

16S rRNA Gene Amplicon Survey: Study Design and Case Study

Considerations for a Longitudinal Case Study of Antibiotic Treatment and Virus Infection

Scott A. Handley, PhD

Assistant Professor

Department of Pathology & Immunology

Washington University School of Medicine

Rationale

- 16S amplicon surveys are extensively used to study the mouse bacterial microbiome in a large variety of contexts
 - e.g. disease, nutrition, sociology, neuroscience, etc.
- **Frequently** fail due to poor study design
 - Batch effects
 - Cage, paternity/breeding, facility, origin effects
 - Co-housed survival studies (specific example)
 - Statistical considerations
 - Detecting signal from noise
 - Minimize variance
 - Filtering out misbehaved data
- Many of these principles apply to other data types (RNAseq)



Image credit: Davide Bonazzi/@Salmanart

"Mouse microbes may make scientific studies harder to replicate" Kelly Servick. Science Aug 16, 2016

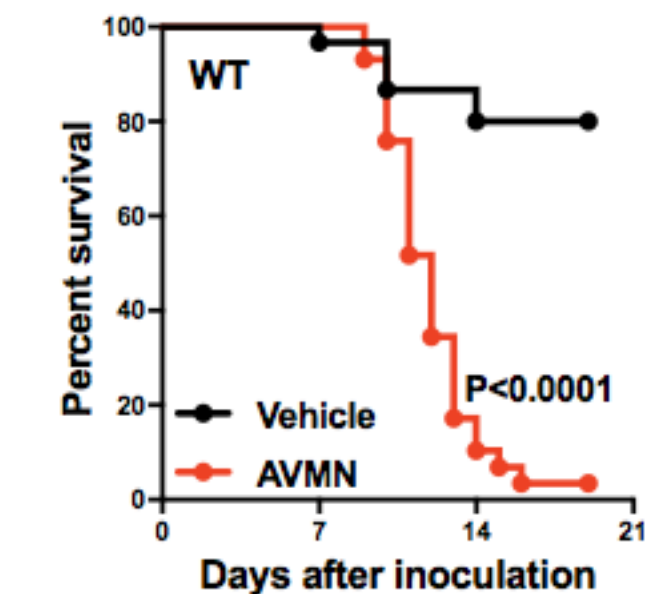
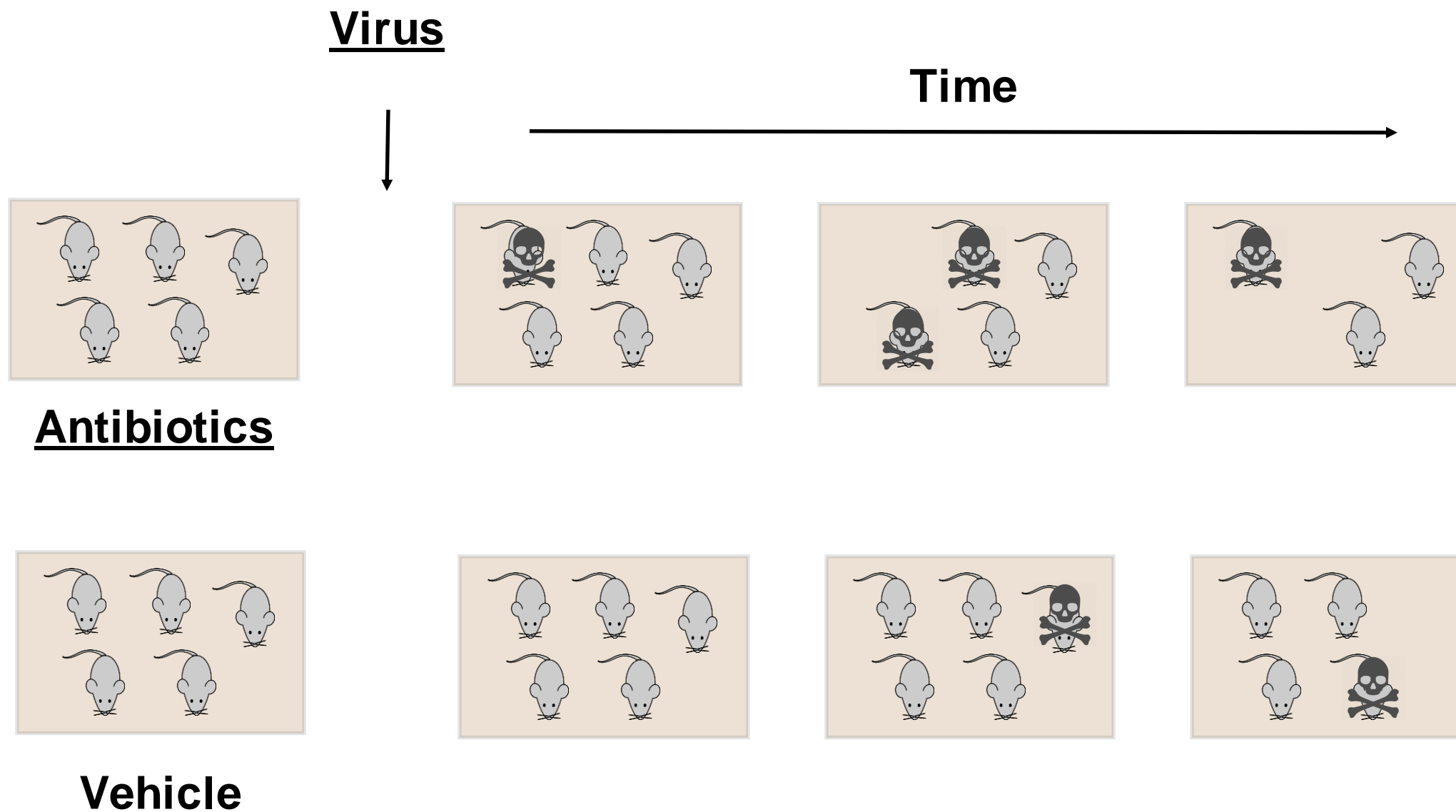
"Accounting for reciprocal host-microbiome interactions in experimental science" Stappenbeck, TS and Virgin HW. Nature. 2016 Jun 9;534(7606):191-9



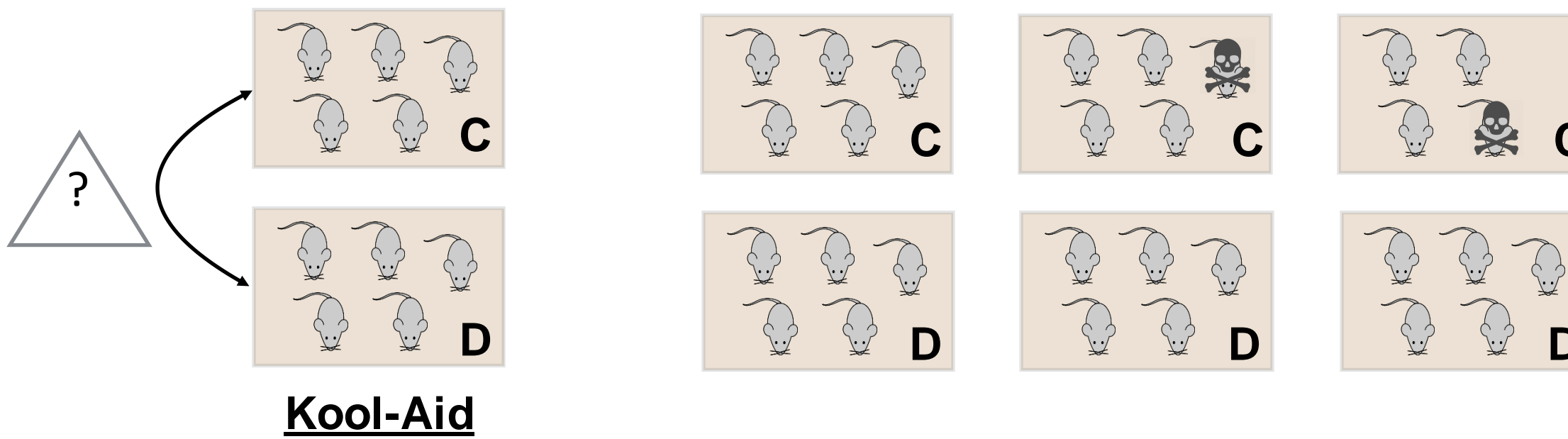
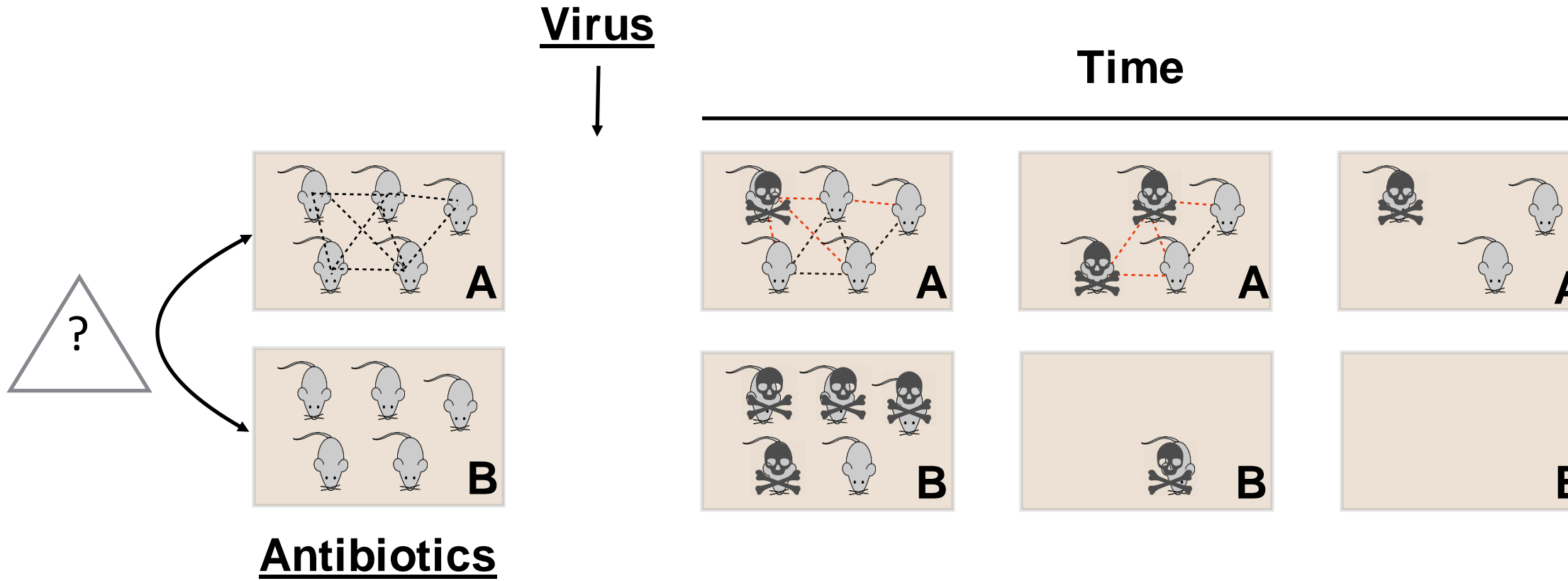
Today's Case Study

Thackray LB, Handley SA, Gorman MJ, Poddar S, Bagadia P, Briseño CG, Theisen DJ, Tan Q, Hykes BL Jr, Lin H, Lucas TM, Desai C, Gordon JI, Murphy KM, Virgin HW, Diamond MS. **Oral Antibiotic Treatment of Mice Exacerbates the Disease Severity of Multiple Flavivirus Infections.** Cell Rep. 2018 Mar 27;22(13):3440-3453.e6. PubMed PMID: 29590614

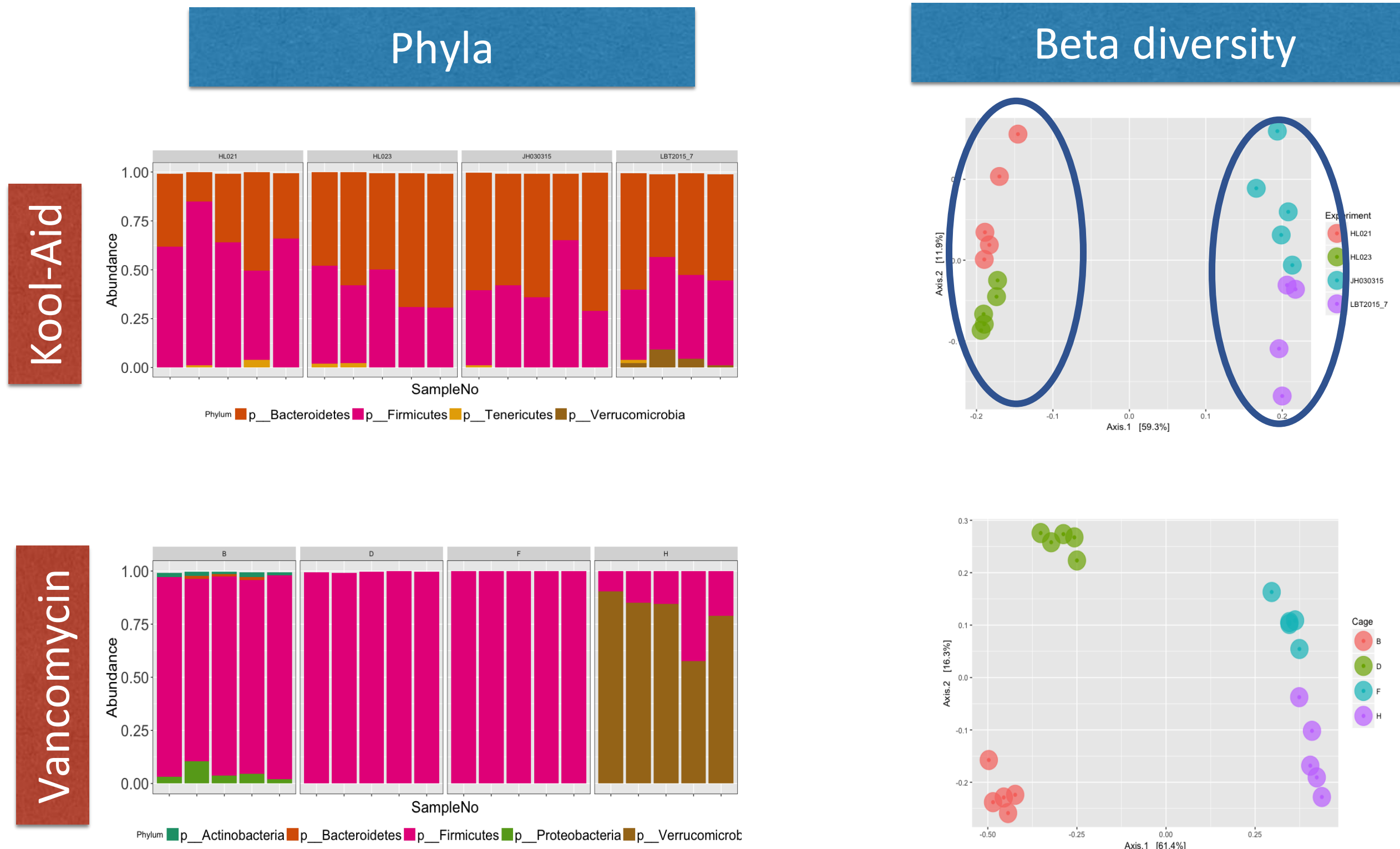
Case Study: Effect of Antibiotics on Flavivirus Pathogenesis



