THE IMPACT OF SUMMER DROUGHT ON THE NITRIFYING MICROBIOME OF REWETTED FEN PEATLANDS

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In partial fulfillment for the degree of Master of Science in Landscape Ecology and Nature Conservation

University of Greifswald, 2024

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 To-do:

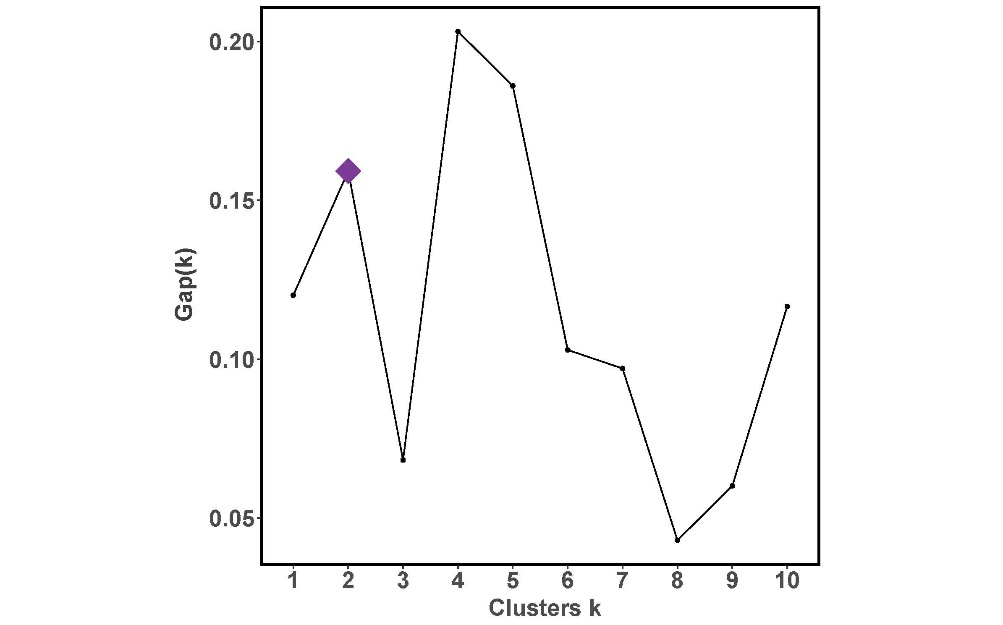
* Manuscript pre-print:
  + Make metabolic pathway map
  + Add rRNA for CW and PW to megafigure
  + Incorporate Haitao’s comments
  + Add redox potential data
  + Look at SEED pathways
* Thesis:
  + Remake clustering figures
  + Write short introduction
  + Edit plasmid cultivation methods
  + Write clustering method details + put in context w/ literature
  + Write phylogenetic tree assembly details
  + Write RT/qPCR details
  + Write brief conclusion

**I. Introduction**

**II. Manuscript pre-print: “Ammonia oxidizing archaea and bacteria respond dynamically to drought in rewetted fen peatlands” (Burns et al., under review)**

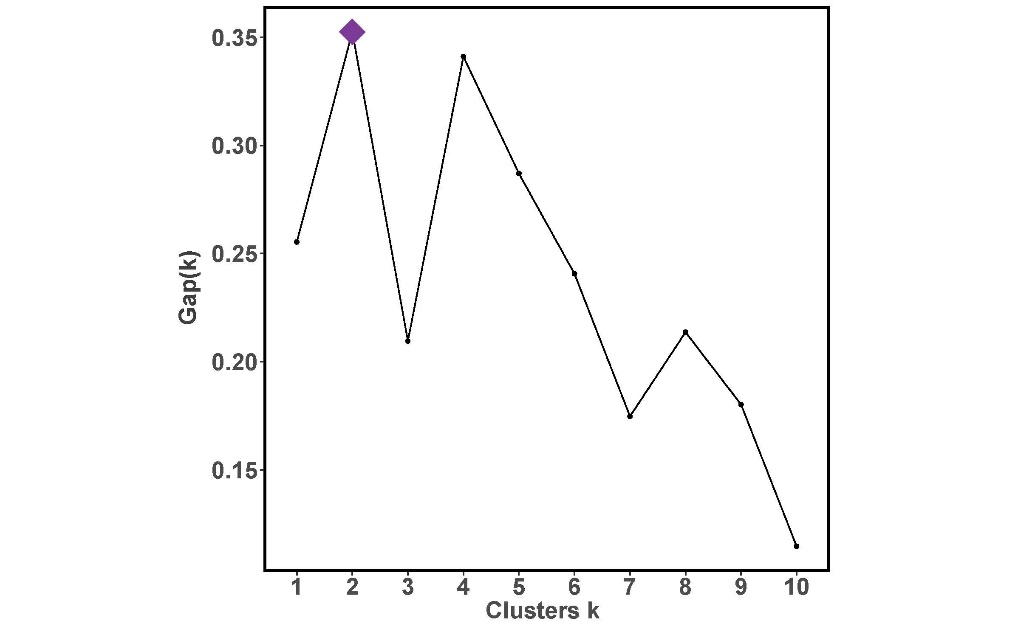
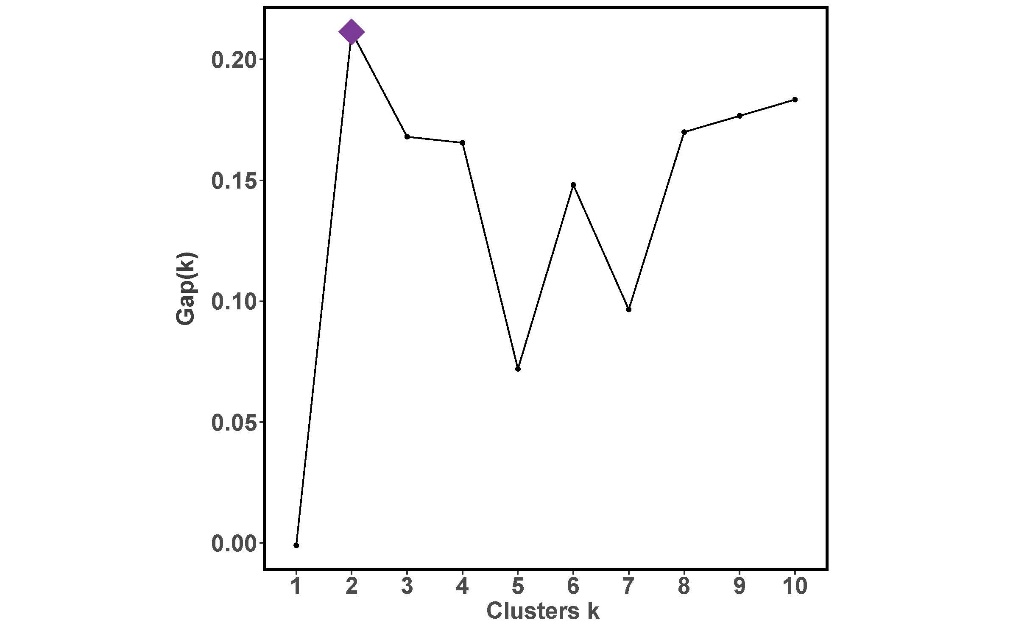
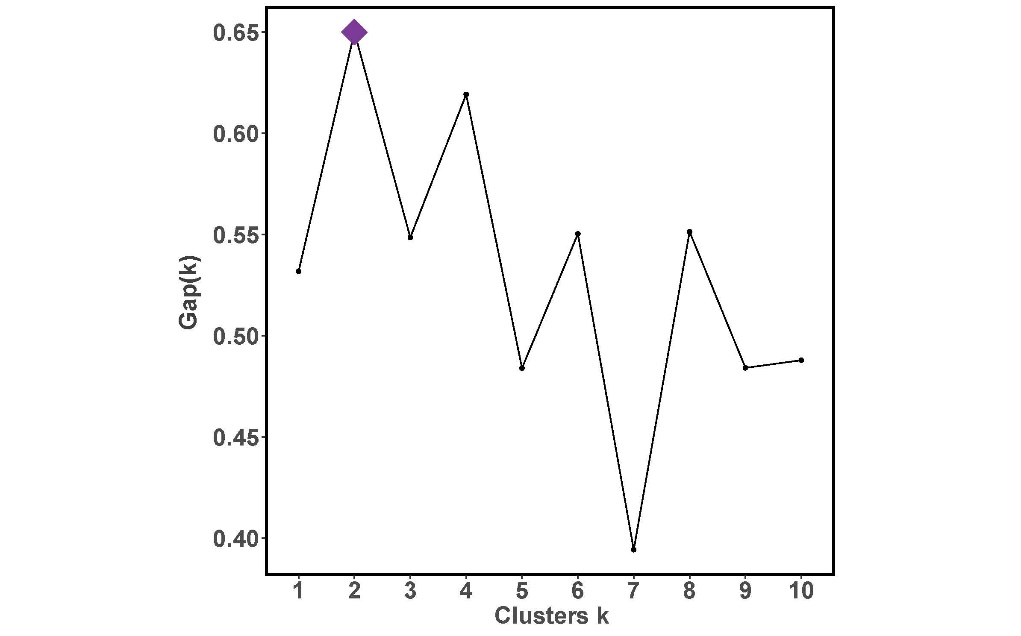
 \*\* there are 6 figures in the publication \*\*

