Quantifying Impacts of an Environmental Intervention Using

Environmental DNA: Supplemental Text 1

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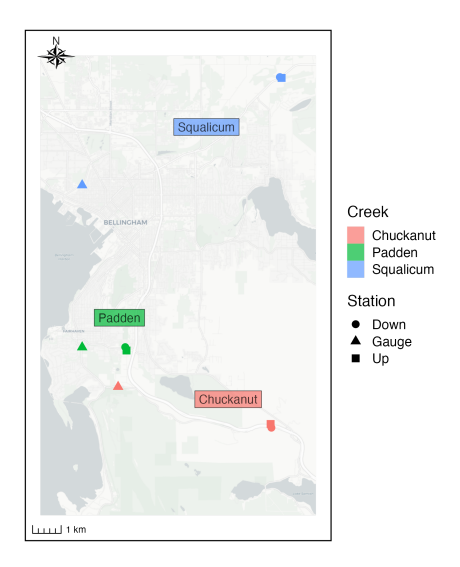
6 Field Sampling

7 Site selection

- 8 The five creeks selected had culverts with various degrees of fish passability: the treatment creek's culvert
- suspected impassible was removed and replaced during the course of the study; two of the four creeks had
- culverts allowing fish passibility, and two had culverts blocking fish passage.
- 11 The average distance between upstream and downstream sampling within a creek was about 160 m; the
- largest distance between downstream and upstream sampling was at Barnes Creek, which was approximately
- 13 330 m, whereas the shortest distance between sampling was at Squalicum Creek at approximately 66 m.

14 Flow variability at sites

- Over the course of the year, flow within each creek varied. USGS flow gauges were located in three of the
- is five creeks, relatively nearby to the sampling locations (Supplementary Figure 1). The closest gauge to
- sampling locations was Padden Creek (~1.5 km); the gauge at Chuckanut Creek was ~5.5 km and the gauge
- at Squalicum Creek was ~7.9 km away (calculated using the Haversine distance in R).
- 19 The flow meters at Squalicum Creek and Chuckanut Creek were offline from November 2021 for the remainder
- 20 of the sampling period. The highest discharge seen during the course of the study from January to November
- ²¹ 2021 occurred in November 2021 at Squalicum Creek. The mean discharge in each creek was: 0.42 m³/s in
- ²² Padden, 0.29 m³/s in Chuckanut, and 1.14 m³/s in Squalicum Creek. The lowest discharge registered by the



 $Figure \ 1: \ Location \ of \ flow \ gauges \ compared \ to \ sampling \ locations \ for \ Chuckanut, \ Padden, \ and \ Squalicum \ Creeks.$

- $_{23}$ flow meters is $0.0028 \text{ m}^3/\text{s}$, which occurred 8.5%, 1.6%, and 0.78% of the time in Padden, Chuckanut, and
- 24 Squalicum, respectively.
- 25 Due to the lack of flow meter in Squalicum and Chuckanut Creeks from November 2021 to February 2022, we
- used historical data from the three flow gauges to calculate the average discharge for each day of the year
- 27 from about 2015-2021 (Supplementary Figure 2). We then used the value for the day of the year that we
- sampled in either 2021 or 2022 when the gauges were offline. For consistency, we also did this at Padden
- ²⁹ Creek despite the gauge there being online for the entire sampling period.

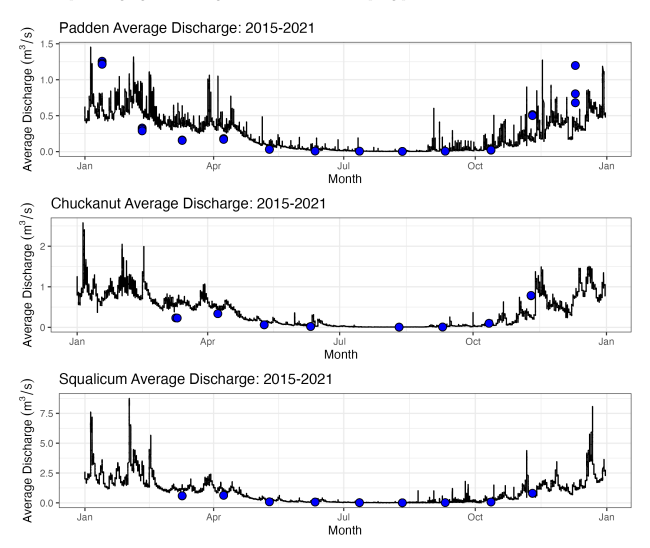


Figure 2: Discharge in creeks with USGS gauges. Note that Chuckanut and Squalicum went offline around November 2021. Lines show the value for each day averaged over 2015-2021. Blue dots show the time of sampling during the course of this study.