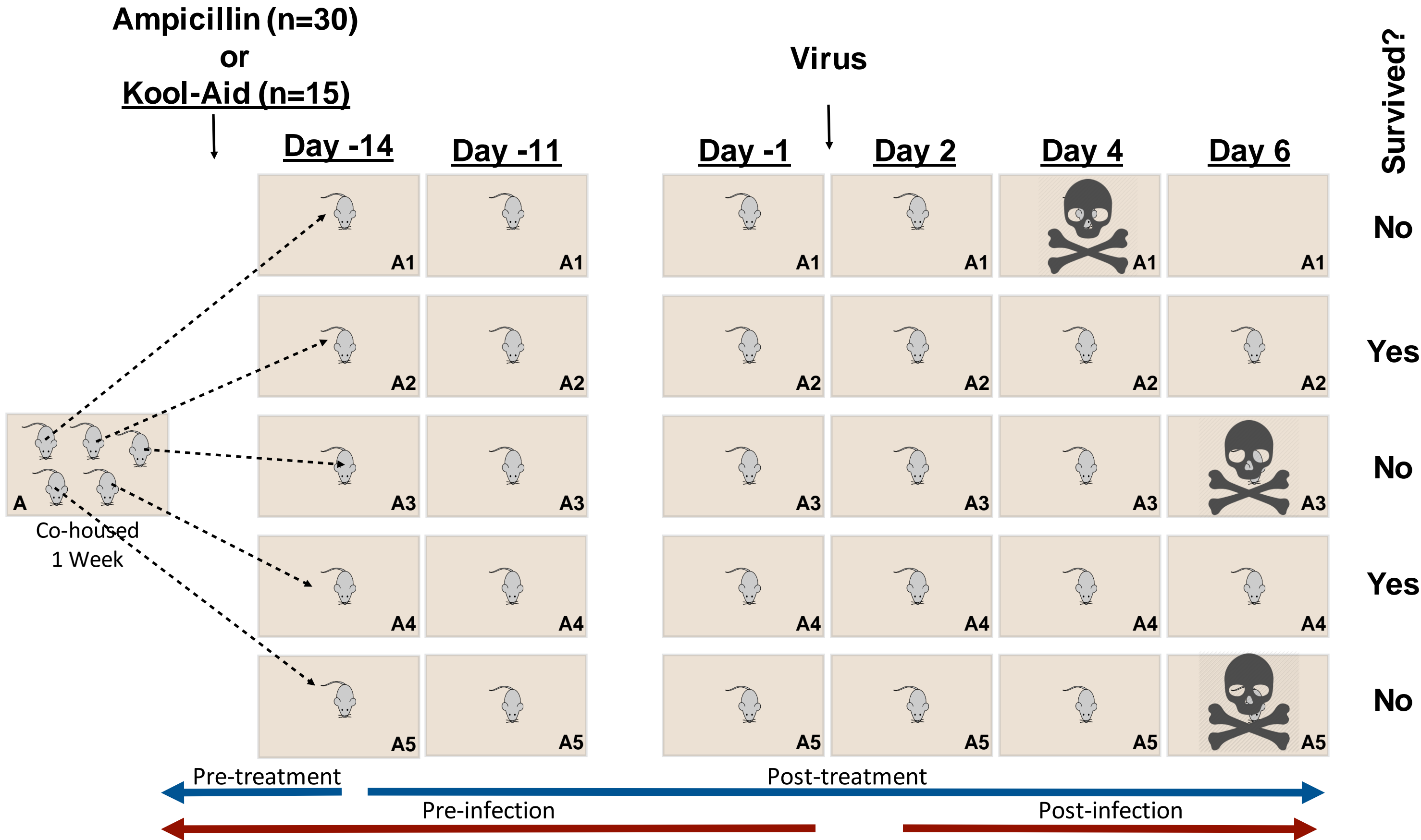
Cage and Mouse-to-Mouse Effects



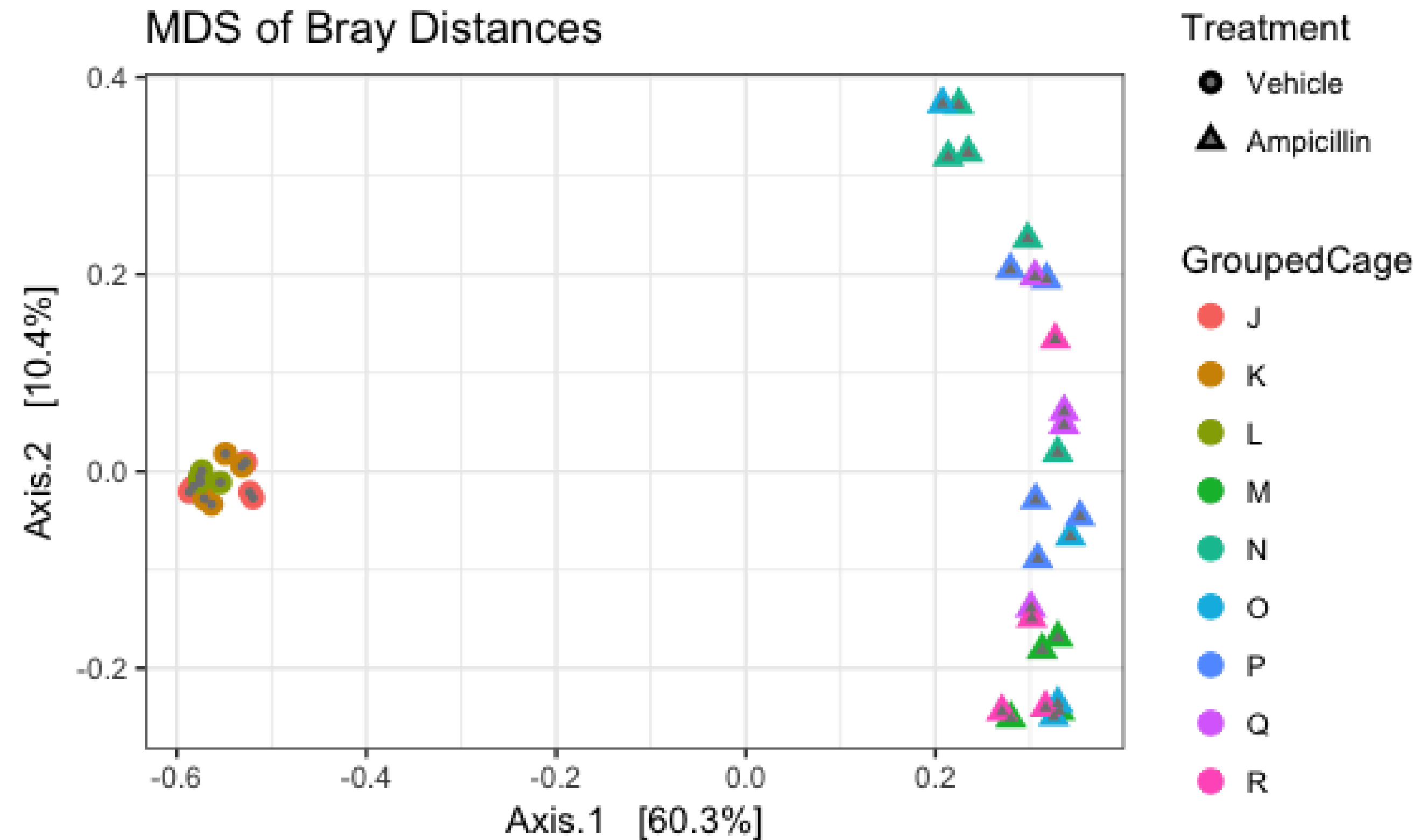
Cage Effects: 14 days post-treatment (pre-infection)



Individual Mouse Isolation Schema

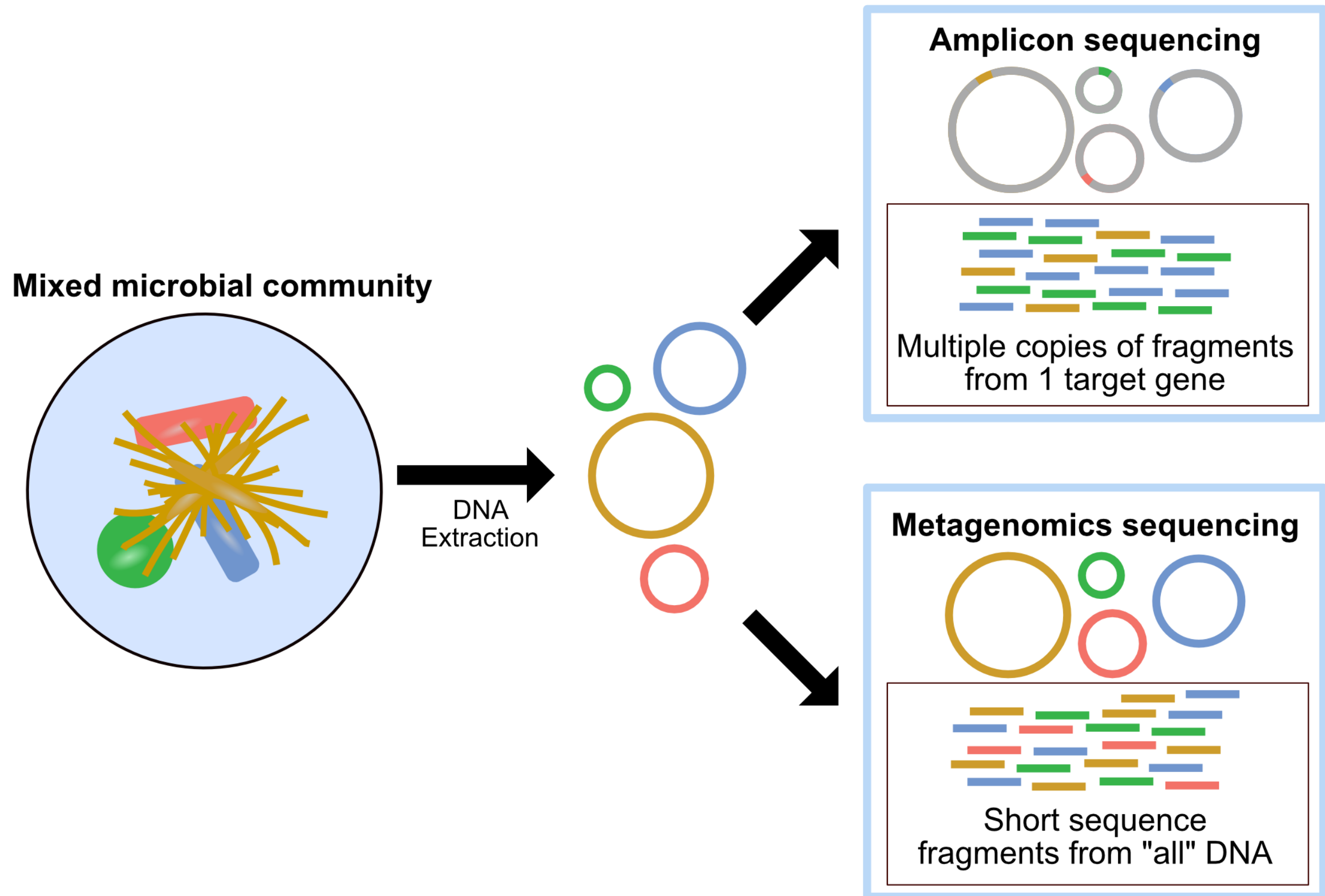


Individual Housing Results

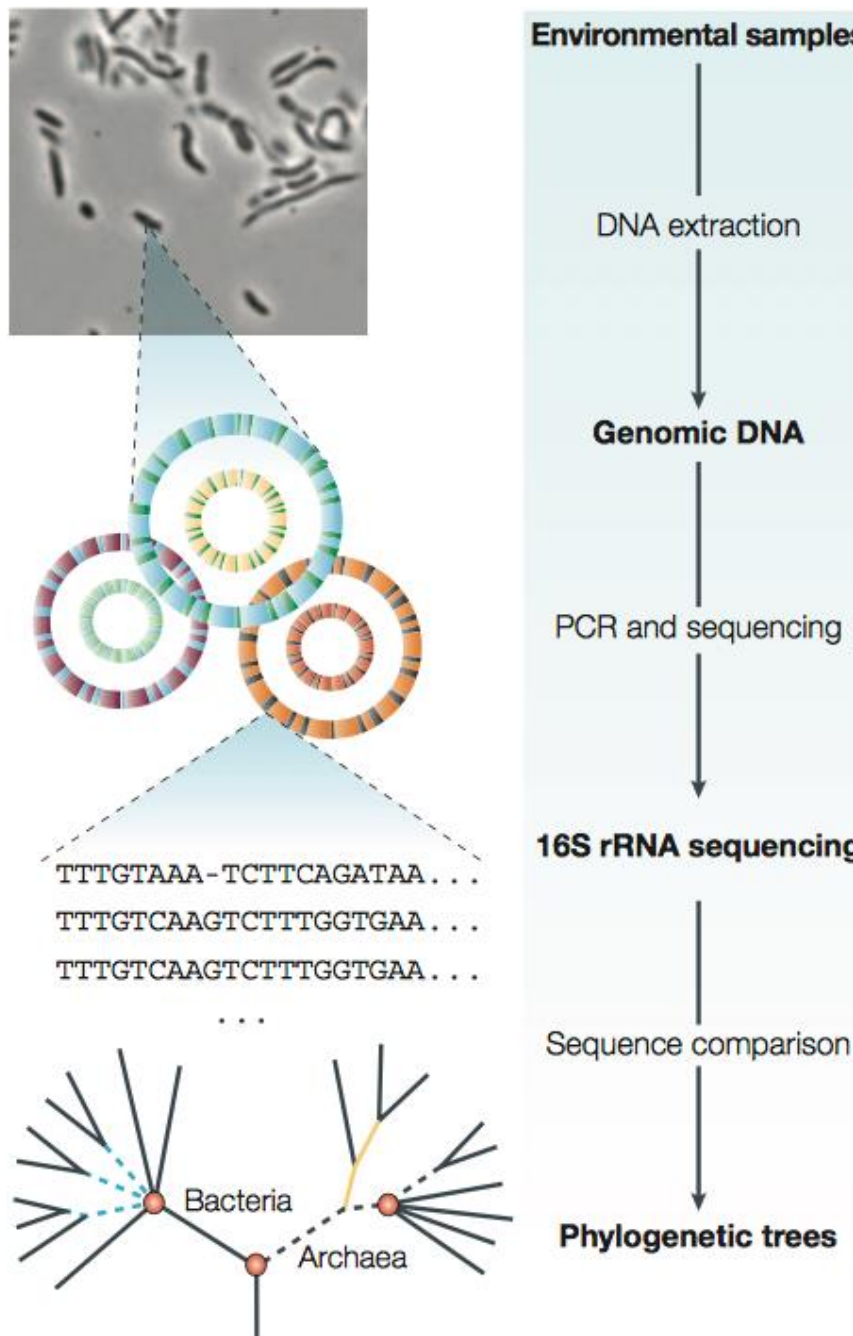


Amplicon Surveys (Highly Opinionated!) Best-practices

It's the classic garbage in, garbage out all over again ...