**III. Appendix I: Clustering Method for Drought Definition**



*Figure 7.* Results of the ClusGap function (K.max = 10, B = 500) on each site’s water table depth values . K = 2 was chosen as the optimal cluster number for each site based on the gap statistic (Gapₖ). This was evident for PW (a), PD (b) and CW (c), where k = 2 also had the maximum Gapₖ; for CD (d), k = 2 was selected heuristically due to the sharp decrease in Gapₖ for k = 3 (despite the subsequent increase when k = 4). Therefore, k = 2 was selected to minimize the instability in the Gapₖ metric.

**CDW AOB**



**CWW AOB**

**d) AOB**

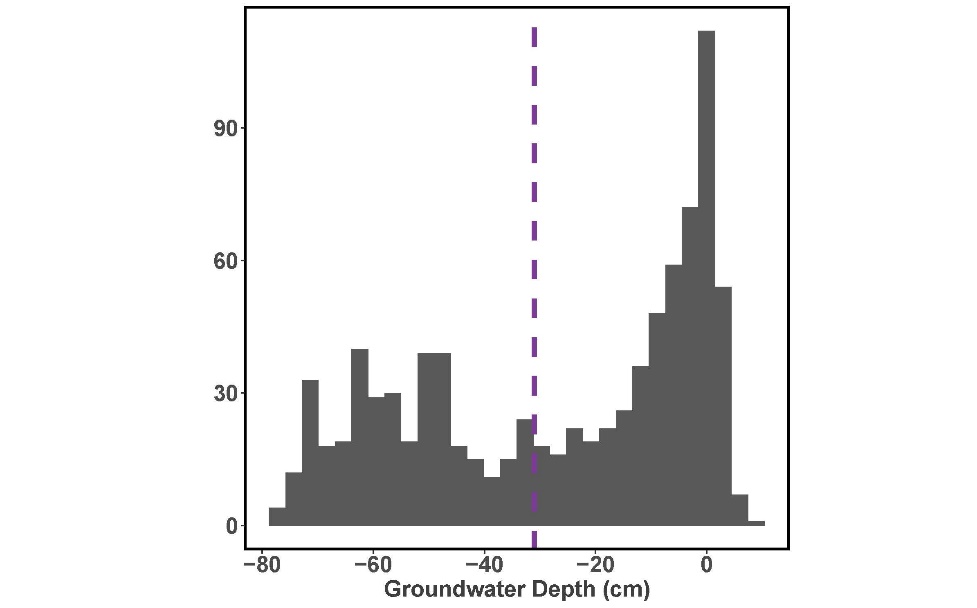
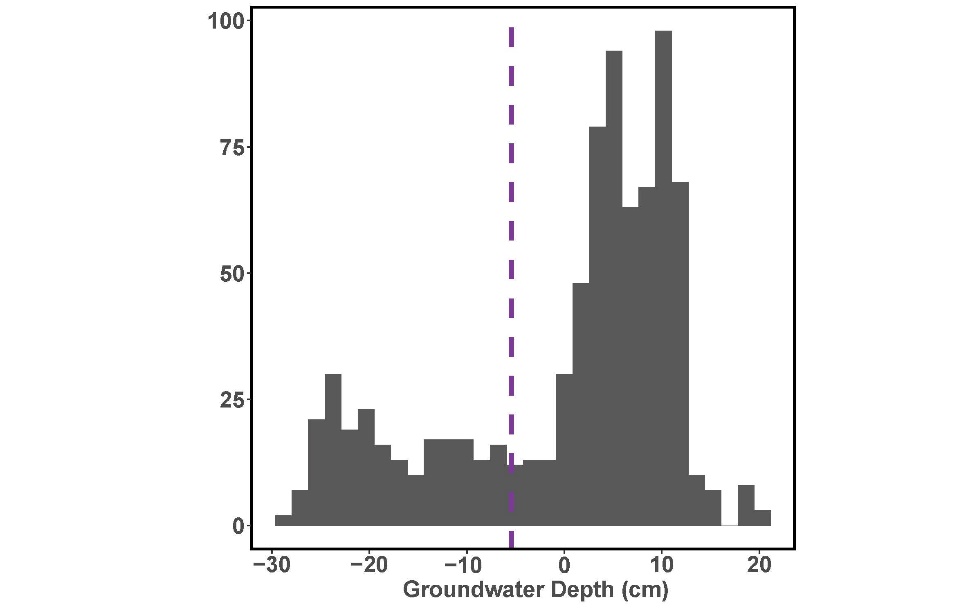
**b) AOB**