30 Stocking in Lake Padden

Padden Lake has historically been stocked with hatchery fish by the Washington Department of Fish and Wildlife (Supplementary Figure 3). During the course of the study, a total of 10,000 rainbow trout were stocked in April and May 2021, 30,000 kokanee salmon were stocked in May 2021, and 10,000 cutthroat trout were stocked in January 2021 two months before sampling began in March. However, Lake Padden is 1.5 km upstream of the sampling locations in Padden Creek and we do not expect to see an increased signal due to the flow from the lake after stocking events.

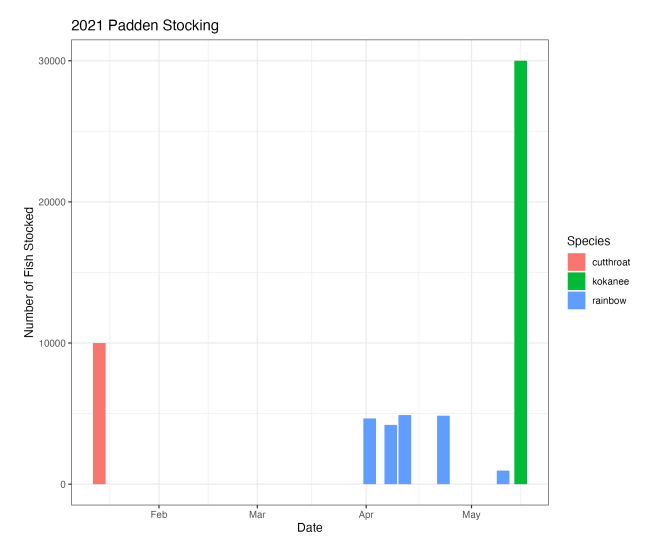


Figure 3: Stocking of Lake Padden in 2021.

37 Water Sampling

- Water samples were collected using Smith Root's eDNA Backpack (Thomas et al. 2018), a portable pumping-
- and-filtering setup set to filter at 1 L/min at 82.7 kPa (12 psi). For most months, a trident sampler was used
- 40 to collect all 3 biological replicates at the exact same time, for a total sampling time of about 5 minutes.
- 41 Otherwise, the three replicates were collected consecutively, for a total sampling time of about 15 minutes.
- 42 The backpack also monitored pressure and flow rate over the course of sampling. The backpack was set to
- 43 have a target flow rate of 1 L/min and a max pressure of 12 psi. Downstream sites were always sampled
- before upstream sites to ensure no potential DNA was introduced into the stream before sampling. In some
- months, less than 2 L of water was filtered due to clogging (Supplemental Figure 4).

46 Laboratory Processing

47 DNA Extractions

- 48 We followed a protocol developed for extracting DNA off the self-preserving Smith Root filters (Thomas et
- ⁴⁹ al. 2019). Filters were removed from their housing with sterile tweezers and cut in half using sterile razor
- 50 blades. One half was archived and the other half was used for extraction. The extraction was performed
- using the Qiagen DNeasy Blood and Tissue Kit (Qiagen, USA) with minor modifications, including adding a
- ⁵² Qiashredder column (Qiagen, USA).