16S rRNA Amplicon Survey



Study Design

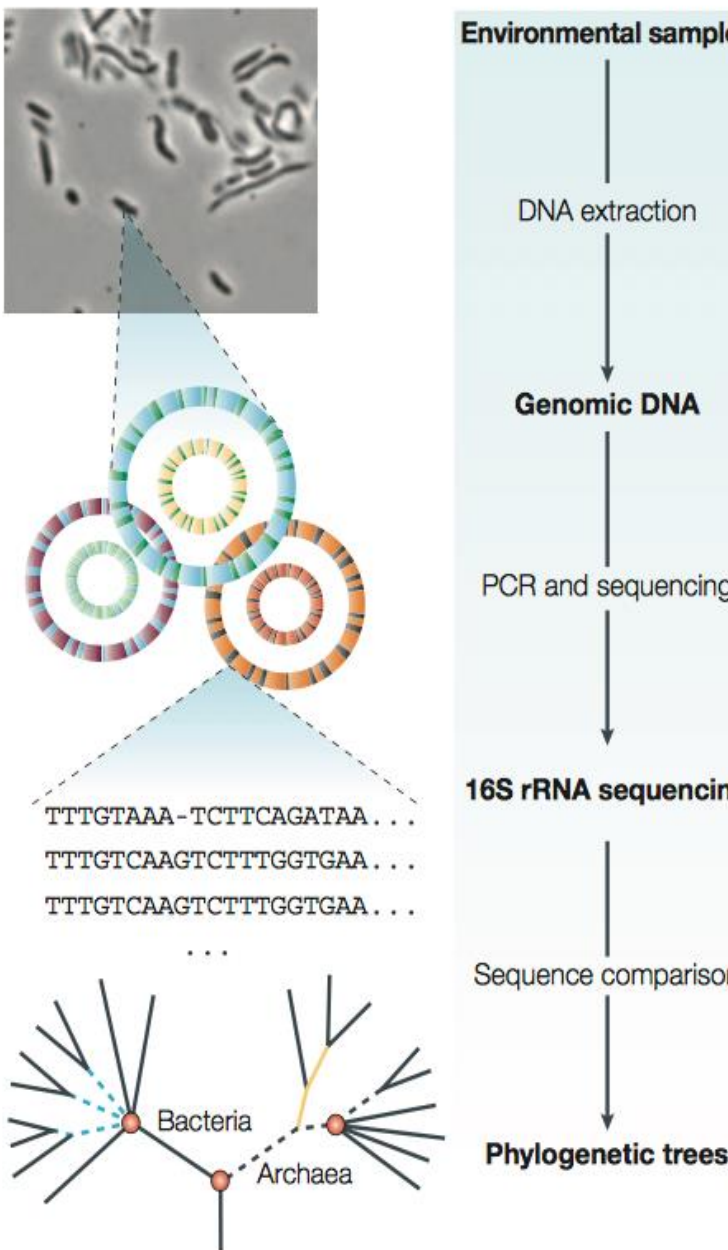
Laboratory

Bioinformatics, Ecological
Analysis and Statistics

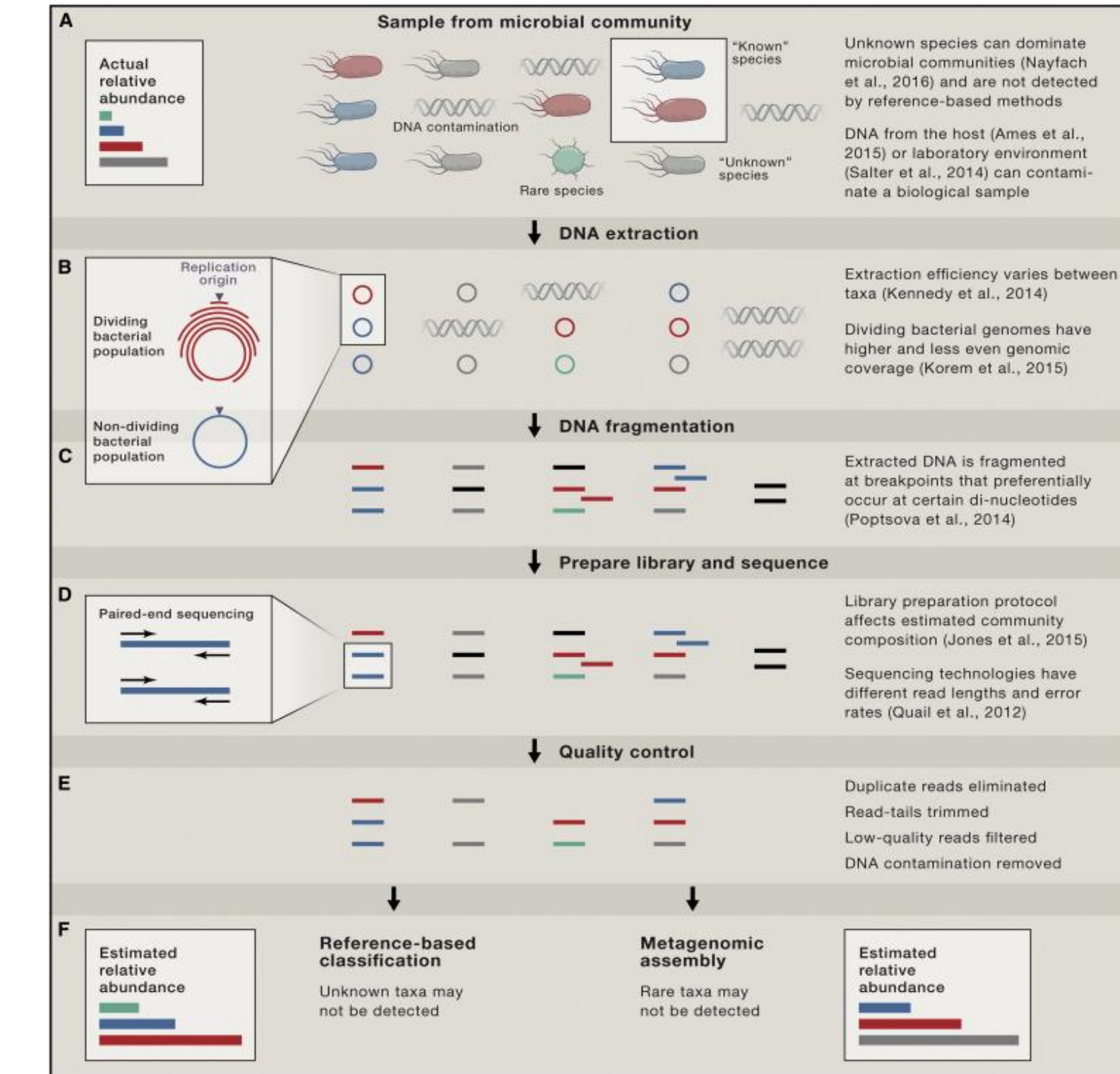
Side note: Amplicon Surveys vs. Metagenomics

Please hold your throwing tomatoes ...

16S Amplicon Surveys vs Metagenomics?



Tringe, S.G., Rubin, E.M. Nat Rev Genet. 2005 Nov;6(11):805-14



Nayfach S., Pollard KS. Cell. Aug 25;166(5):1103-16

Most of Your Decision Will Boil Down to \$\$\$ and information type

- Our labs per sample costs:

- 16S = \$17.50 per sample
- Metagenome = \$225.00 per sample

• Student Resources

• [Methods and Protocols](#) | Novel Systems Biology Techniques

• Other Publications

Evaluating the Information Content of Shallow Shotgun Metagenomics

Benjamin Hillmann, Gabriel A. Al-Ghalith, Robin R. Shields-Cutler, Qiyun Zhu, Daryl M. Gohl, Kenneth B. Beckman, Rob Knight, Dan Knights

- Understanding analytical space
- Data storage
- What type of information do you need? Taxonomic or functional



The mSystems journal logo features the word "mSystems" in a large, bold, red sans-serif font. To the left of the "m" is a stylized blue and white circular graphic. Below the main title, it says "AN OPEN ACCESS JOURNAL PUBLISHED BY THE AMERICAN SOCIETY FOR MICROBIOLOGY".

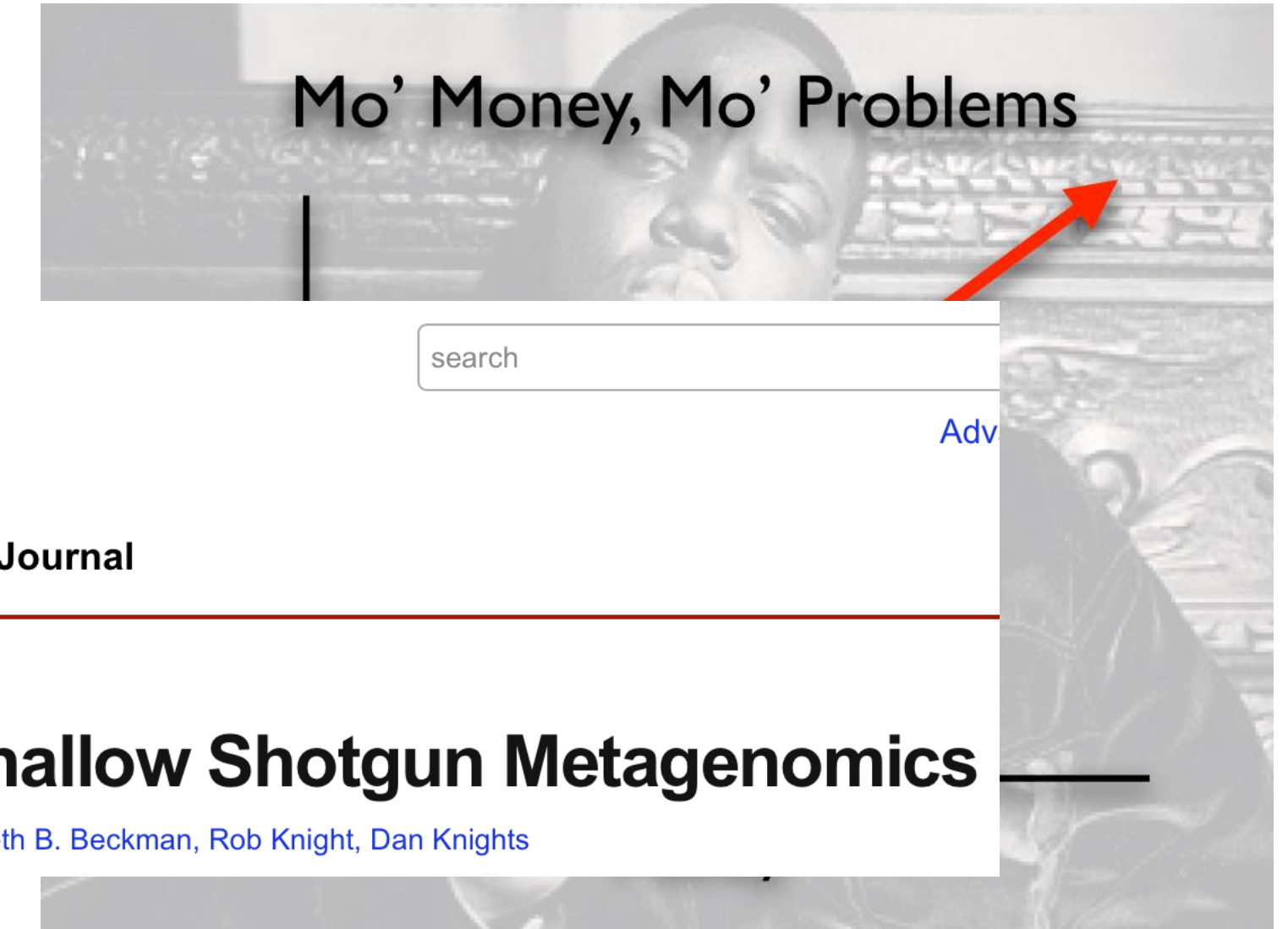
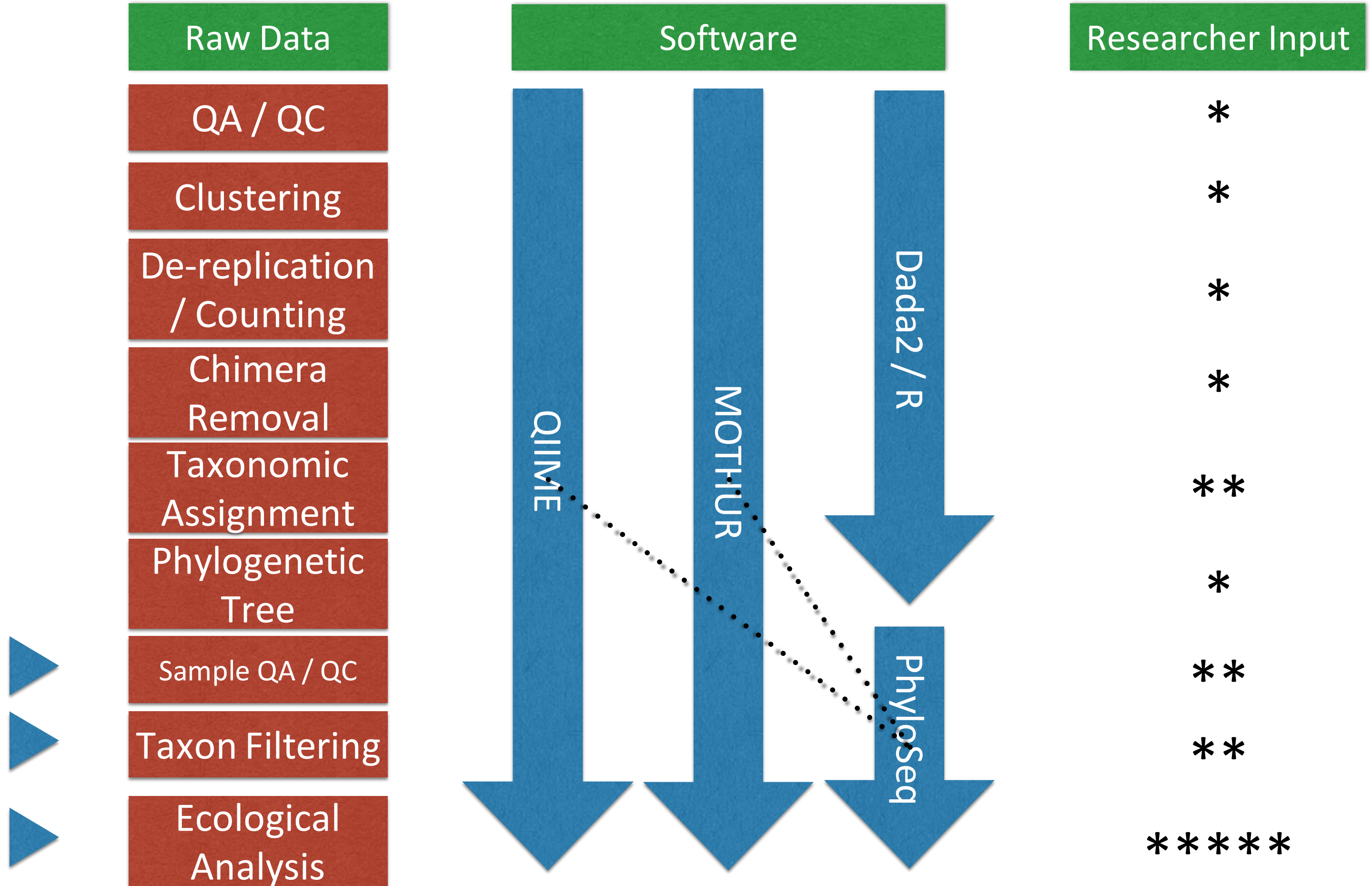


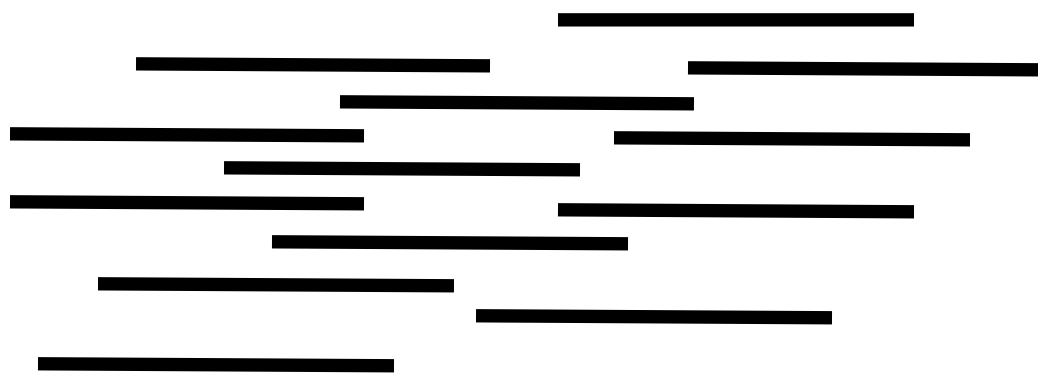
Image credit: The Internet
Quote credit: Notorious B.I.G.