**PD AOB**

**PWW AOB**

**c) AOB**

**a) AOB**

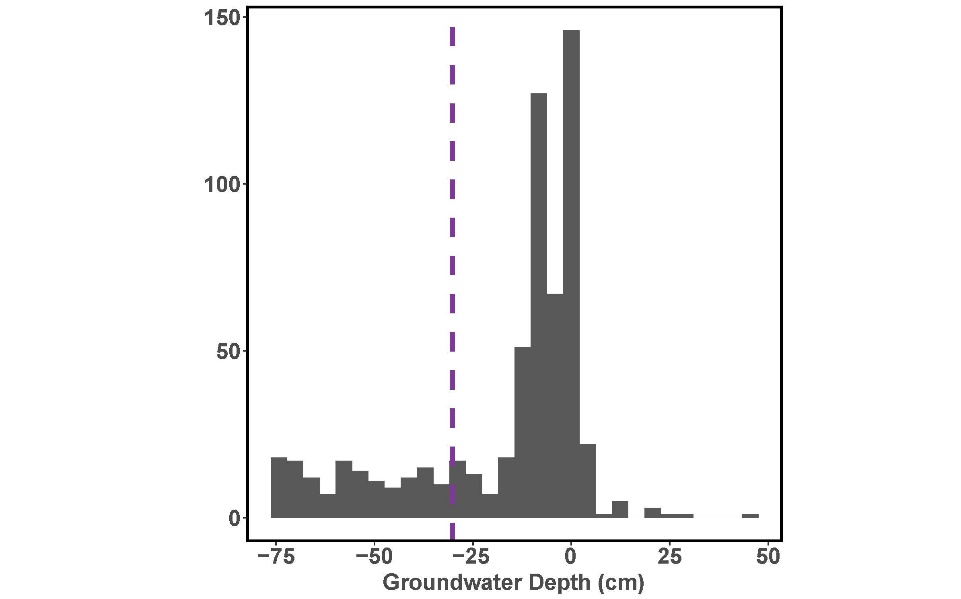


**b) AOB**

**PD AOB**

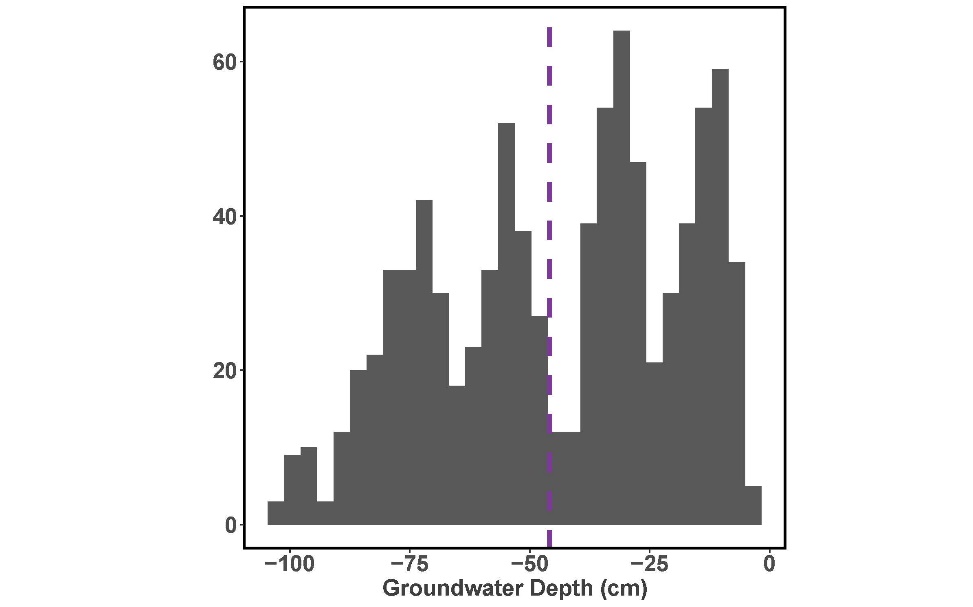
**PWW AOB**

**a) AOB**



**CWW AOB**

**c) AOB**



**CD AOB**

**d) AOB**

*Figure 8.* Histograms of water table values in each site. The k-means cluster-identified drought threshold for each site included as follows: (a) PW drought threshold -5.45 cm; (b) PD, -31.05 cm; (c) CW, -30.24 cm; (d) CD, -45.88 cm.