53 PCR Amplification

- PCR reactions included 10 μ L of 5X Platinum ii Buffer, 0.4 μ L of Platinum ii Taq, 1.25 μ L of 8 mM dNTPS,
- ⁵⁵ 1.25 μ L of 10 μ M F primer, 1.25 μ L of 10 μ M R primer, 5 μ L of template, and 30.85 μ L of molecular grade
- water, for a total reaction volume of 50 μ L. Cycling conditions were as follows: 95°C for 2 min, 35 cycles of
- ⁵⁷ 95°C for 30 sec, 60°C for 30 sec, 72°C for 30 sec, followed by a final extension of 72°C for 5 min.
- For indexing, 10 ng of PCR product was used as template in a final volume of 11.25 μ L For samples with
- concentrations less than 0.88 ng/ μ L, 11.25 μ L was added despite being less than 10 ng of amplicon. Each
- sample received a unique index; Nextera index sets A and B were used in combination to avoid using the
- same index for more than one sample on a single sequencing run. The PCR reaction included the 11.25 µL of
- template, 12.5 μ L of Kapa HiFi MMX (Roche, USA), and 1.25 μ L of indexed primer. Cycling conditions
- were as follows: 95°C for 5 min, 8 cycles of 98°C for 20 sec, 56°C for 30 sec, 72°C for 3 min, and a final
- extension of 72°C for 5 min.

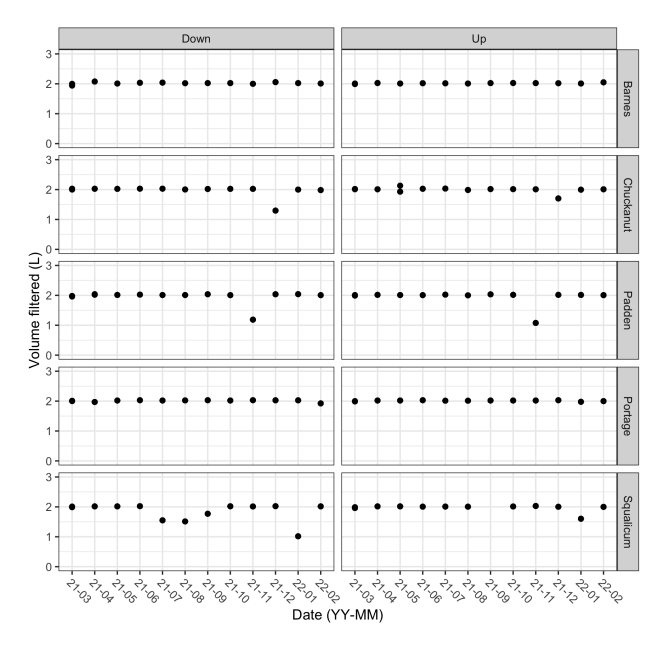


Figure 4: Volume filtered (L) for each sample. Note that Squalicum upstream in September 2021 was completely dry and therefore no water was filtered.

65 Species Specific qPCR

- 66 Each DNA sample was run in triplicate using Gene Expression Mastermix (ThermoFisher, USA), a final
- concentration of 0.375 μ M F primer, 0.375 μ M R primer, and 0.105 μ M probe, as well as 1X EXO-IPC mix,
- 68 1X EXO-IPC DNA, 3.5 μ L of template for a final reaction volume of 12 μ L. The EXO-IPC mix includes the
- ₆₉ primers and probe for the EXI-IPC DNA, with the probe having a VIC reporter, allowing it to be multiplexed
- with the *O. clarkii* assay, which has a FAM reporter.
- 71 Thermocycling was as follows: 50°C for 2 min, 95°C for 10 min, followed by 45 cycles of 95°C for 15 sec, 60°C
- ₇₂ for 1 min. The cycle threshold (Ct) value determined for the EXO-IPC assay from the NTC was compared to
- ₇₃ the Ct value for the EXO-IPC assay in each of the environmental samples. If the Ct value was >0.5 Ct values
- ₇₄ from the mean Ct for the NTCs, the sample was deemed inhibited and diluted 1:10 and re-assayed until
- ⁷⁵ the Ct value fell within the accepted range (Supplementary Figure 5). After converting Ct values to DNA
- concentrations using the standard curve (see below), the concentration was multiplied by the dilution factor.

77 Bioinformatics Processing

- Primers were removed with cutadapt (Martin 2011) and then reads were de-noised, filtered, merged, and
- ⁷⁹ ASVs were generated using dada2 (Callahan et al. 2016). For each MiSeq run, the trimming lengths were
- 80 determined by visually assessing the quality score plots. After ASVs were generated, taxonomy was assigned
- using the "classify" function in the insect package in R using the classifier published by the authors of the
- package (Wilkinson et al. 2018).