What are the stages of a 16S amplicon computational workflow and how can we create optimal data for analysis?



Sequence Clustering

16S RNA Amplicons

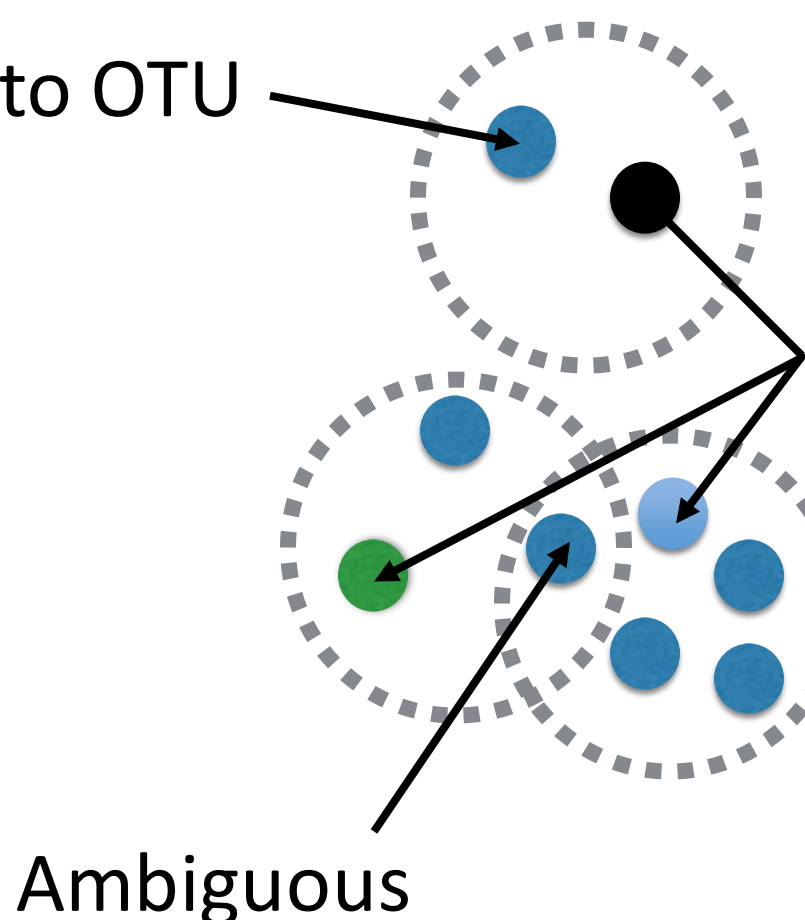


97% Similarity

Amplicon Clusters



> 97% identical to OTU

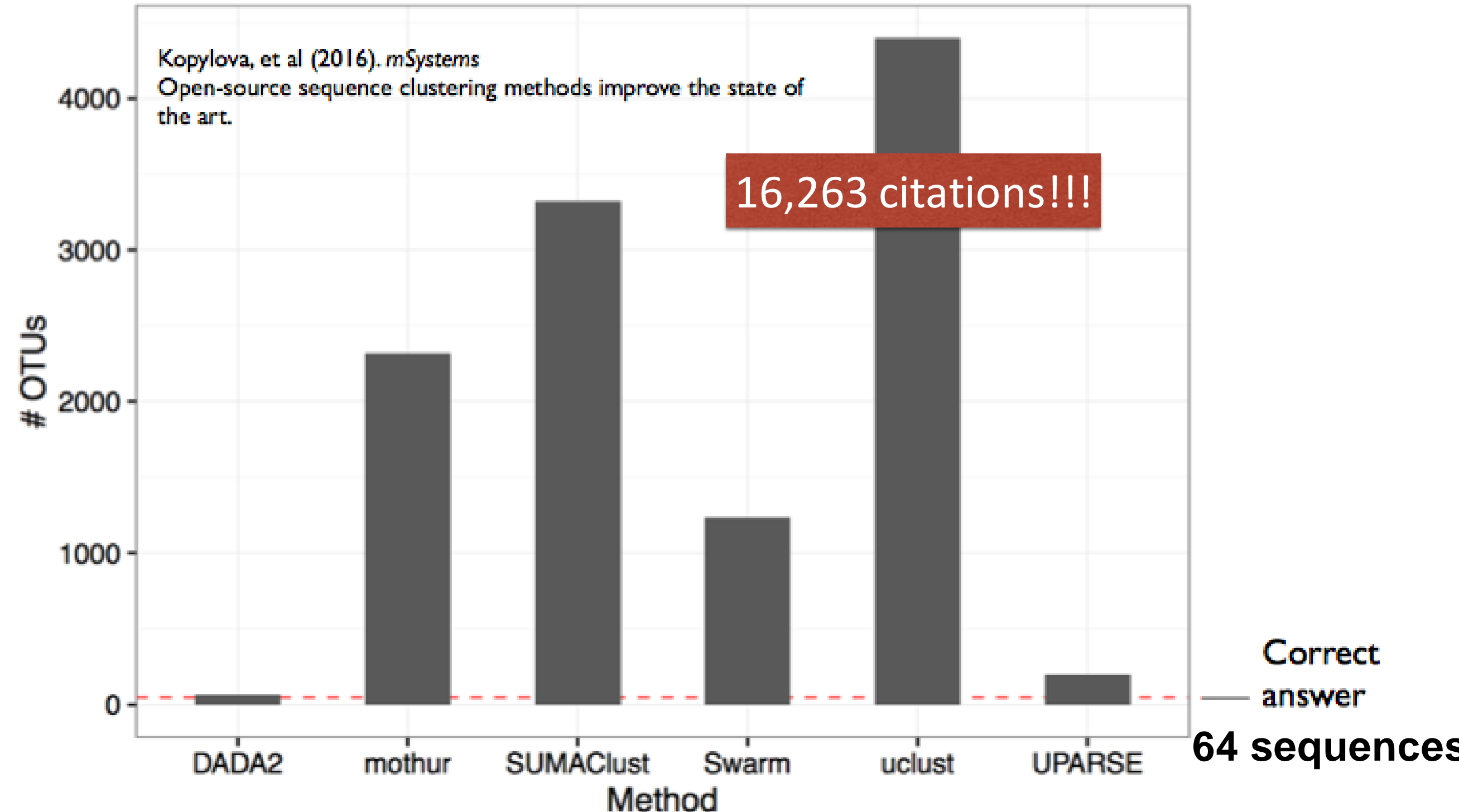


- **UCLUST**
- **UPARSE**
- **SWARM**
- **SUMACLUST**
- **OTHERS**

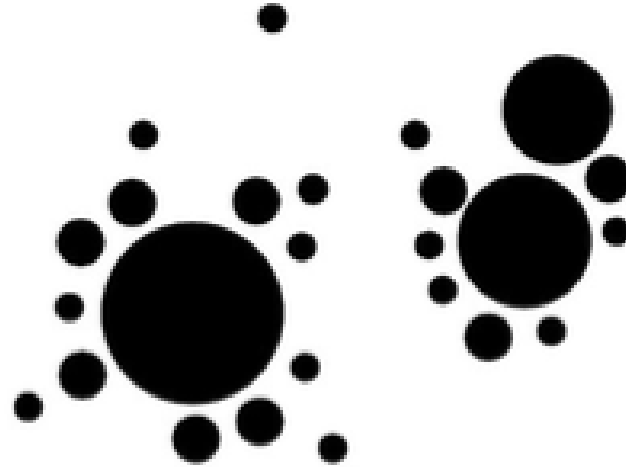
Recognized Problems with Sequence Clustering

- **False-positives:** 1,000s of OTUs when only 10s of sequences are present
 - Due to clustering artifact / noisy sequences
 - Inflates richness (# of species)
 - Sparse matrices
- **Poor taxonomic resolution** defined by arbitrary radius (e.g. 97%)
- **Increased financial cost:** poor data efficiency
- **Increased computational cost:** Clustering is quadratic
- **Unstable:** Sequence and count frequently depend on input order

There is some hope



<http://benjjneb.github.io/dada2/R/SotA.html>

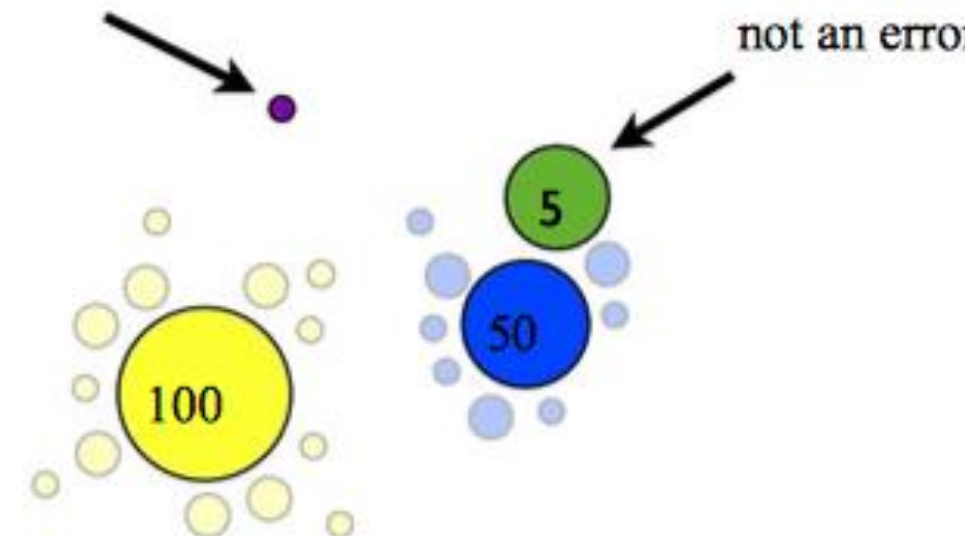


Step 1: Initial guess.
All sequences + errors

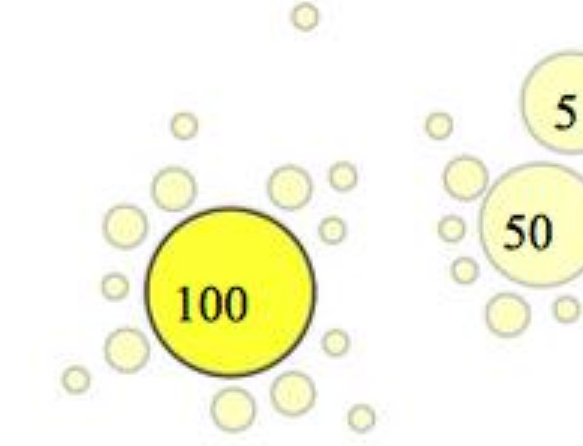
$$\Pr(i \rightarrow j) =$$

	A	C	G	T
A	0.97	10 ⁻²	10 ⁻²	10 ⁻²
C	10 ⁻²	0.97	10 ⁻²	10 ⁻²
G	10 ⁻²	10 ⁻²	0.97	10 ⁻²
T	10 ⁻²	10 ⁻²	10 ⁻²	0.97

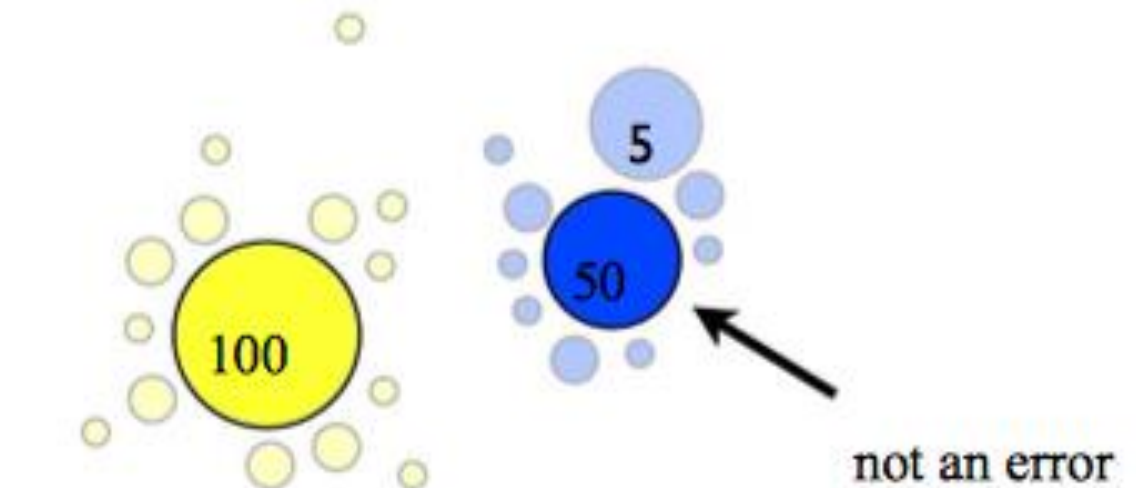
not an error



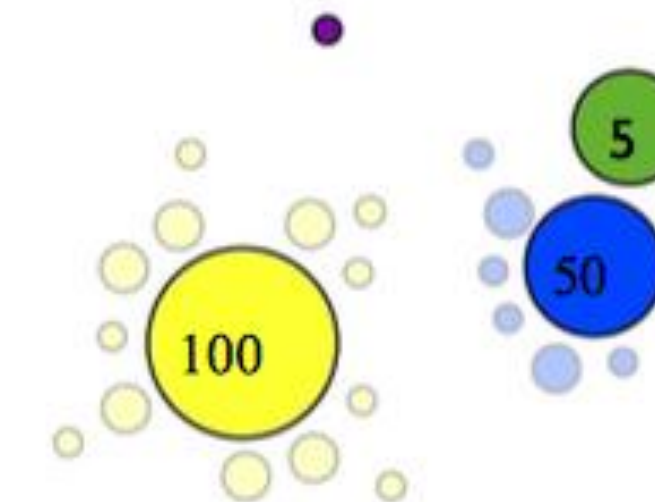
Step 3: Reject more sequences
under new model & update



Step 2: Initial error model



Step 3: Unlikely error under model.
Recruit errors. Update the model

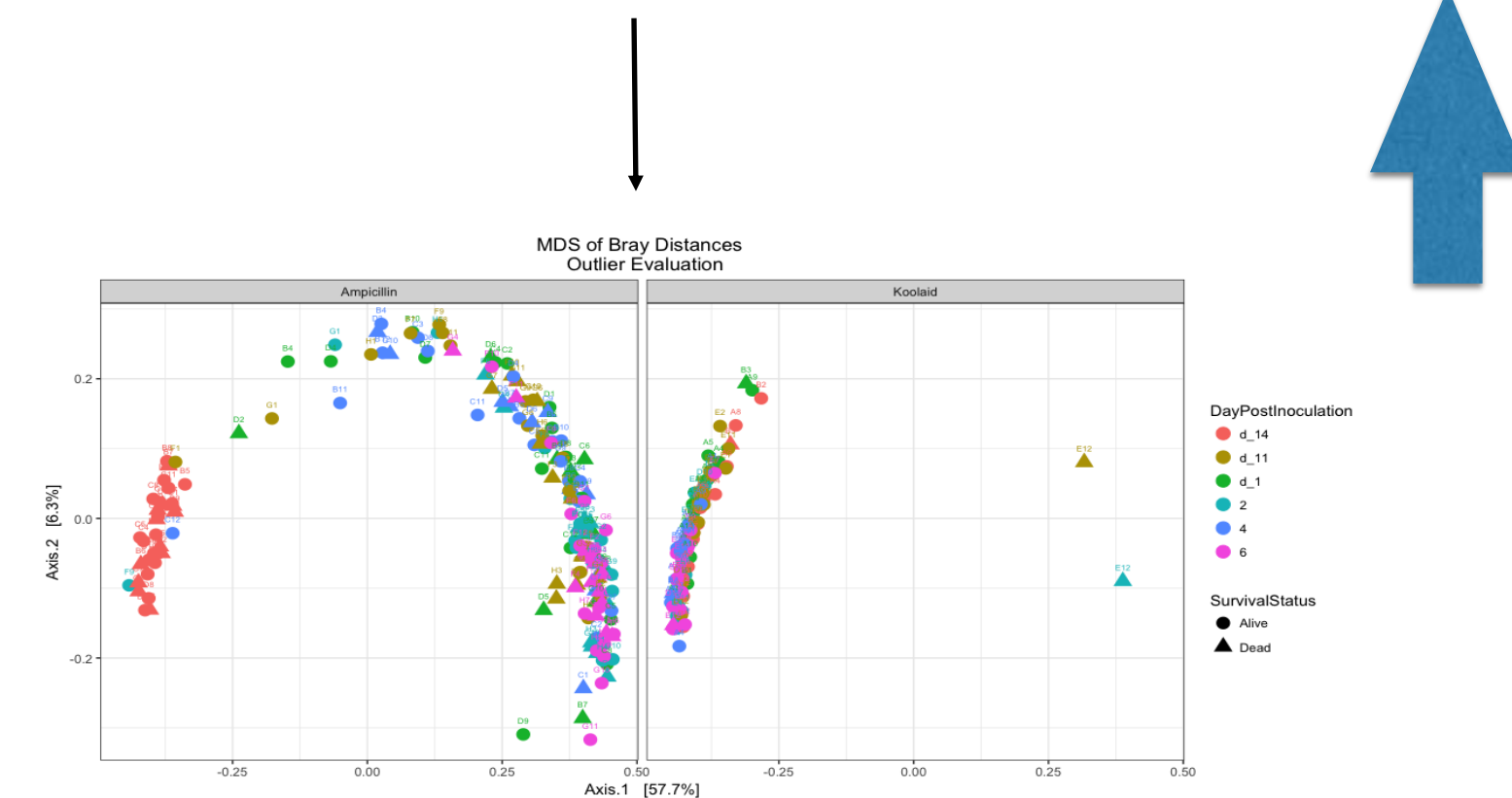


Convergence: All errors are plausible

What does all of this work get you?