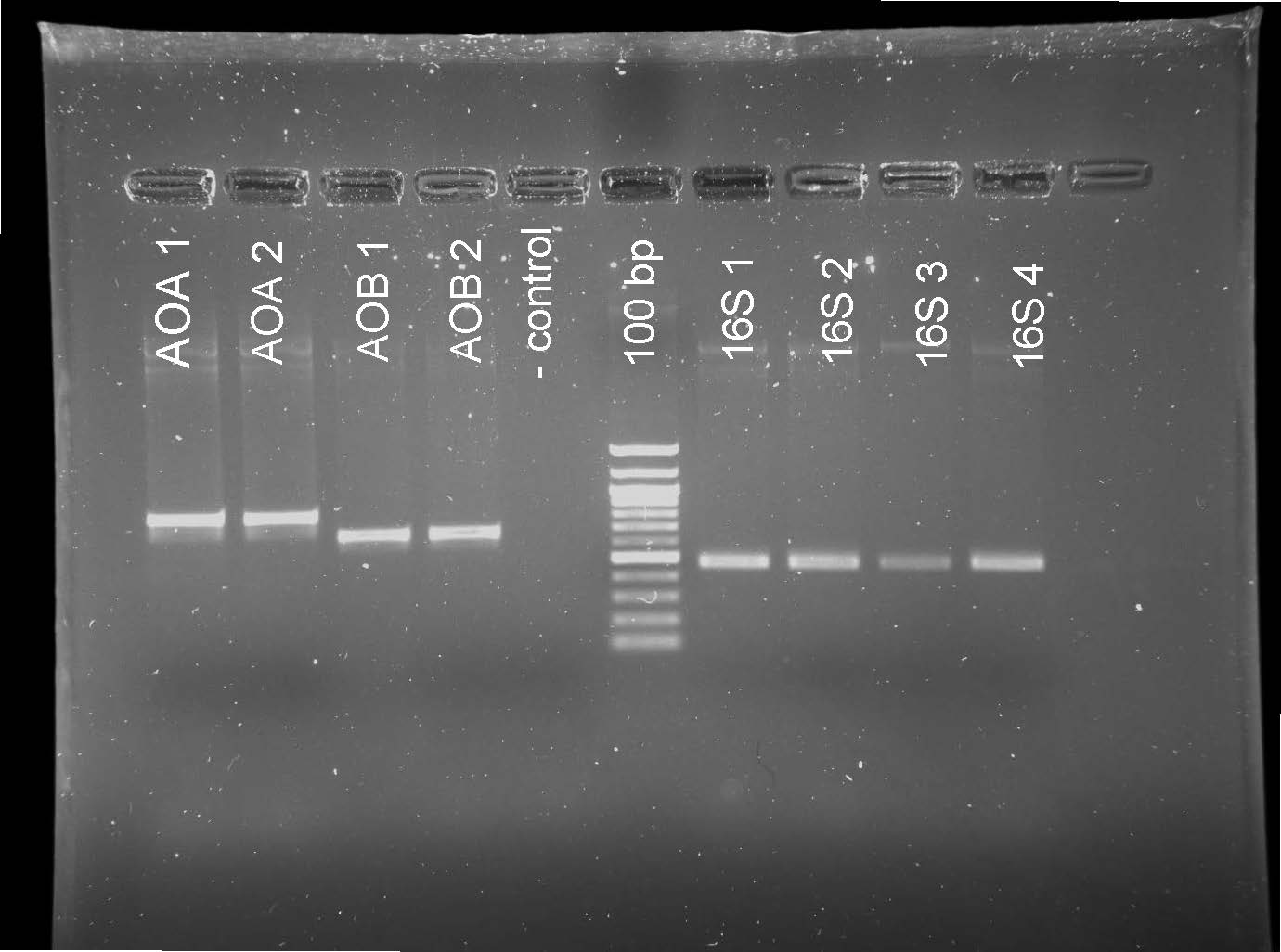
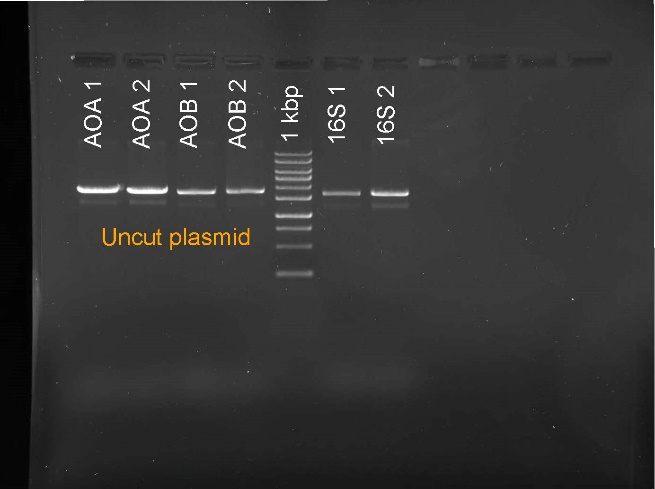
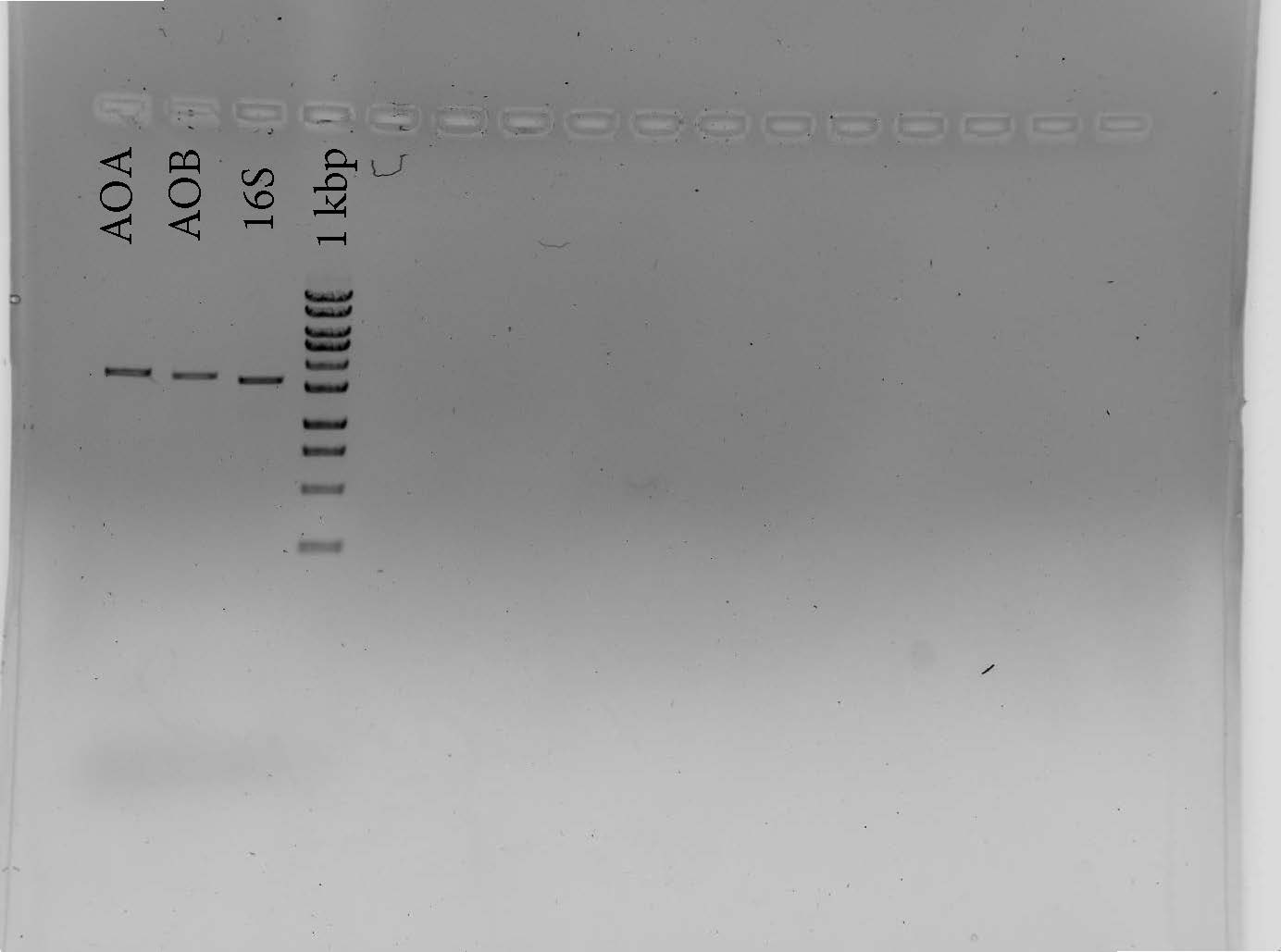
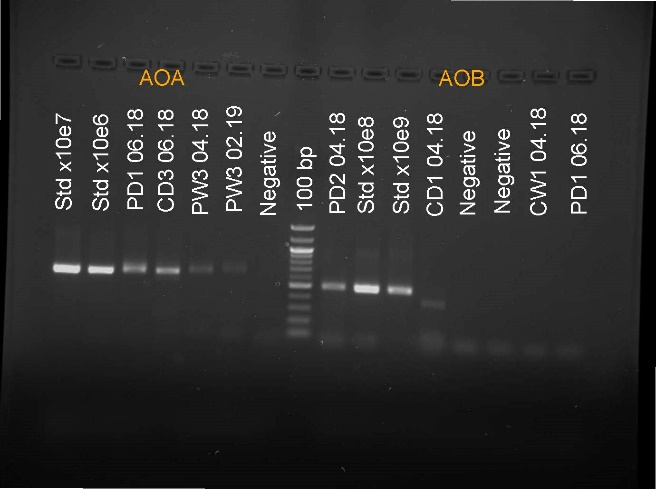
**IV. Appendix II: AOA and AOB Standard Cultivation**

