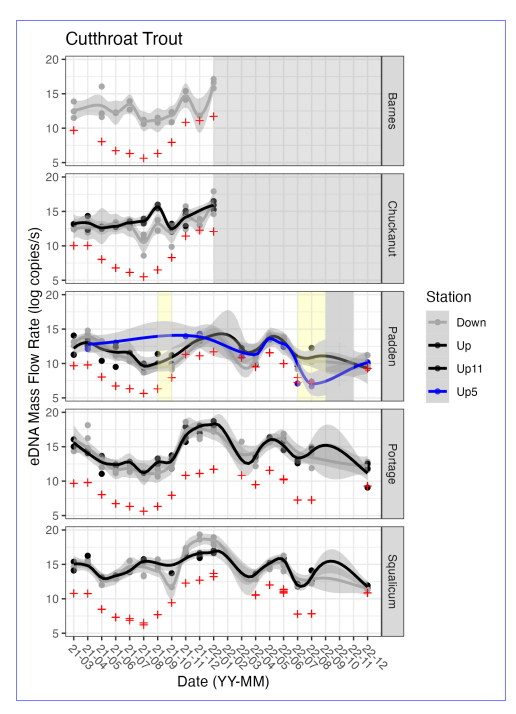


Figure 6: Absolute concentration—mass flow rate (log copies/L of waters) of O. clarkii (cutthroat trout (O. clarkii) as measured by qPCR before after flow correction. Note that Barnes Creek and Chuckanut Creek were not sampled after February 2022. Red crosses show the limit of detection for each species and time point, which changes with flow rate and total volume filtered per sample.

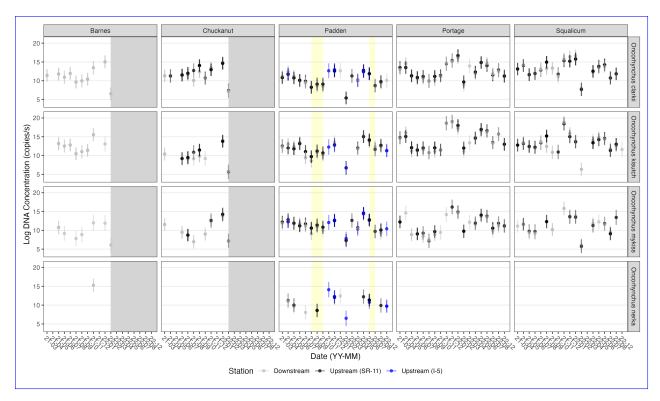


Figure 7: Trends across creeks and across time in mass flow rate (log copies/s) for each of four salmonid species across creeks and across time as estimated by eDNA analysis. Light colored dots are Points represent posterior means derived by expanding the calibrated metabarcoding proportions as described in the main text; darker colored dots are posterior means for the time series linear mixed effects model of and error bars represent the same 95% posterior confidence interval. Colors indicate station upstream (black) or downstream (grey) of the culvert. Padden has an under-road additional sampling site upstream of the second culvert (I-5; blue). 75% and 95% posterior CI plotted for each time point. Grey Yellow shading indicates the time period in which the culvert culverts in the treatment creek (Padden Creek) was were replaced. Grey shading indicates time points that were not sampled (Barnes and Chuckanut after February 2022). Time points with no data had no sequencing reads corresponding to that species or no quantifiable cutthroat DNA by qPCR.