ID	Sample 1	Sample 2	Sample 3	Sample 4
ASV 1	0	0	2	0
ASV 2	12	8	8	456
ASV 3	112	101	98	10
ASV 4	435	435	382	3
ASV 5	76	83	68	145



Sparse Matrix OTU Clustering

Less Sparse Matrix Sequence Resolution

ID	Sample 1	Sample 2	Sample 3
OTU 1	0	0	1
OTU 2	1	0	0
OTU 3	1	0	0
OTU 4	1	1	1

ID	Sample 1	Sample 2	Sample 3
ASV 1	0	1	1
ASV 2	1	1	0
ASV 3	1	0	1
ASV 4	1	1	1

- More noisy than reality
- Bad for statistical inference
 - Multiple hypothesis testing
 - Poorly defined, difficult to separate distributions

Raw Data

QA / QC

Clustering

De-replication
/ Counting

Chimera
Removal

Taxonomic
Assignment

Phylogenetic
Tree

Sample QA / QC

Taxon Filtering

Ecological
Analysis

Sample Outlier Detection

ID	Sample 1	Sample 2	Sample 3	Sample 4
ASV 1	0	0	2	0
ASV 2	12	8	8	456
ASV 3	112	101	98	10
ASV 4	435	435	382	3
ASV 5	76	83	68	145

...

n=724

... n=270



Individual Mouse Isolation Schema

Ampicillin (n=30)

or

Kool-Aid (n=15)

Virus

Survived?

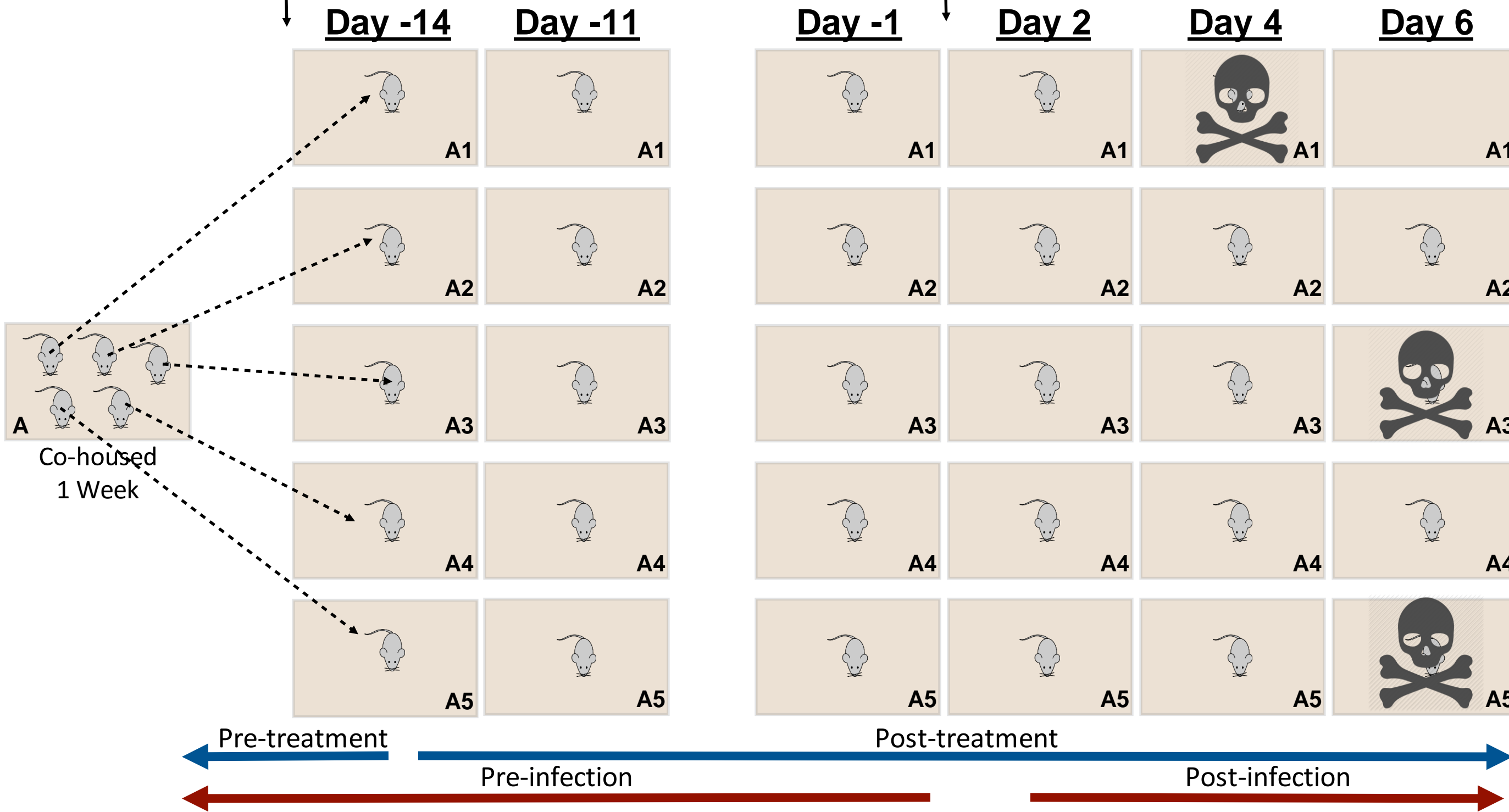
No

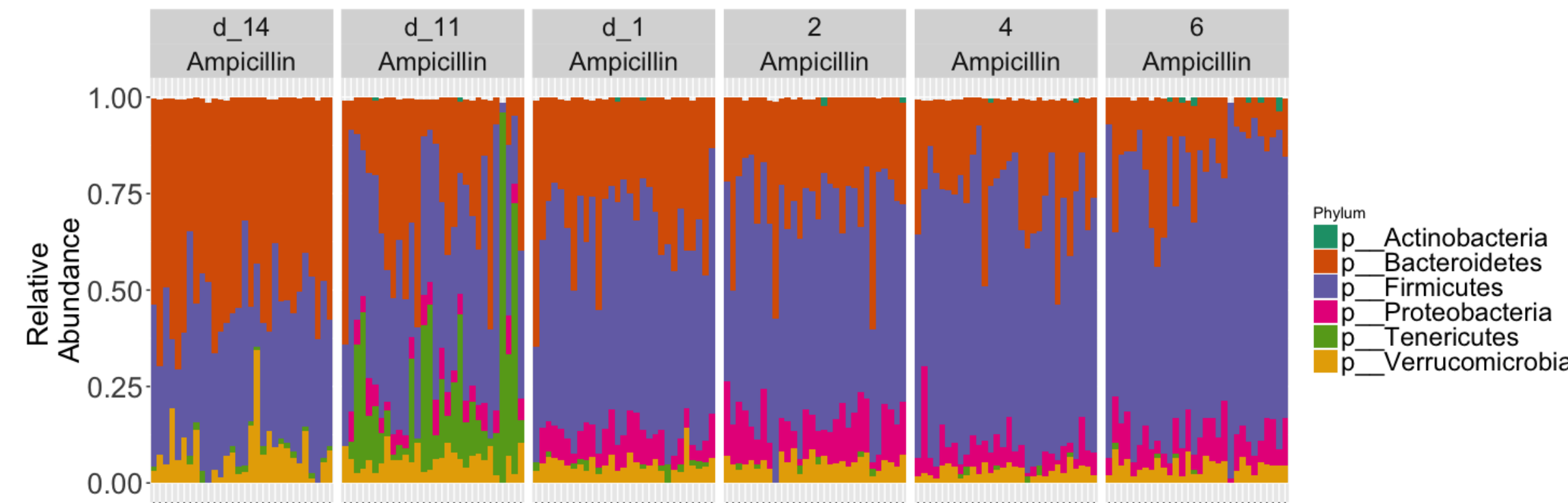
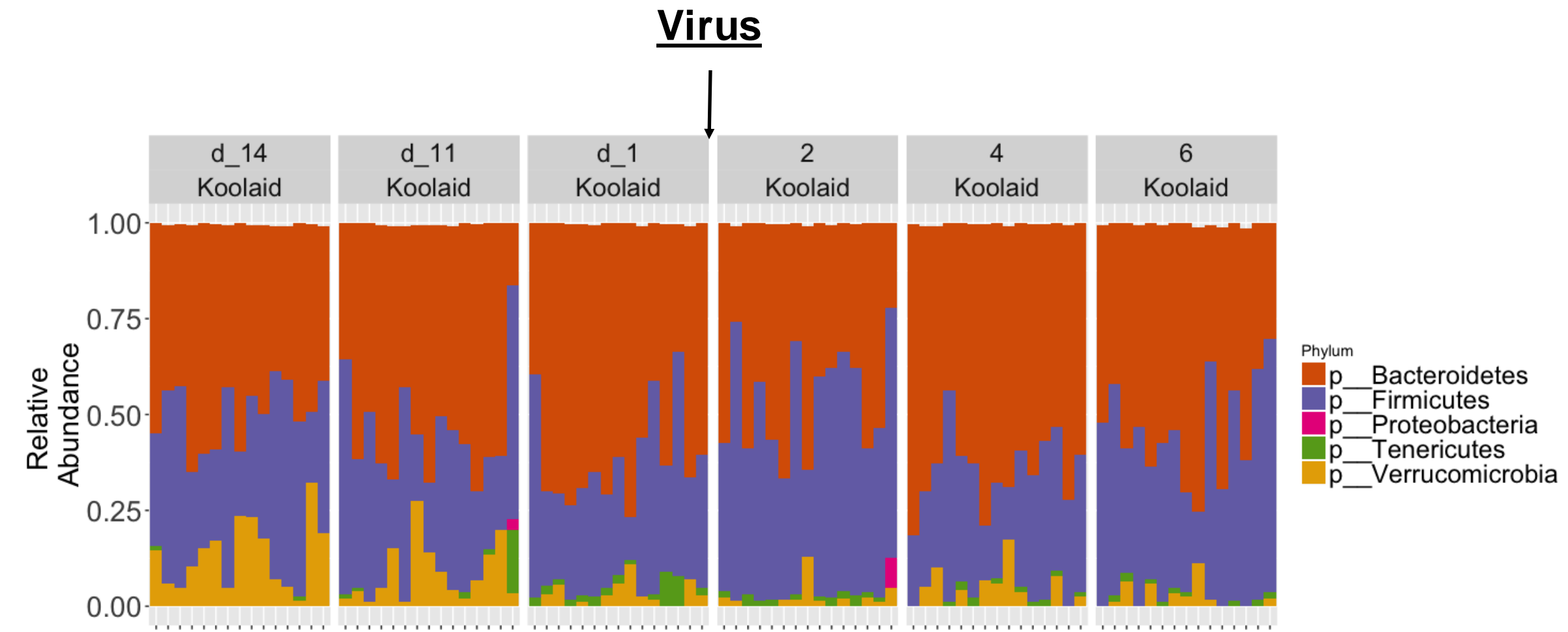
Yes