The purified cultures chosen to create standards for downstream qPCR analysis were the ammonia-oxidizing bacteria *Nitrosopira multiformis* [1]*,* ammonia oxidizing archaea *Nitrosophaera viennensis* [2], and *Escherichia coli* [citation needed?]. First, the target genes were amplified via PCR with the following 25 ul reaction: 0.75 ul GoTaq® DNA Polymerase (Promega, Wisconsin, USA), 5 ul 5x GoTaq® Reaction Buffer (Promega), 2.5 ul MgCl₂, 0.5 ul DNTPs, 13.375 ul H₂O, 0.125 ul each forward and reverse primer, and 1 ul DNA template. The archaeal amoA gene was amplified from *N. multiformis* with the primer pair camoA-19F [2, 3]/tamoA-629R [4], using a PCR cycle with a 5 minute initialization phase at 95°C, followed by 35 rounds of a 45 second denaturation stage at 95°C, a 45 second annealing stage at 55°C, and a 1 minute elongation phase at 72°C, before a final 10 minute finalization stage at 72°C. The bacterial amoA gene was amplified from *N. viennensis* using the primer pair amoA-1F/2R [5], and the 16S rRNA gene was amplified from *E. Coli* using the 515F [6] /806R [7] primers . Both the AOB and 16S rRNA templates were amplified using the following PCR protocol: a 2 minute initialization stage at 95°C, then 35 rounds of a 30s elongation stage at 95°C, a 30s annealing stage at 55°C, and a 45s elongation stage at 72°C, before a 5 minute finalization stage at 72°C.

The lengths of the resulting PCR products of the genes of interest were then verified via gel electrophoresis before being cleaned with the Zymo DNA Clean & Concentrator kit following the standard protocol (California, USA). The purity of the resulting gene fragments were assessed with a NanoDrop Spectrophotometer (ThermoFisher, Massachusetts, USA), with the following 260/230 ratios: 1.94 for the 16S rRNA product, 1.66 for the A-amoA product, and 1.74 for the B-amoA product.

The PCR products were then ligased to prepare for plasmid insertion with 1 ul of the Promega T4 DNA Ligase (California, USA), 1 ul Promega pGEM®-T Easy Vector, and 5 ul of the Promega 2x Rapid Ligation Buffer. Additionally, the following volumes of the cleaned PCR products were added: 2 ul of the B-amoA product with 1 ul H2O, 1 ul of the 16S rRNA product with 2 ul H2O, and 1.5 ul of the A-amoA product with 1.5 ul H2O, resulting in a 10 ul reaction. 2 ul of the ligase solution was then mixed with 50 ul competent *E. Coli* and placed on ice for 20 minutes. The solution was then heat shocked at 42°C for 45-50 seconds before being placed on ice for a further 2 minutes. Then, 950 ul room-temperature SOC medium was added to the solution and incubated at 37°C with 150 rpm shaking for 1.5 hours. Finally, 100 ul of the solution was plated onto an agar medium with 166 mg NaCl, 166 mg yeast, 330 mg Tryptone, 500 mg agar, 0.1 ul ampicillin, 0.04 g X-Gal and 1.67 ml water. The agar cultures were then incubated overnight at 37°C.

Colonies that displayed successful plasmid insertion were picked for each gene (as indicated by the white colony color) and amplified via the same PCR protocol as detailed above for the initial gene amplification. The quality of the plasmids was then visually investigated via gel electrophoresis, and two colonies were selected for each inserted gene to amplify in liquid SOC medium with 8.33 ul ampicillin overnight at 37°C with 150 rpm shaking. At this step, cryostocks for each plasmid gene were frozen at -80°C with a 1:1 glycerin mixture. Two samples for each gene were selected for plasmid extraction and immediate use.



*Figure 9.* Annotated gel electrophoresis printouts from throughout the standard cultivation protocol:

*9a*. Extracted plasma inserts, with two replicates each of inserts for extracted archaeal amoA (AOA 1&2) and bacterial amoA (AOB 1&2). Additionally, there is a negative control and 100 bp ladder for reference. Four replicates of extracted inserts for the 16S rRNA gene (16S 1-4) are included.

*9b.* Linearized plasmid inserts, with two replicates each of linearized archaeal amoA (AOA 1&2), bacterial amoA (AOB 1&2) compared to a 1kbp ladder. Two plasmid inserts of the 16S rRNA gene (16S 1&2) are also included.

*9c.* Double-cut linearized plasmid inserts, after cut plasmid were recovered from the gel in 9b and resuspended. Archaeal amoA (AOA), bacterial amoA (AOB), the 16S rRNA gene (16S) and a 1kbp ladder are pictured.

*9d.* Results from first qPCR plate runs, with samples from the plate with the archaeal amoA standard on the right (including two standards with x10e7 and x10e6 archaeal amoA copies, four samples and one negative control). Samples from the bacterial amoA plate are on the left, and includes 2 standards (x10e8 and x10e9 copies of bacterial amoA), three samples and two negative controls. Between is a reference 100 bp ladder.

**d) AOB**

**c) AOB**

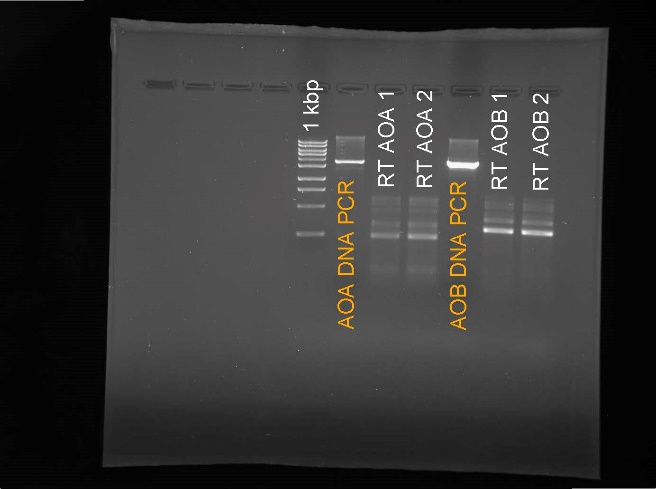
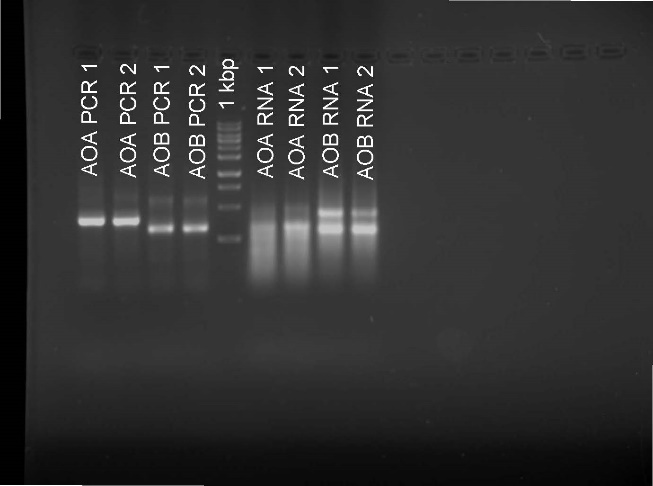
**b) AOB**