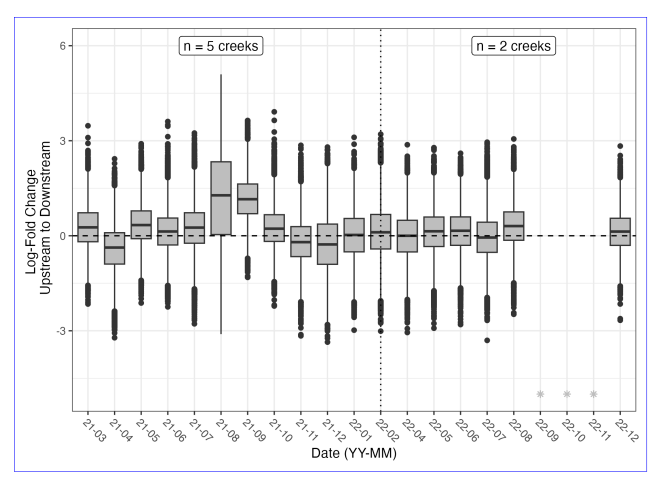


Figure 8: The effect of culvert on salmonid abundance summed across all species and creeks by time. The y-axis shows the difference log-fold change in eDNA mass flow rate (copies/s) between upstream and downstream-concentrations, normalized by downstream-concentration upstream mass flow rate. The box boundaries correspond to the 25th and 75th percentiles; the whiskers correspond to 1.5 times the interquartile range. Here, negative values imply that eDNA concentrations mass flow rates are higher downstream than upstream. Samples with very low eDNA mass flow rates (< 150 copies/s) were removed before plotting to remove extreme proportional values due to large denominators. Grey stars indicate times when no samples were taken.

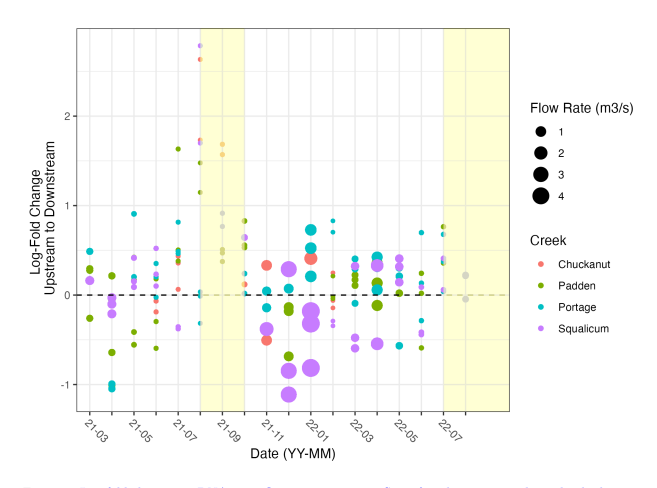


Figure 9: Log-fold change in eDNA mass flow rate over time. Size of circles corresponds to the discharge in each creek at each time point. Color of circles corresponds to each creek. Each creek and time point has up to four circles of the same color for the four salmonid species.

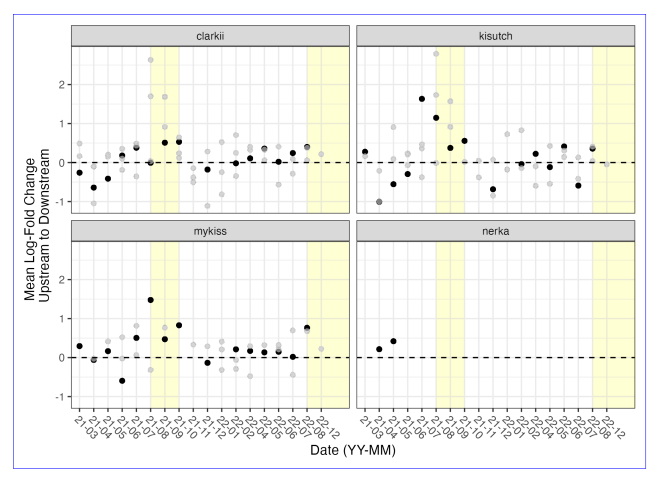


Figure 10: Effect of Construction on Salmonid DNA Concentrations Log-Fold Change in eDNA Flow Rate Upstream to Downstream in Padden Creek. Error bars show 95% confidence intervals of the normalized difference between upstream and downstream DNA concentrations. Grey Yellow shading shows when construction started through the end for each of samplingthe two culverts. Construction ended Grey points show the corresponding log-fold changes in early October 2022. control creeks and black points show Padden Creek. Sockeye/kokanee salmon (O. nerka) was only found in Padden Creek so other creeks are not shown. Samples with very low eDNA mass flow rates (< 150 copies/s) were removed before plotting to remove extreme proportional values due to large denomenators.