No

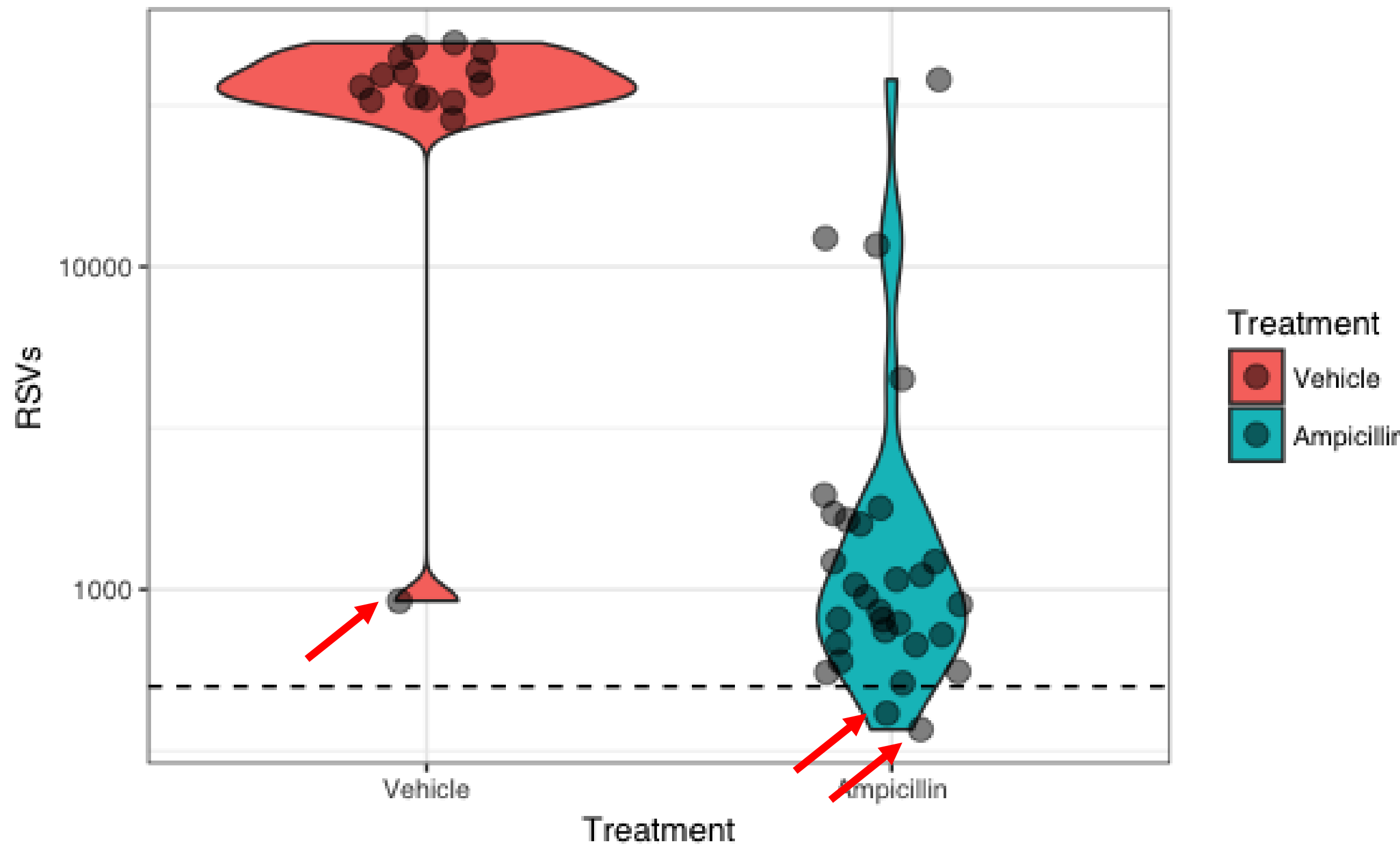
Yes

No



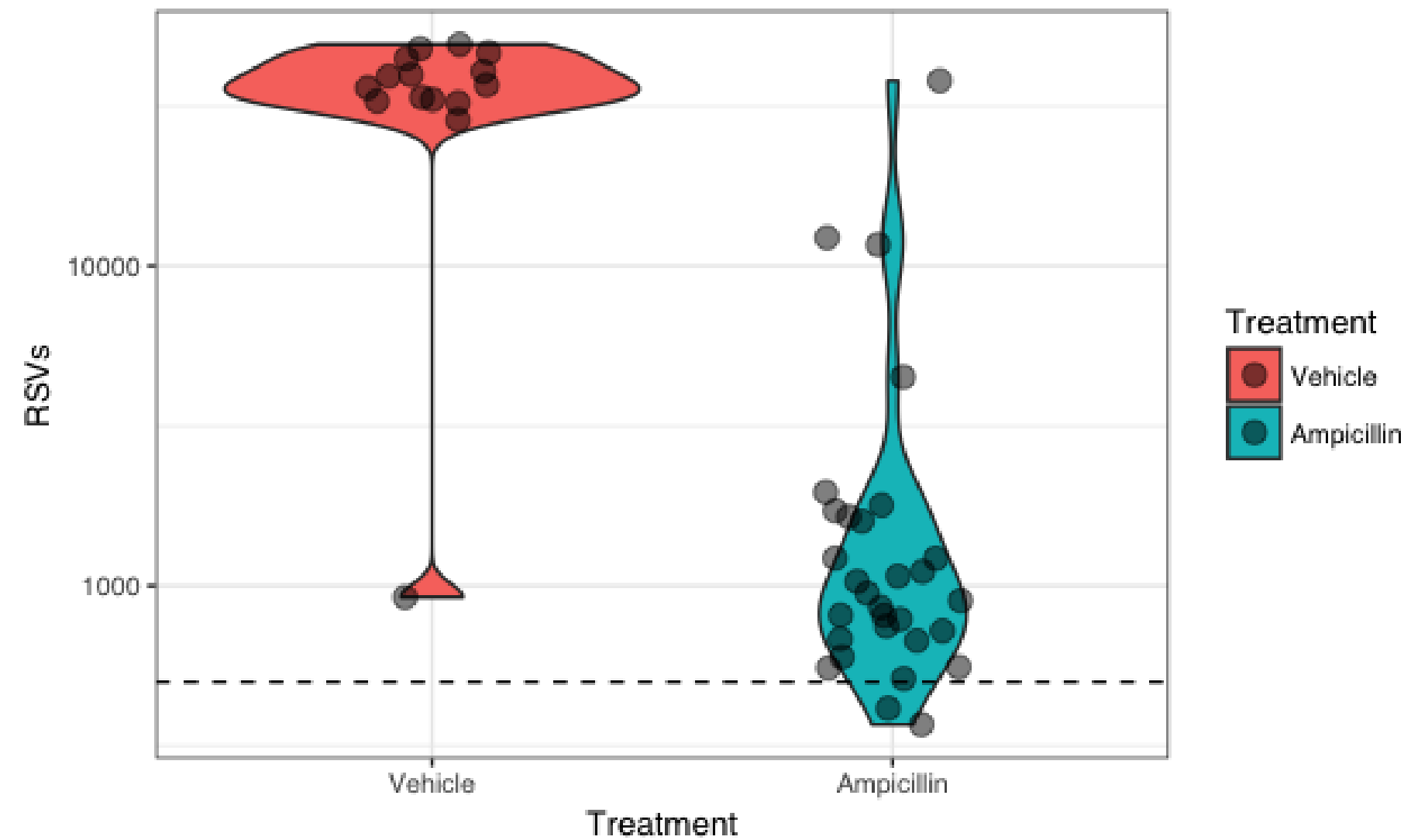
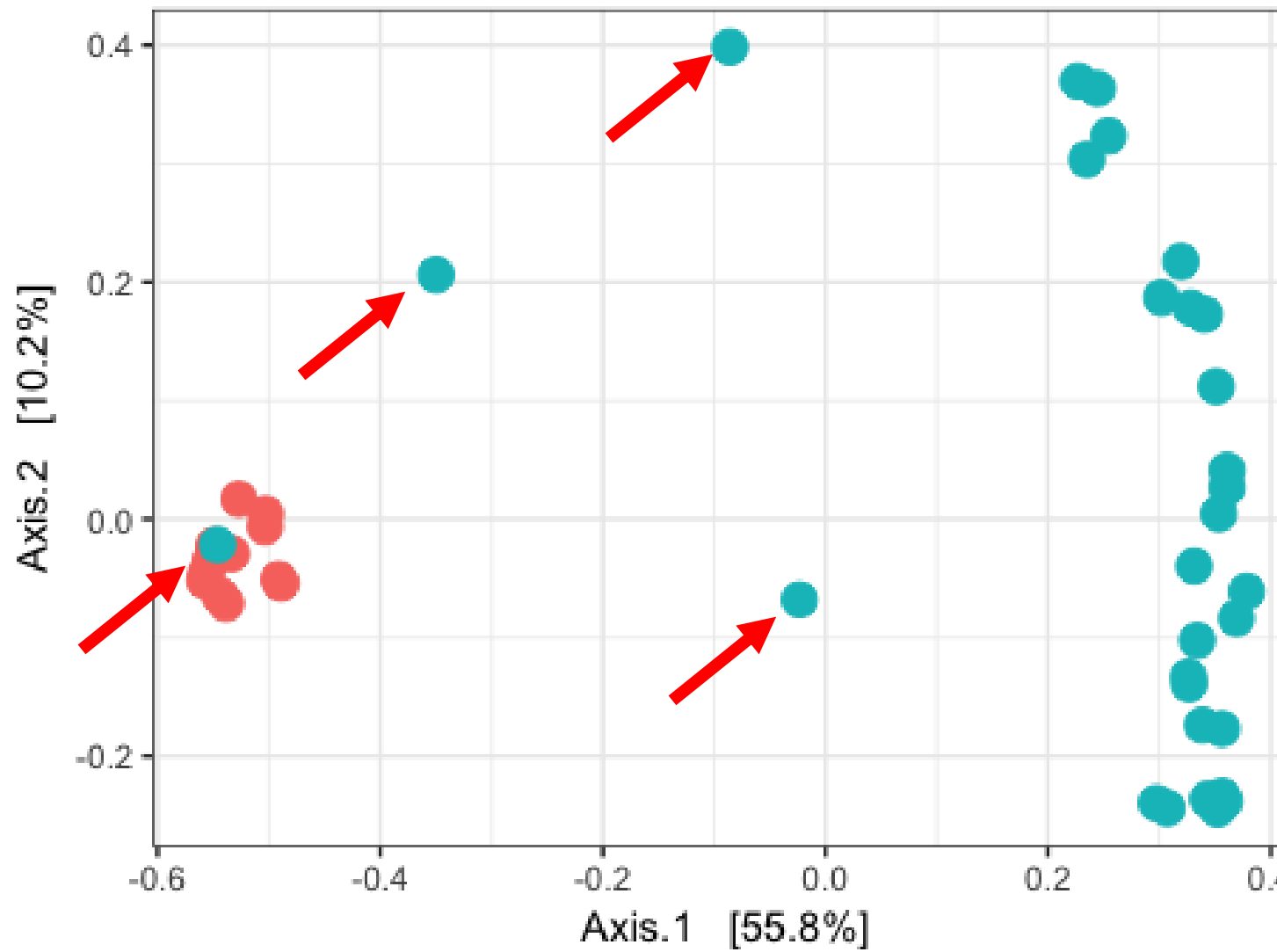


Sample Outlier Detection – Unexpectedly Low # of Sequences



Samples that “perform” unexpectedly

MDS of Bray Distances
Outlier Evaluation

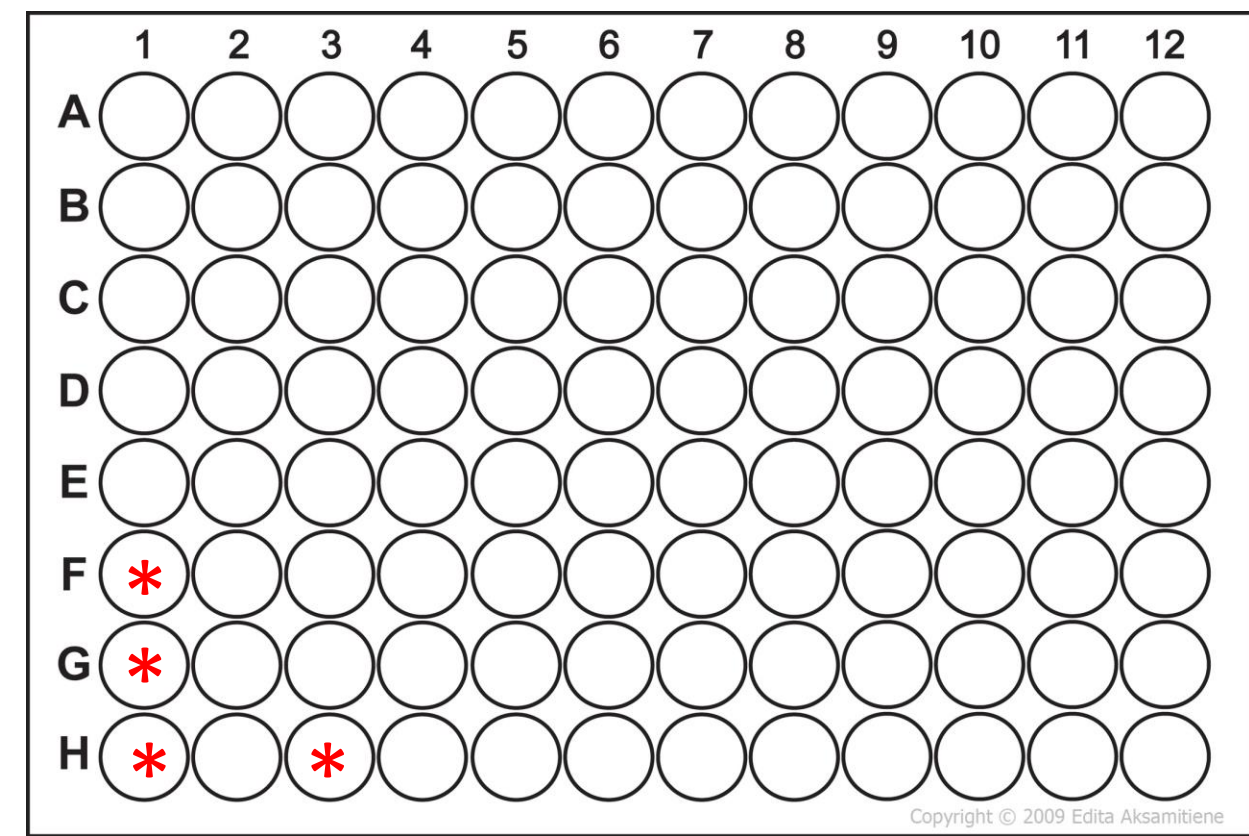
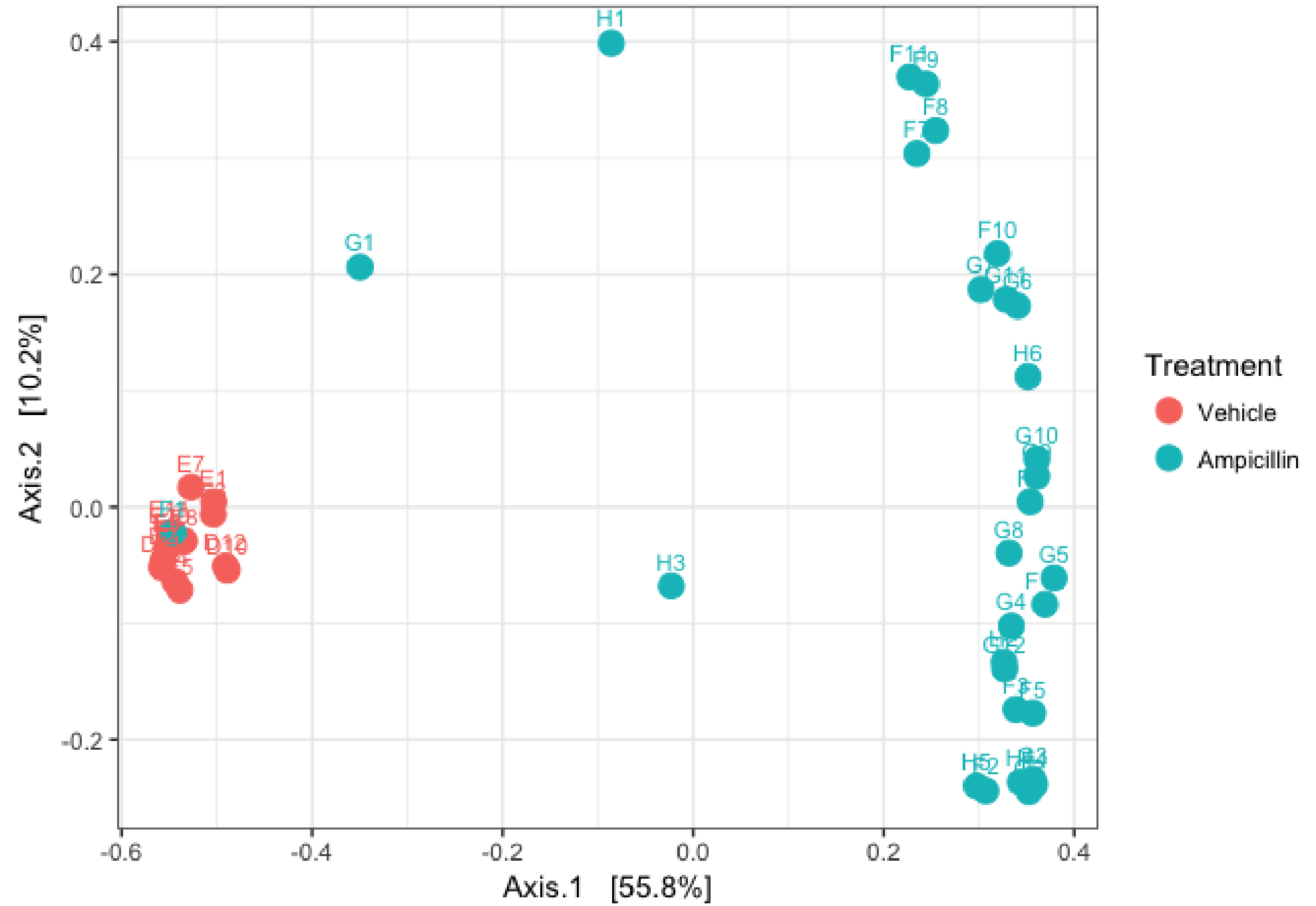


Rules of Thumb for Sample Detection and Removal

- **Justify and document!!!**
- Except in extreme cases, test how sample removal alters your downstream results. Do the experiment!
- Know your data. When are you comfortable removing a sample based on your knowledge of the system
- Explore using multiple plot types
- Include enough detail to make analysis interpretable and reproducible