**a) AOB**

The selected plasmids were extracted using the Zymo plasmid extraction kit, which resulted in the following elution concentrations measured with NanoDrop: 201.955 and 124.844 ng/ul for A-amoA, 118.291 and 108.291 ng/ul for B-amoA, and 92.125 and 72.237 ng/ul for 16S rRNA. A subsample of each extracted plasmid was then amplified via the same PCR protocols as above and visually verified with a gel (Fig. 9b). The extracted genes were then linearized in 50 ul reactions with 1 ug DNA, 5 ul 10x NE buffer, 1 ul enzyme and the rest of the volume with H2O. The plasmids inserts were linearized at 37°C for 1 hour, and then heat inactivated at 65°C for 20 minutes.

A gel of the linearized plasmid inserts demonstrated that there was still uncut plasmid in the extracted products. Therefore, the products at the correct fragment lengths (635 bp for A-amoA, 491 bp for B-amoA and 253 bp for 16S rRNA, with the 3015 bp plasmid) were cut from the gels. The cut gel products were resuspended with the Zymoclean Gel DNA Recovery Kit and verified with another gel for purity. Then, the copies/ul of each gene was calculated via the following formula:

The ultimate copy numbers of the plasmid products were 3.32e9 for A-amoA, 1.44e9 for B-amoA, and 1.65e8 for 16S rRNA. The A-amoA and B-amoA plasmid products were aliquoted to standards with 1e9 copies/ul of the inserted gene in 5 ul of H2O, and stored at -80°C until use for qPCR applications.



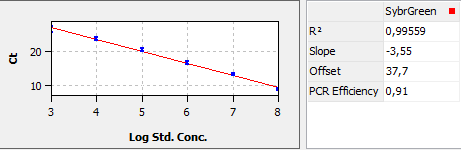
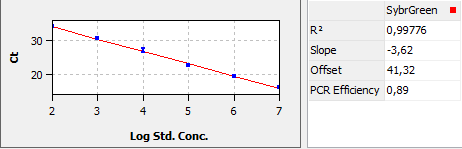
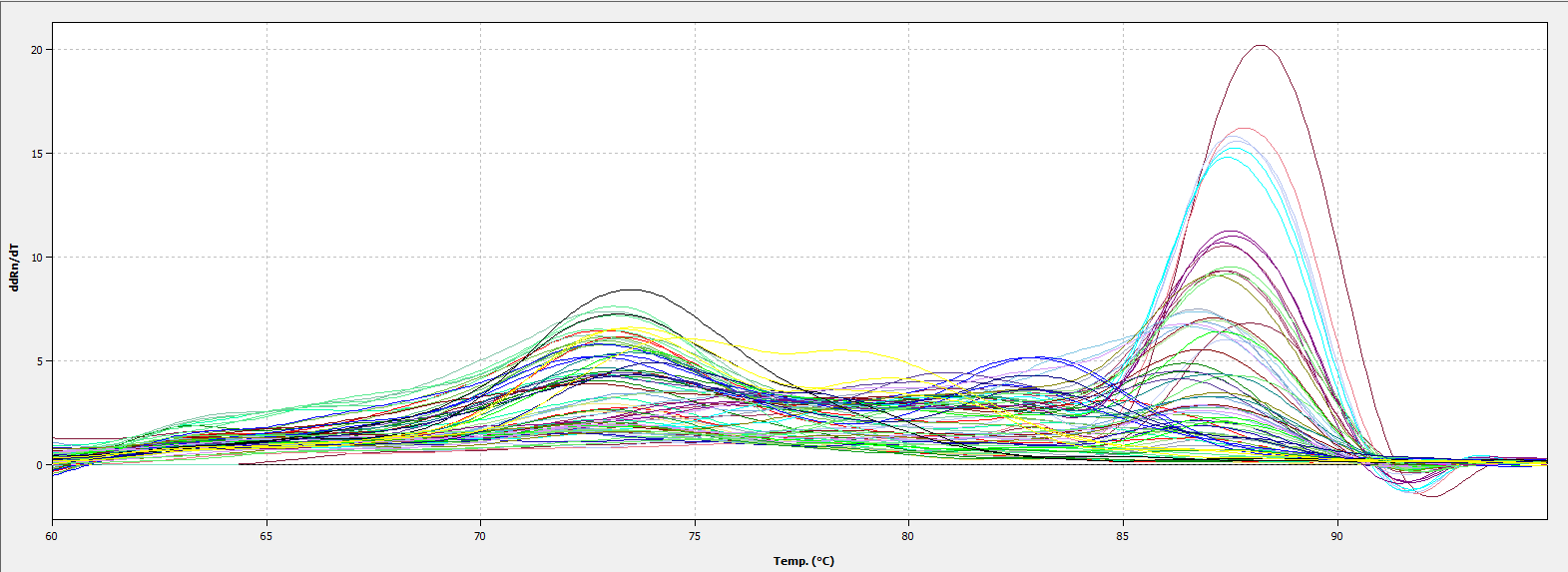
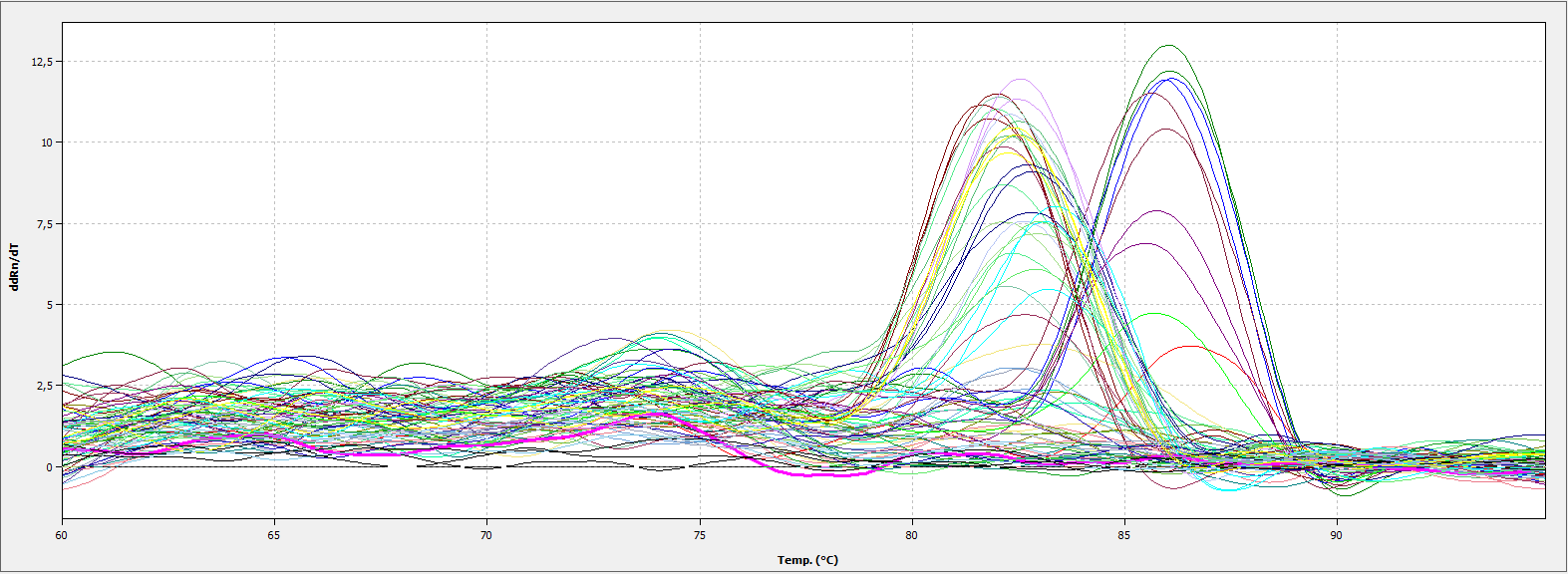
**b) AOB**