83 Quality Controls

- Positive controls were included on each sequencing run to monitor for cross contamination that might have
- 85 occurred in the laboratory or due to "tag jumping". With 13 MiSeq runs, we included one sample of kangaroo
- tissue on each run and then measured how many reads of kangaroo were found in environmental samples and
- ₈₇ how many reads of non-kangaroo were found in kangaroo samples (Supplementary Figure 6).
- 88 We can also check to make sure that no reads assigning to kangaroo were in the environmental samples. We
- ₈₉ processed X environmental samples and only 2 had any reads assigned to kangaroo (Supplementary Figure 7).

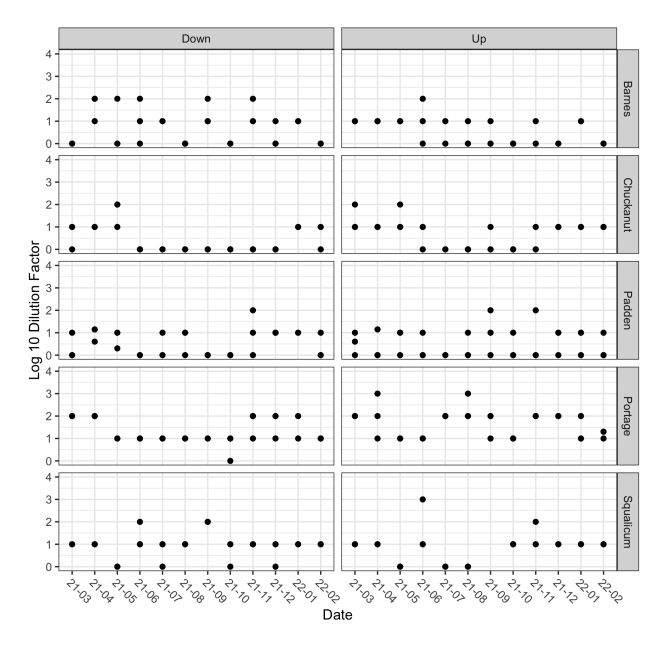


Figure 5: Dilution factor required to remove inhibition as measured by an internal positive control (IPC) qPCR assay.

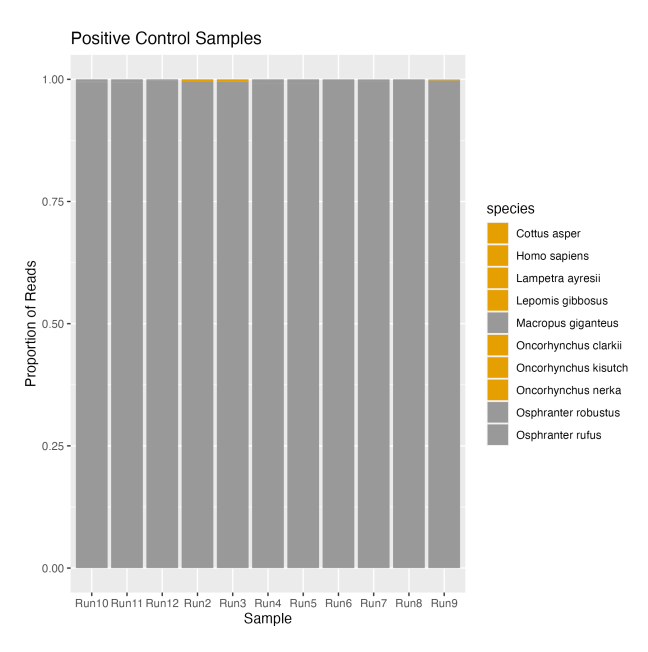


Figure 6: Proporiton of annotated reads found in positive controls. Grey colors are the three species of kangaroo used for positive controls and are what should be in each sample. Orange species should not be in the positive controls and indicate low level contamination from environmental samples.

