



Helbing Laboratory eDNA Technical Bulletin

All eDNA tools are validated through a rigorous multi-step evaluation protocol that includes tests of DNA target specificity and amplification sensitivity¹⁻³.

General eDNA Assay Information

Target Species: Sockeye Salmon (*Oncorhynchus nerka*) eDNA qPCR Tool: eONNE2 Gene Target: MT-CYB
Species Code: te-ONNE eDNA qPCR Format: TaqMan Published in:

eDNA Assay Sensitivity Test Summary using gBlocks™ Synthetic DNA

LOD 0.5 95% CI 0.4-0.9 Copies/Rxn LOQ 1.9 95% CI 1.4-3.2 Copies/Rxn LOB 0 hits/8
LOQ_{continuous} 4 Copies/Rxn

Binomial-Poisson model for 8 technical replicates determined using eLowQuant R code⁴. When the LOQ < LOD, use the LOD for the LOQ. Enzyme: QIAcuity

eDNA Assay Specificity Test Information

Each qPCR reaction in the specificity assay contained 10 picograms of voucher target gDNA (n=25 technical replicates)

Species	Common Name (<i>Species</i>)	Detection	# Voucher		Sample Sources/Locations
			Specimens		
ma-HOSA	Human (<i>Homo sapiens</i>)	No	1		Netherlands
te-ONCL	Cutthroat Trout (<i>Oncorhynchus clarkii</i>)	No	1		Alberta or British Columbia
te-ONGO	Pink Salmon (<i>Oncorhynchus gorbuscha</i>)	No	1		British Columbia
te-ONKE	Chum Salmon (<i>Oncorhynchus keta</i>)	No	1		British Columbia
te-ONKI	Coho Salmon (<i>Oncorhynchus kisutch</i>)	No	1		British Columbia
te-ONMY	Rainbow Trout (<i>Oncorhynchus mykiss</i>)	No	1		British Columbia
te-ONNE	Sockeye Salmon (<i>Oncorhynchus nerka</i>)	Yes	1		British Columbia
te-ONTS	Chinook Salmon (<i>Oncorhynchus tshawytscha</i>)	No	1		British Columbia
te-SASA	Atlantic Salmon (<i>Salmo salar</i>)	No	1		Nova Scotia

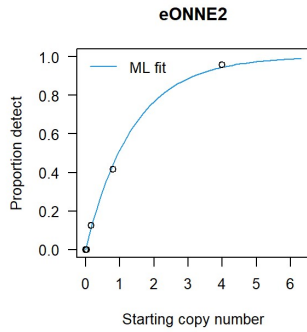
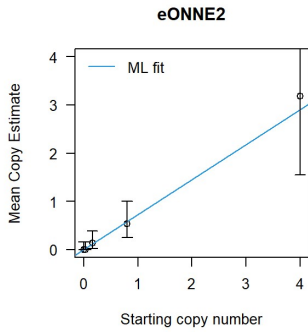
References

- Hobbs, J, Adams, IT, Round, JM, Goldberg, CS, Allison, MJ, Bergman, LC, Mirabzadeh, A, Allen, H, Helbing, CC (2020) Revising the range of Rocky Mountain tailed frog, *Ascaphus montanus*, in British Columbia, Canada, using environmental DNA methods. *Environmental DNA*. 2020; 2: 350-361. <https://doi.org/10.1002/edn3.82>
- Hobbs, J, Round, JM, Allison, MJ, Helbing, CC (2019) Expansion of the known distribution of the coastal tailed frog, *Ascaphus truei*, in British Columbia, Canada, using robust eDNA detection methods. *PLOS ONE* 14(3): e0213849. <https://doi.org/10.1371/journal.pone.0213849>
- Langlois, VS, Allison, MJ, Bergman, LC, To, TA, and Helbing, CC (2020) The need for robust qPCR-based eDNA detection assays in environmental monitoring and risk assessments. *Environmental DNA*, 3: 519-527. doi: 10.1002/edn3.164
- Lesperance, M, Allison, MJ, Bergman, LC, Hocking, MD, and Helbing, CC (2021) A statistical model for calibration and computation of detection and quantification limits for low copy number environmental DNA samples. *Environmental DNA*, 00: 1-12. doi: 10.1002/edn3.220



eDNA Assay Sensitivity Test Details using gBlocks™ synthetic DNA

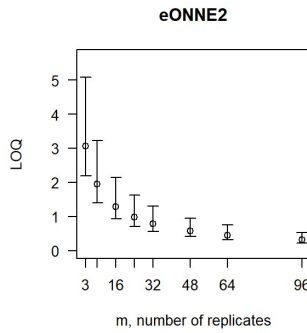
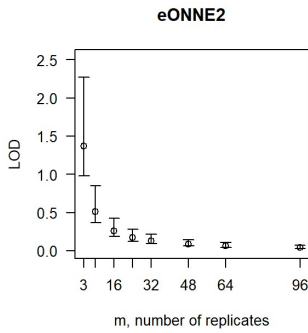
To calculate tables for different numbers of replicates, raw csv data files can be accessed here:
<https://onlineacademiccommunity.uvic.ca/helbinglab/edna/>



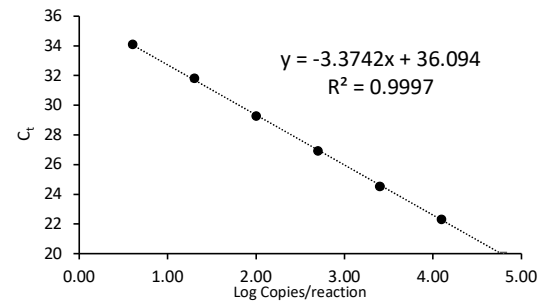
From 8 Technical Replicates

# Detects	# Copies	SE
0	0.00	0.00
1	0.18	0.19
2	0.40	0.29
3	0.65	0.40
4	0.95	0.52
5	1.35	0.68
6	1.91	0.93
7	2.86	1.41

Determined using eLowQuant R code⁴.



Applied to reactions with 100% positive hits



Efficiency 98%

Binomial-Poisson model: no intercept
Determined using eLowQuant R code⁴.
Based on a 2 µL DNA input in a total 15 µL reaction

Field Sample Validation

Known
Sample Type Presence # Samples Detected Location

Abbreviations

95% CI	95% Confidence interval	LOQ	Limit of quantification
eDNA	Environmental DNA	MT-CYB	Mitochondrial cytochrome B gene
gDNA	Total genomic DNA extracted from voucher specimen	NTC	qPCR no template control
LOB	Limit of blank	qPCR	Quantitative real-time polymerase chain reaction
LOD	Limit of detection	SE	Standard error