*Figure 10.* Annotated gel electrophoresis printouts from the standard transcription protocol:

*10a.* First (unsuccessful) attempt at transcribing the amoA standards, with PCR products of DNA amplified from each transcribed product. Archaeal amoA products are on the left (AOA) and bacterial amoA products are on the right (AOB), compared to a 1 kbp ladder.

*10b.* Second (and successful) attempt to transcribe amoA standards and fully remove remnant DNA to avoid DNA replication contaminating the RT-qPCR results. On the left are the PCR products amplified with the SP6/T7 primer pair and later reverse transcribed (with two replicates each from archaeal amoA, AOA PCR 1&2, and bacterial amoA, AOB PCR 1&2). On the right are the transcribed RNA standard products, with two replicates each of archaeal amoA (AOA RNA 1&2) and bacterial amoA (AOB RNA 1&2). They are separated by a 1kbp reference ladder.

**a) AOB**

**V. (RT-)qPCR results**

*Figure 11.*

*11a.* Output of one of three qPCR plates using the bacterial amoA standard. The R2 value was 0.996 and the slope was -3.55 with an efficiency of 0.91. The standard curve used to calculate copy numbers is inlaid on the upper left of the figure.

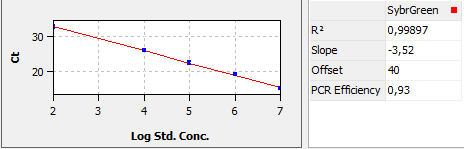
*11b.* Output of one of three qPCR plates using the archaeal amoA standard. The R2 value was 0.998 and the slope was -3.62 with an efficiency of 0.89. The standard curve used to calculate copy numbers is inlaid on the upper left of the figure.

**b) AOB**

**qPCR AOA**

**qPCR AOB**

**a) AOB**



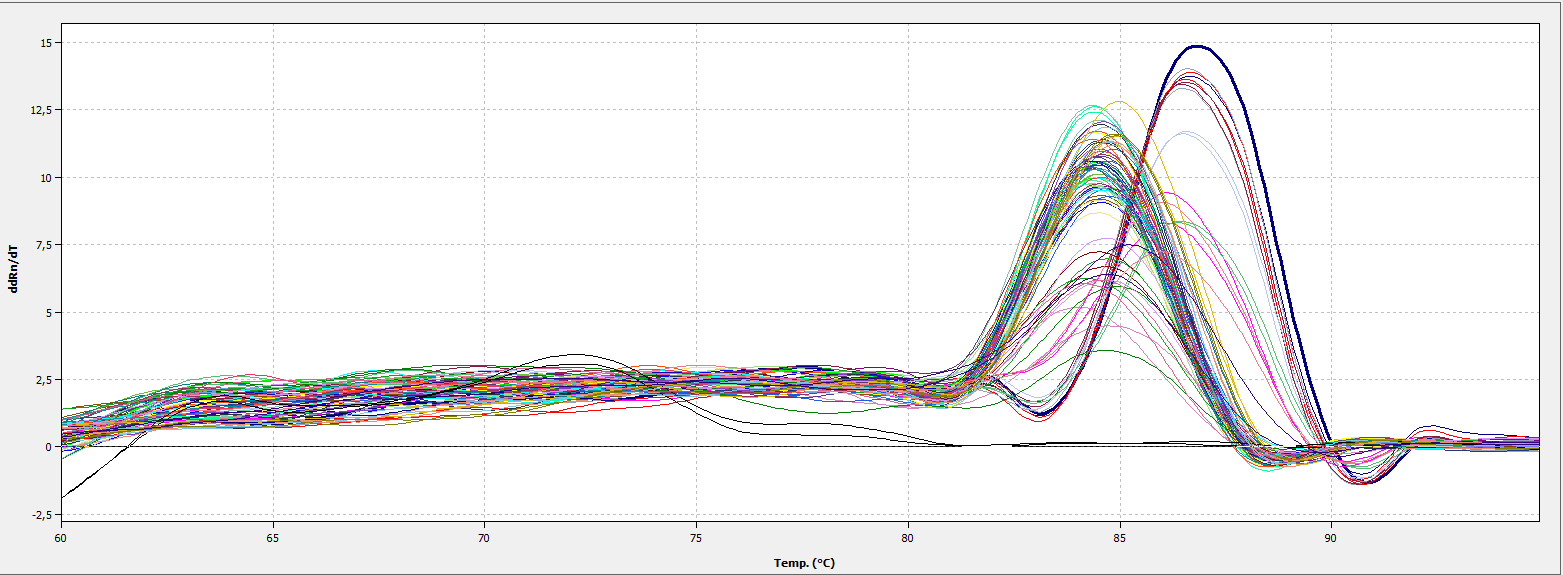
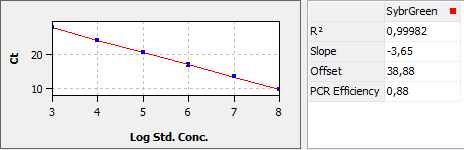
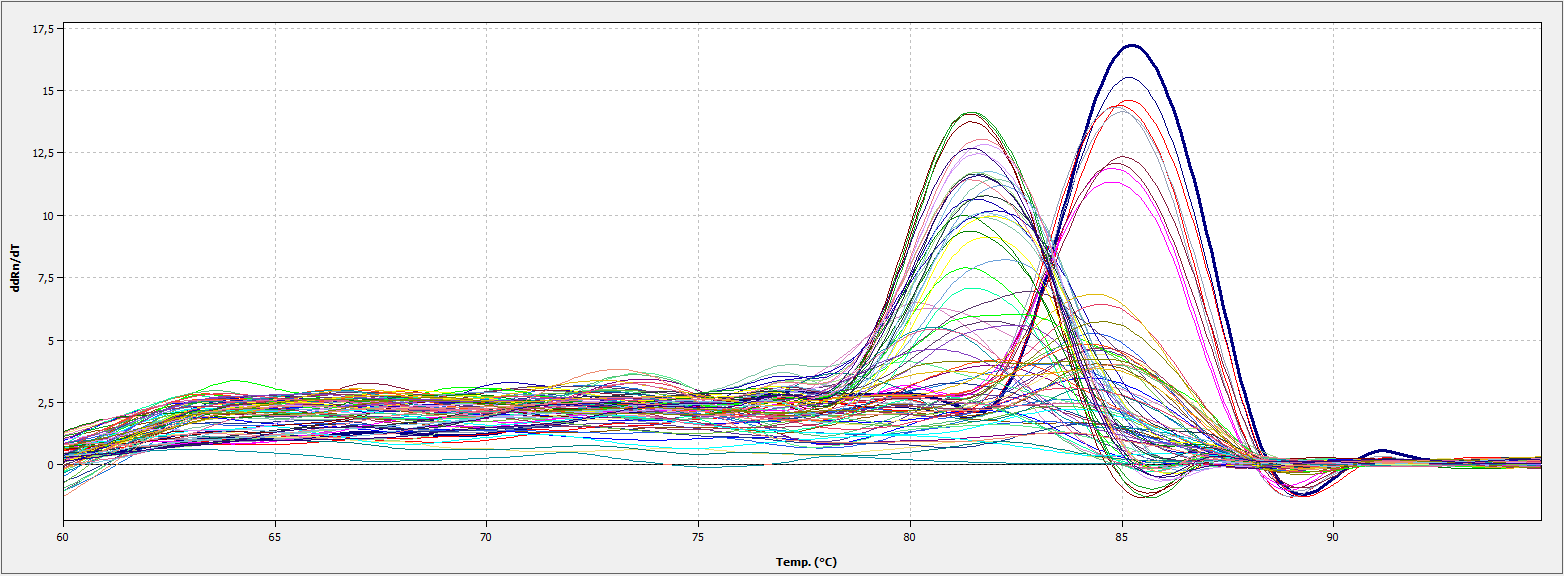
*Figure 12.*