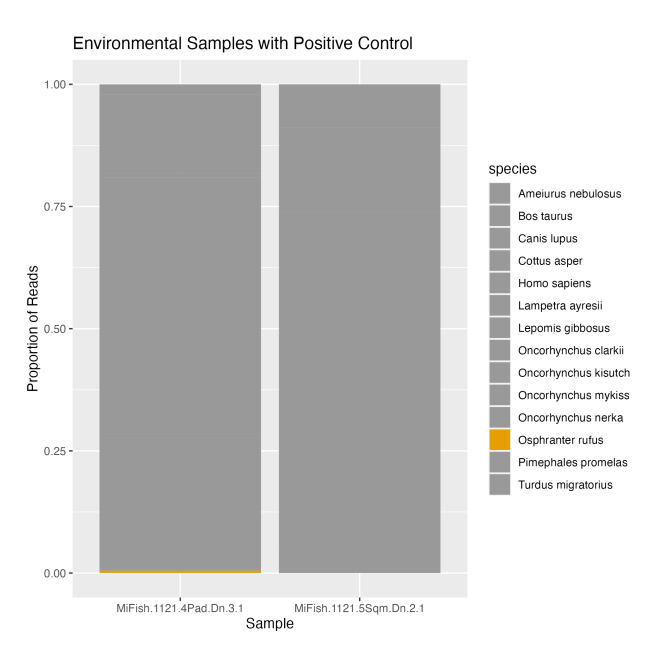


Figure 7: Proporition of annotated reads found in environmental samples with positive control. Grey colors are non-kangaroo reads and therefore and are what should be in each sample. Orange species are kangaroo reads and therefore should not be in the environmental samples and indicate low level contamination from positive controls.

90 Annotation

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(Supplementary Figure 7).

- 91 We first used a tree-based annotation method (insect package) and then followed up with a BLAST search
- 92 for all ASVs that were not annotated to species level by insect.

93 Correcting metabarcoding data for amplification bias

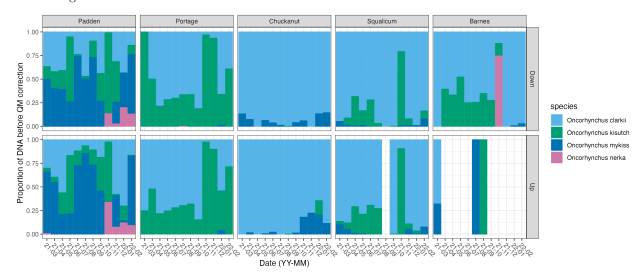
Using our six mock communities (three different taxa compositions x two proportions [even and skewed]), we can first check how well the quantitative metabarcoding model corrects for amplification bias. In one case, we consider the even mock communities as the mock community data and the skewed mock communities as unknown. We can then re-create what the model believes to be the original starting proportions of the skewed mock community given the proportions of reads found in the skewed mock communities and the proportion of DNA as compared to the proportion of reads found in the even mock communities. We can

We can also check how well the calibration is working by comparing the alpha values by using different

subsets of mock community data as true and unknown (Supplementary Figure 8).

also do the same treating the skewed mock communities as known and even mock communities as unknown

We can then use the mock communities to correct the data from the MiSeq (Supplementary Figure 9) to account for the different alpha values (Supplementary Figure 8). The corrected results are shown in the main text as Figure 3.



Effects of culverts In the main text, we show the effect of culverts averaged over creeks and species. Here, we show them separate (Supplementary Figure 10). The y-axis scale changes for each creek, demonstrating

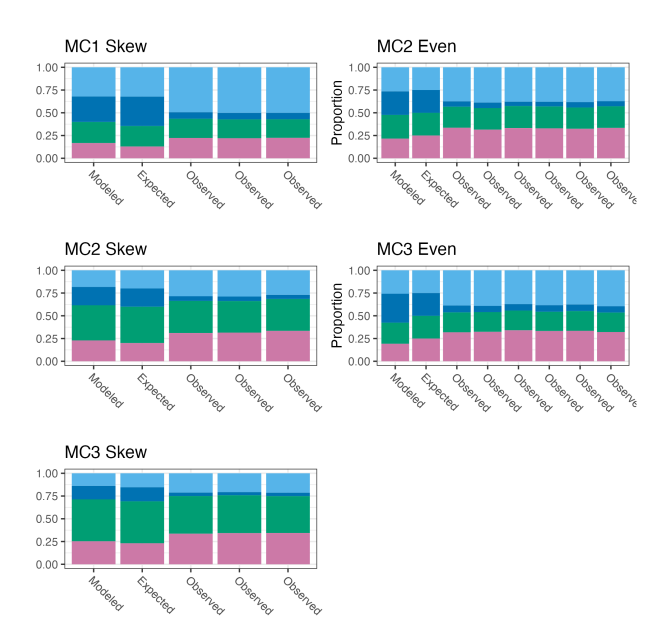


Figure 8: Intercalibration of mock communities used to correct environmental samples for amplification bias.