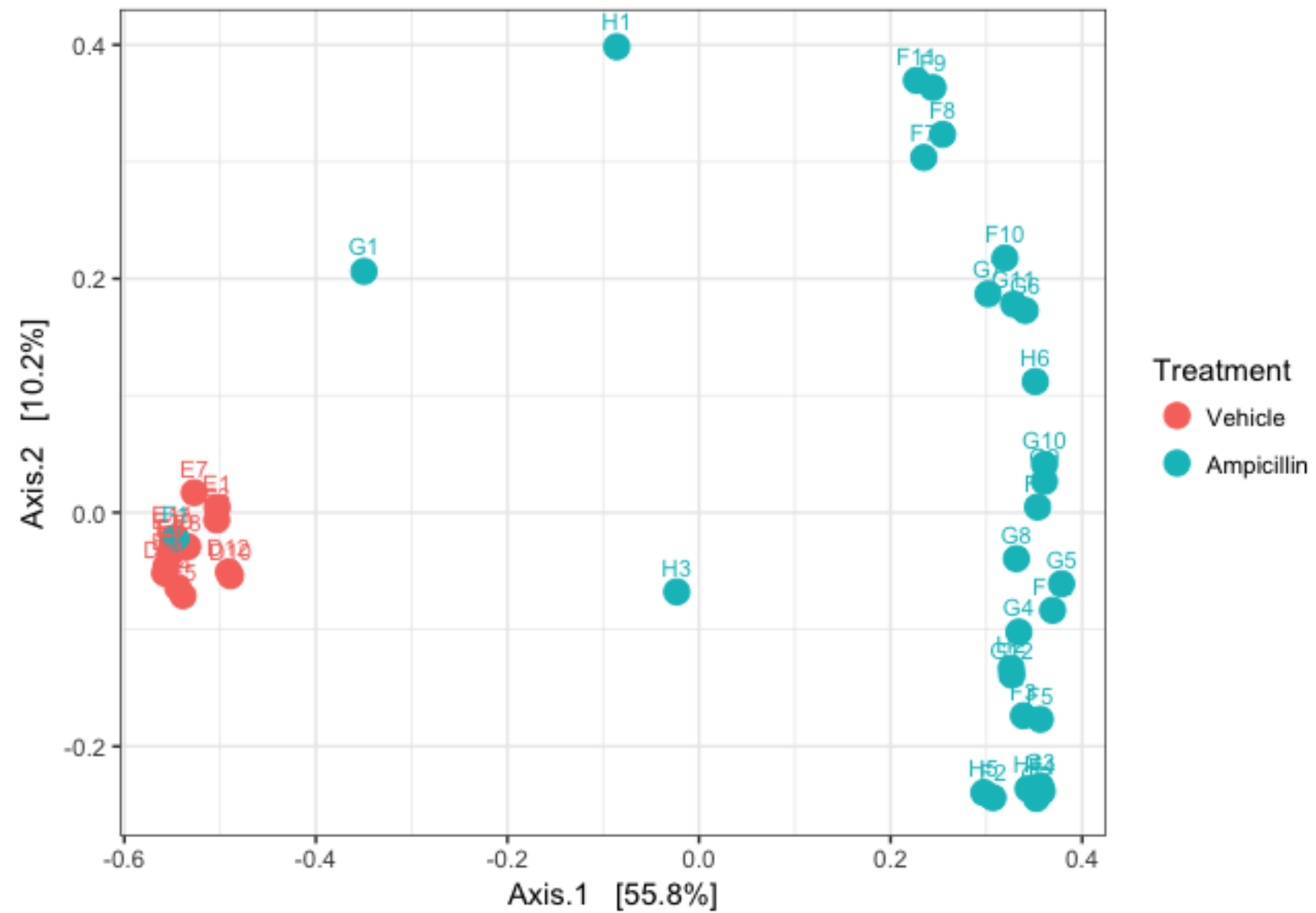
Understand your data better

MDS of Bray Distances
Outlier Evaluation

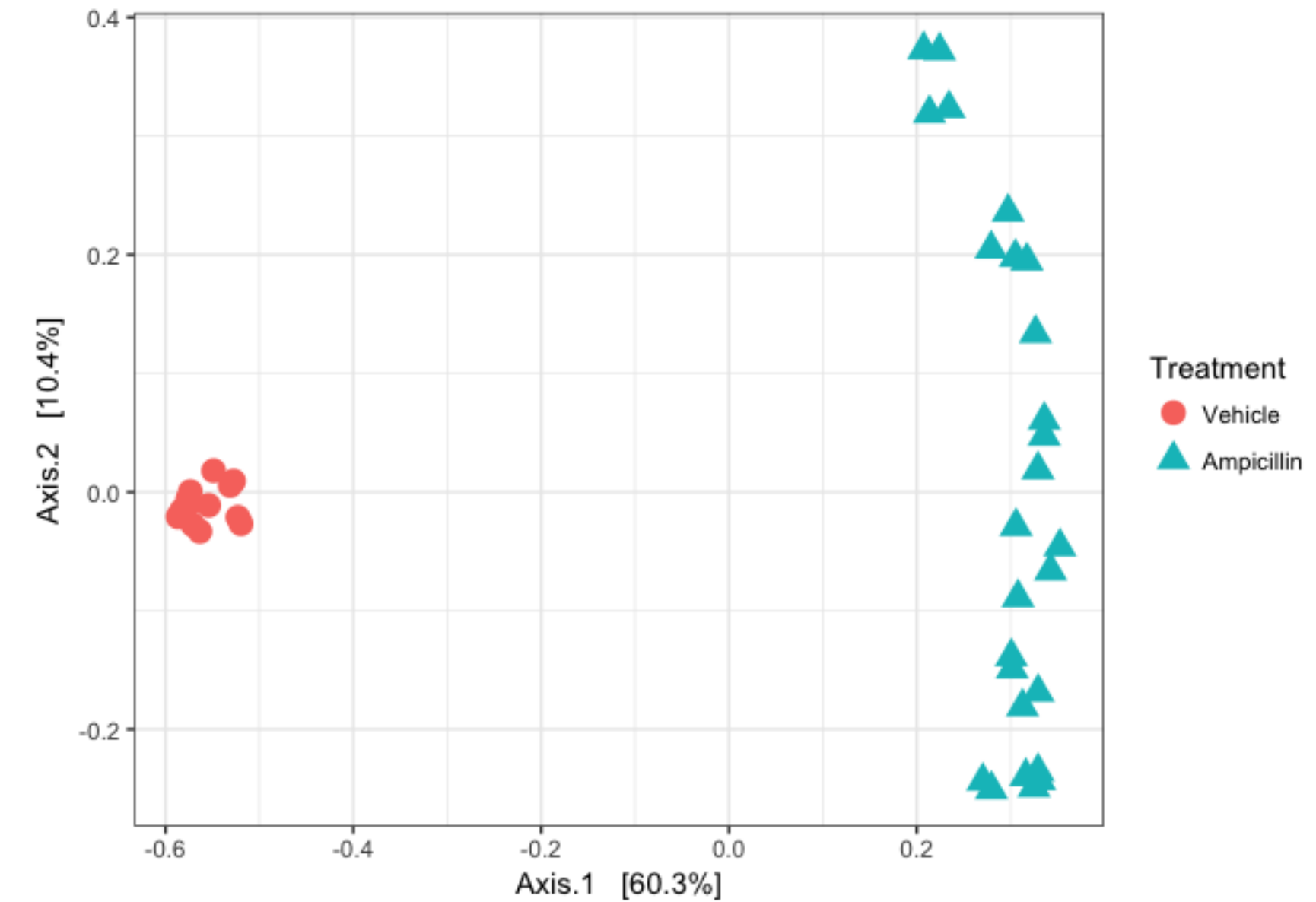


Cleaned Data

MDS of Bray Distances
Outlier Evaluation

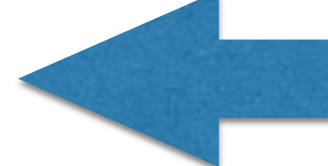


MDS of Bray Distances
Outliers Removed



Feature Outlier Detection

ID	Sample 1	Sample 2	Sample 3	Sample 4
ASV 1	0	0	2	0
ASV 2	12	8	8	456
ASV 3	112	101	98	10
ASV 4	435	435	382	3
ASV 5	76	83	68	145

... n=270


...

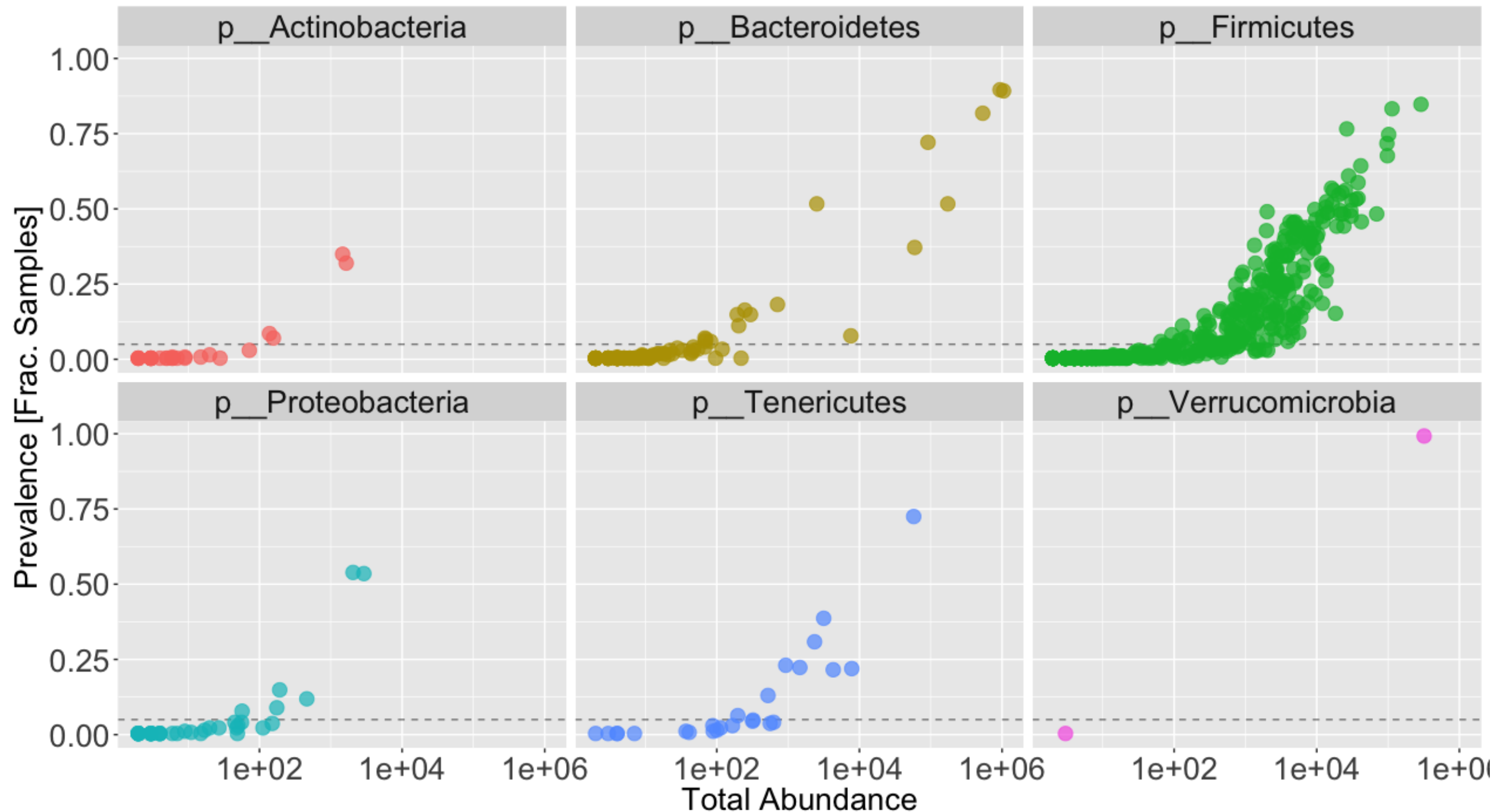
n=724

Low-abundant feature removal is commonplace

- “*We removed all taxa that were under 1% relative abundance and present in less than 3% of all samples.*”

Sequence/Taxa Outlier Detection

Filtering out low impact information



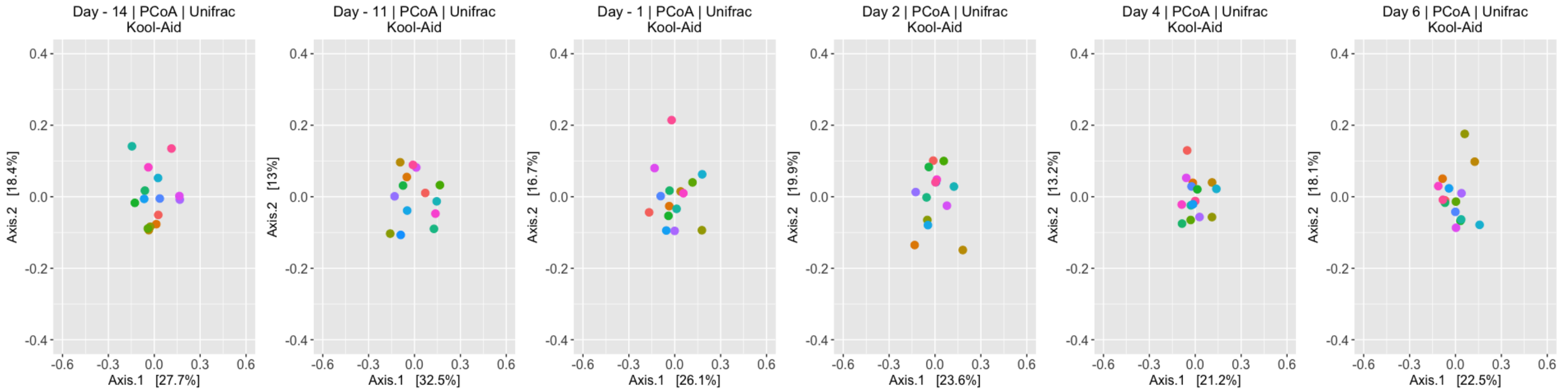
Rules of Thumb for Feature Detection and Removal

- **Justify and document!!!**
- Except in extreme cases, test how feature removal alters your downstream results. Do the experiment!
- Know your data. When are you comfortable removing a feature based on your knowledge of the system
- Explore using multiple plot types
- Include enough detail to make analysis interpretable and reproducible

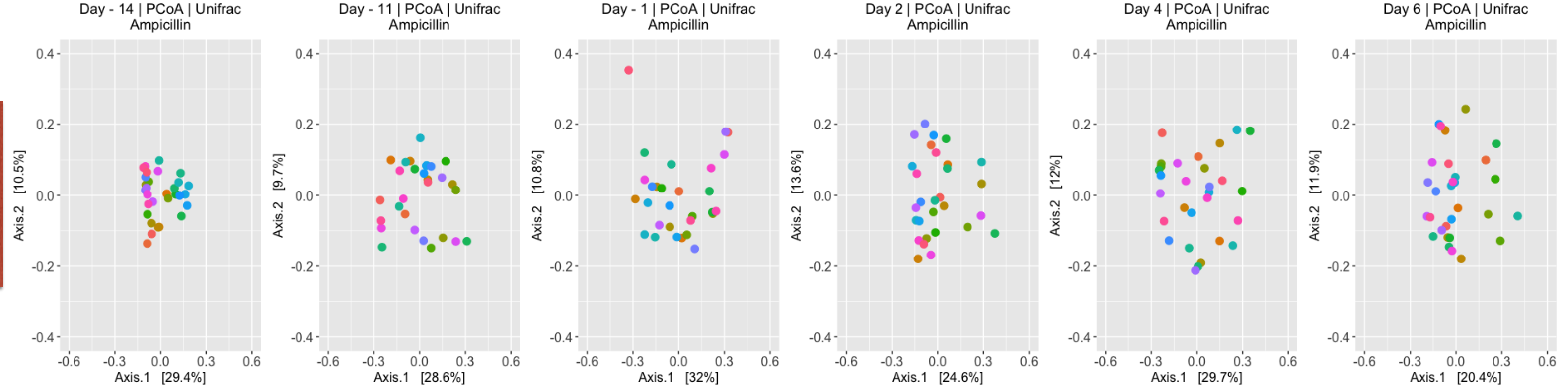
Beta Diversity Throughout the Course of the Experiment