*12a.* Output of RT-qPCR plate using the bacterial amoA standard. The R2 value was 0.999 and the slope was -3.65 with an efficiency of 0.88. The standard curve used to calculate copy numbers is inlaid on the upper left of the figure.

*12b.* Output of RT-qPCR plate using the archaeal amoA standard. The R2 value was 0.999 and the slope was -3.52 with an efficiency of 0.93. The standard curve used to calculate copy numbers is inlaid on the upper left of the figure.

**RT-qPCR**

**AOA**



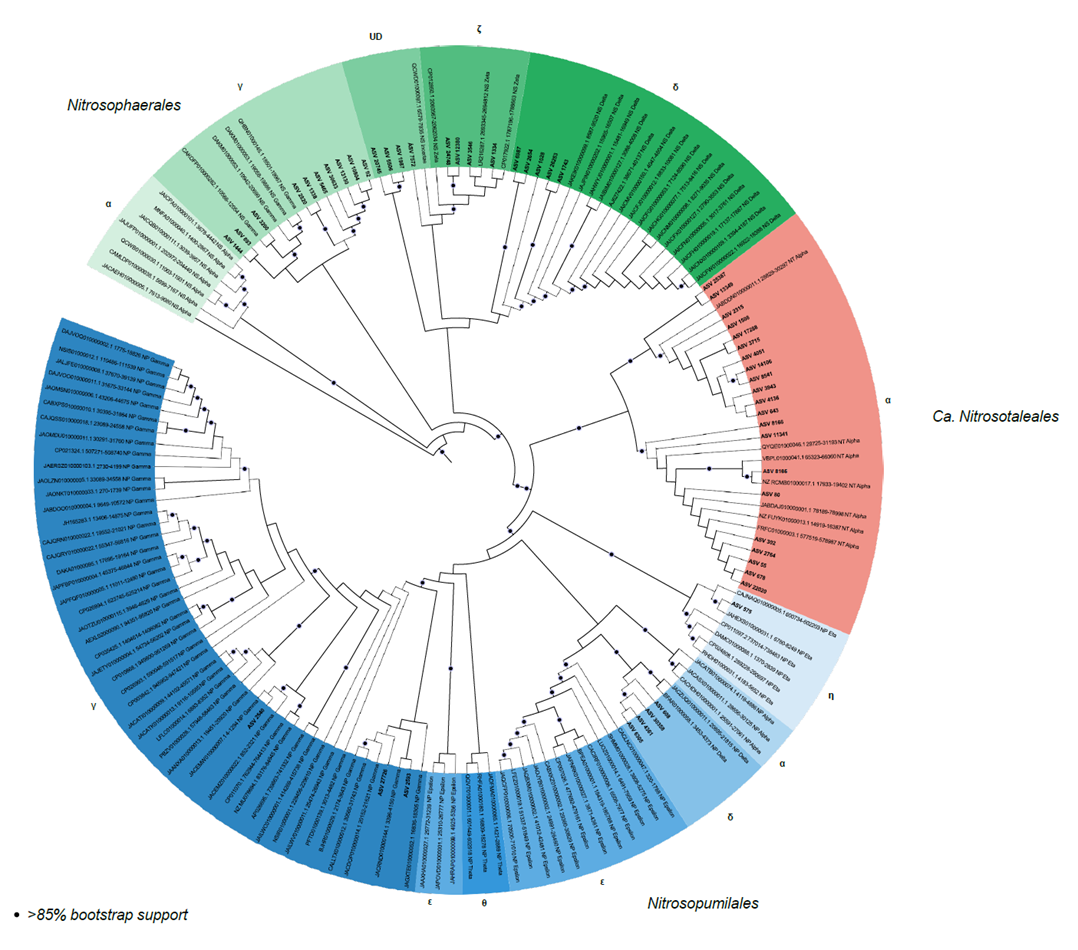
**b) AOB**

**a) AOB**

**RT-qPCR**

**AOB**

**VI. Appendix III: AOA Phylogenetic Tree Assembly**



*Figure 13.* Phylogenetic tree

**VI. Conclusion**

**VII. Works Cited**

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