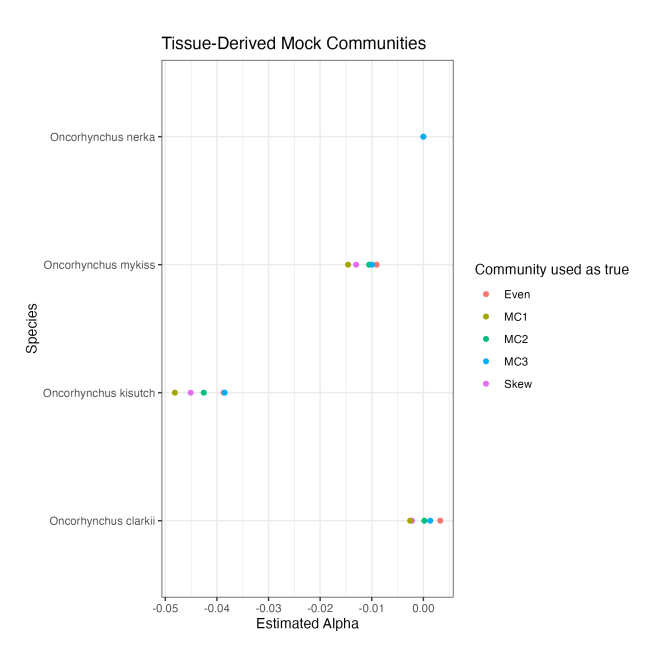


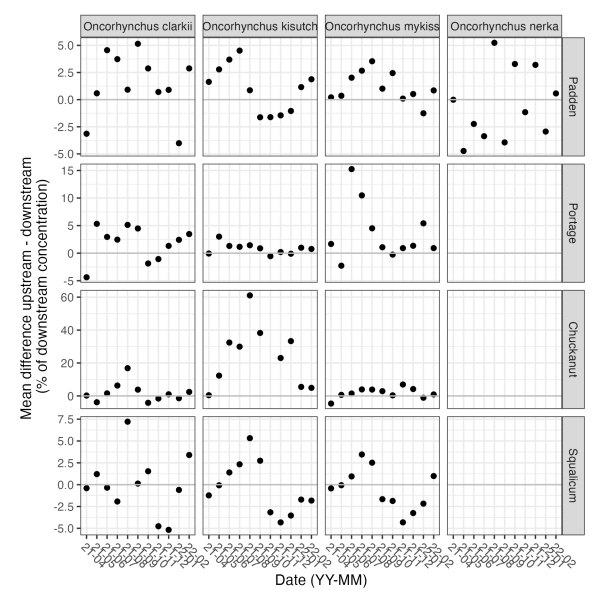
Figure 9: Estimated alpha values of salmonid species with different calibrations of the mock communities. Each color represents a different subset of mock community data treated as 'true' to calibrate the remainder of the mock community data.

that some culverts have a larger effect on the difference between upstream and downstream than others.

Namely, O. kisutch in Chuckanut Creek sees the largest percent difference. There also seems to be a slight

effect of season on the culverts, where summer months often have the most highest difference between

upstream and downstream (i.e., higher eDNA concentrations upstream).



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115 References

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Callahan, B. J., P. J. McMurdie, M. J. Rosen, A. W. Han, A. J. A. Johnson, and S. P. Holmes. 2016. DADA2: High resolution sample inference from illumina amplicon data. Nature methods 13:581–583.

Martin, M. 2011. Cutadapt removes adapter sequences from high-throughput sequencing reads. EMBnet.journal 17:10.

Thomas, A. C., J. Howard, P. L. Nguyen, T. A. Seimon, and C. S. Goldberg. 2018. ANDe TM: A fully

- integrated environmental DNA sampling system. Methods in Ecology and Evolution 9:13791385.
- Thomas, A. C., P. L. Nguyen, J. Howard, and C. S. Goldberg. 2019. A self-preserving, partially biodegradable eDNA filter. Methods in Ecology and Evolution 10:1136–1141.
- Wilkinson, S. P., S. K. Davy, M. Bunce, and M. Stat. 2018. Taxonomic identification of environmental DNA
 with informatic sequence classification trees.