Colored by Cage

Kool-Aid



Ampicillin



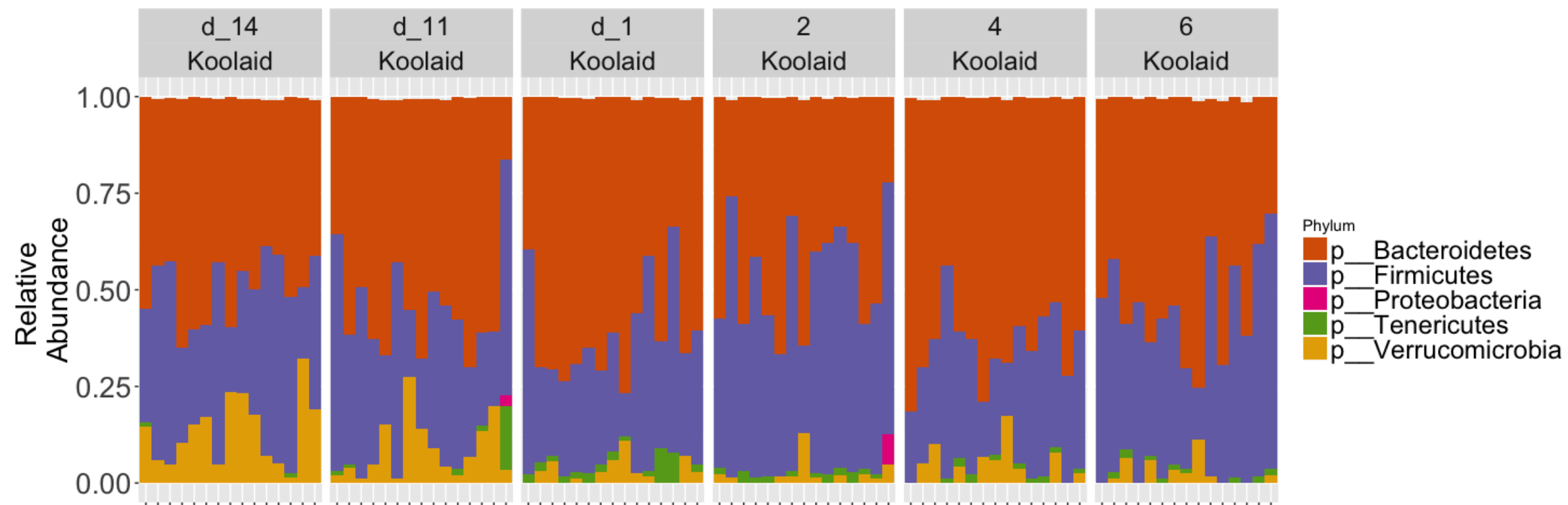
Summary

- Explore -> Document -> Test
- Does any of this really matter?
 - Sometimes?
 - Less so for community ecology measurements
 - More so for detection of differentially abundant taxa
 - Detailed exploration provides more opportunities for insights
 - Don't publish garbage data

Frequently Used 16S Analysis Techniques

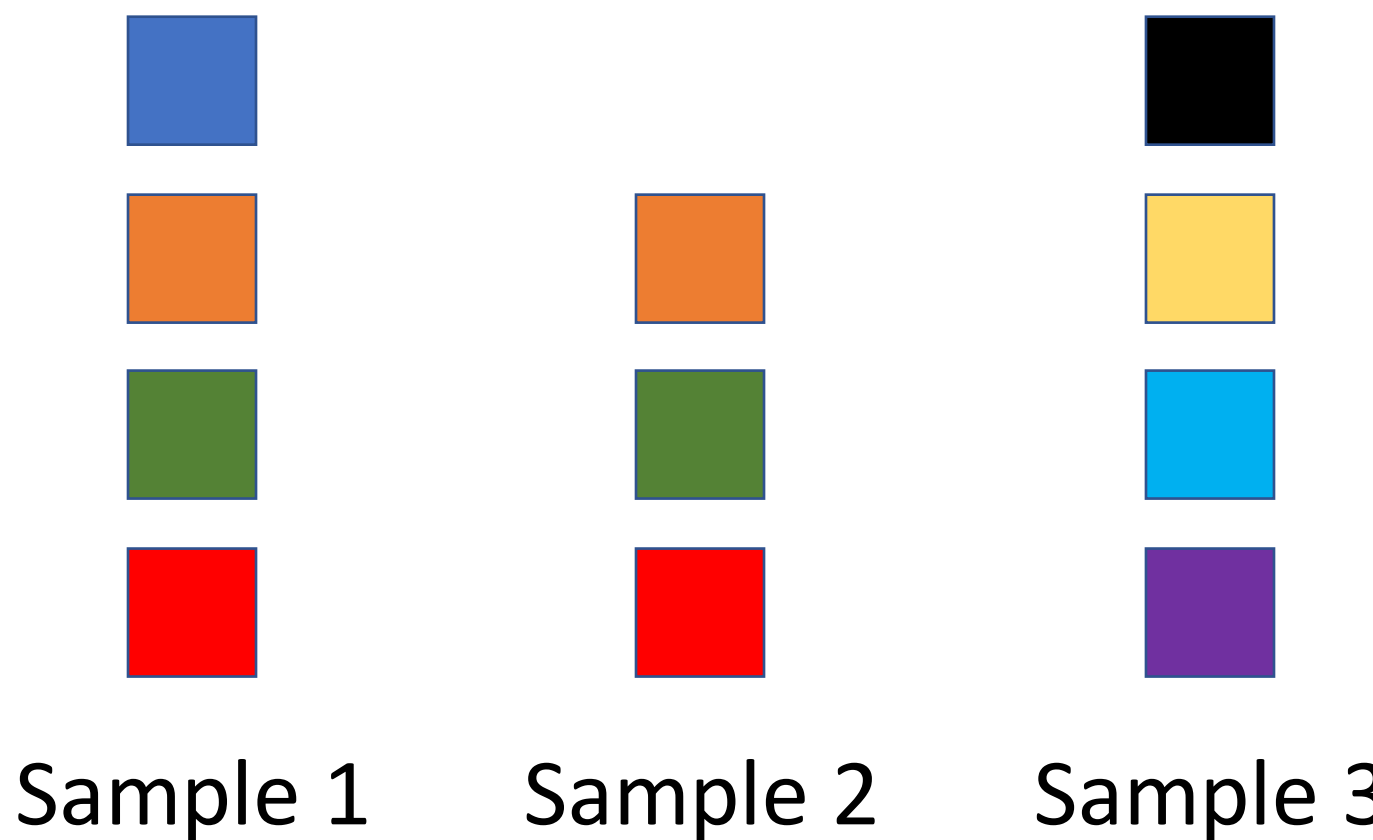
Community Composition

- Broad overview
- Nothing statistical

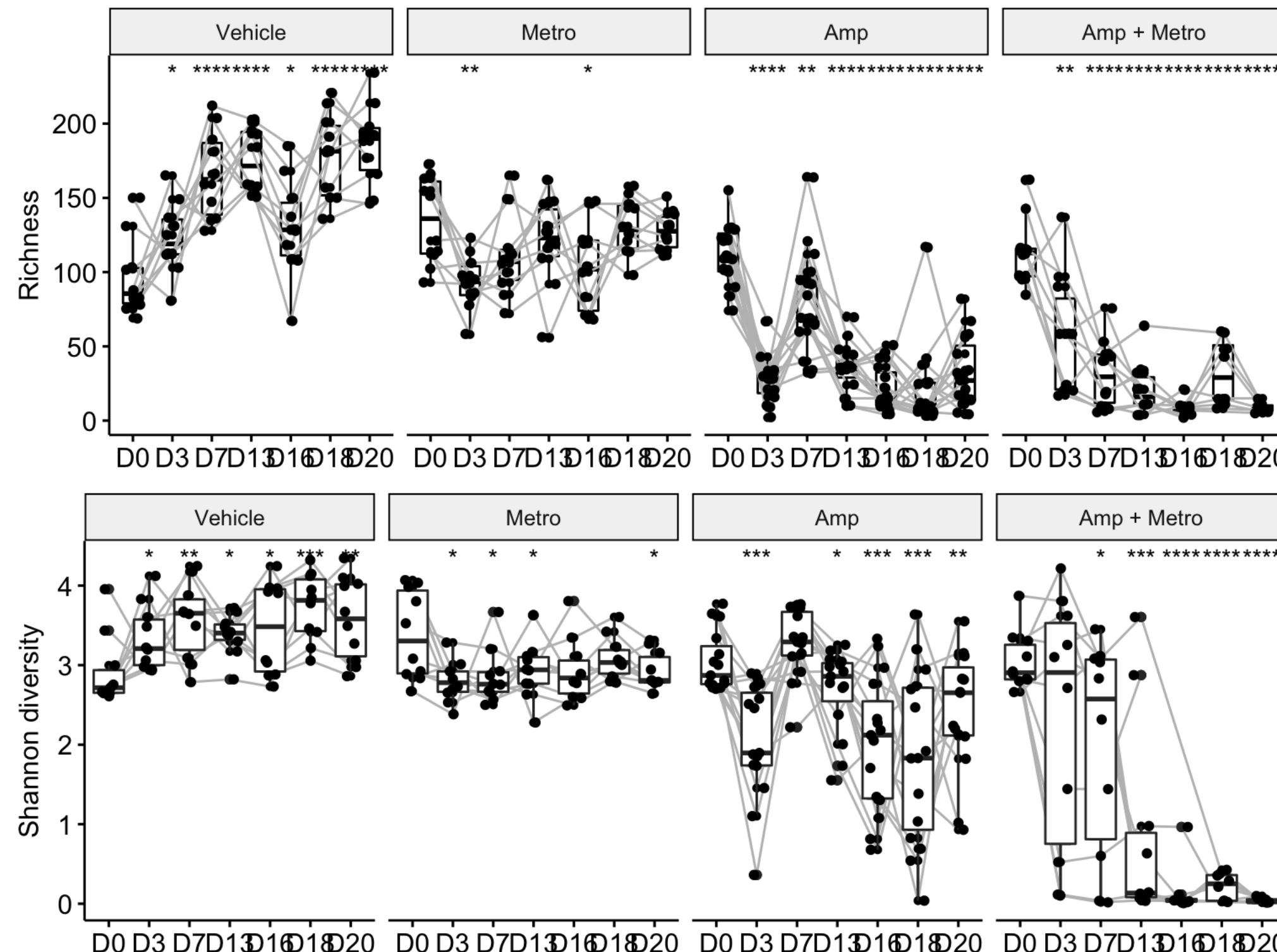


Alpha Diversity: Richness

- Richness: Number of unique taxa (ASVs) that are observed in a sample
 - Taxonomy independent
 - Abundance independent (presence / absence)
- Loads of other Alpha diversity measures (Chao1, Shannon, Simpsons, etc.)

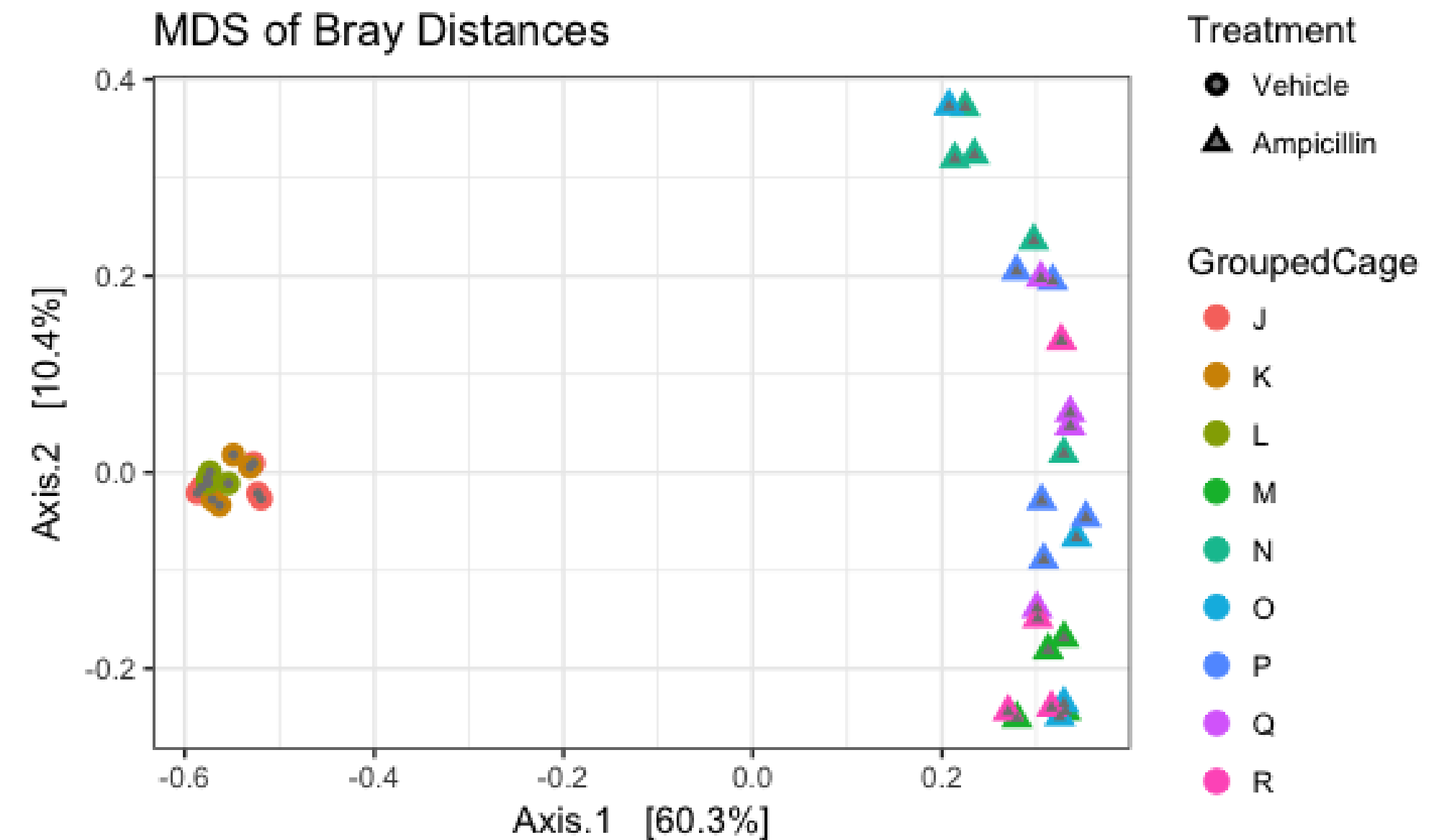


Richness Example



Beta Diversity

- Between sample similarity
 - Distance between one sample to all other samples
 - Multivariate
 - Can incorporate relative abundances or not
 - Can incorporate phylogenetic relatedness or not
 - Most frequently displayed in an ordination plot



To learn about distance measures and ordination:
<https://sites.google.com/site/mb3gustame/home>

Differential Abundance Analysis

- What specific taxa are different between study groups?
 - Lots of methods
 - DeSeq2
 - Random Forest
 - LeFse
 - ANCOM
 - Gneiss
 - ...

