Lecture course on environmental DNA metabarcoding using Claident and R: From nucleotide sequence data processing 田辺晶史 to ecological analyses

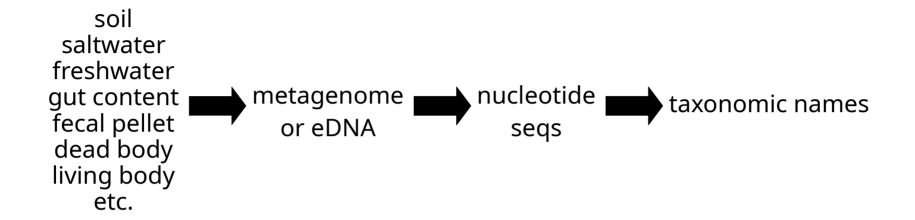
Akifumi S. Tanabe

## ClaidentとRによる

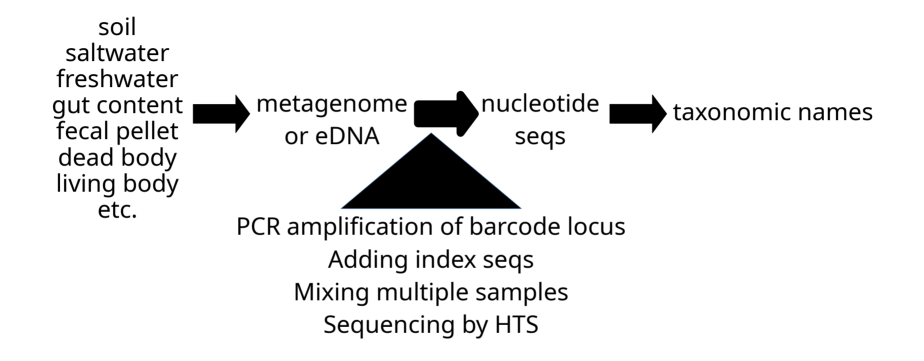
環境DNAメタバーコーディング分析講座:

塩基配列データ処理から生態学的分析まで

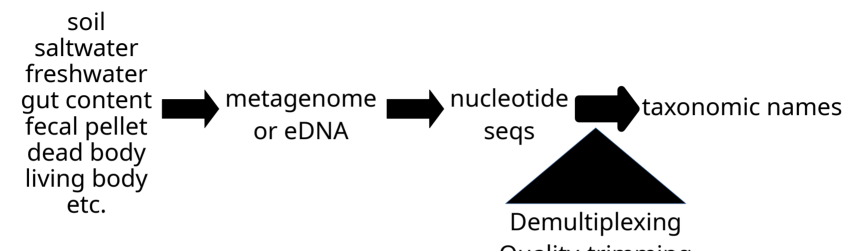
#### **Workflow of metabarcoding**



#### Molecular laboratory processes of metabarcoding



#### **Computational processes of metabarcoding**



# Claident

https://www.claident.org/

Quality-trimming
Quality-filtering
Denoising
Chimera removal
Decontamination
Clustering
Taxonomic assignment

#### Single-end sequence data analysis in Claident

- 1. Demultiplexing by clsplitseq
- 2. Evaluate sequence quality by VSEARCH via clcalcfastqstatv
- 3. Quality-trimming&filtering by VSEARCH via clfilterseqv
- 4. Denoising by DADA2 via cldenoiseseqd
- 5. Removing chimeras by UCHIME3 via clremovechimev
- 6. Removing contaminants by clremovecontam
- 7. Additional clustering by VSEARCH via clclassseqv (Optional)
- 8. Assigning taxonomy by clmakecachedb, clidentseq, classigntax
- 9. Additional taxonomy processing by clmergeassign, clfillassign
- 10.Summarizing results by clsumclass, clsumtaxa

#### Overlapped paired-end sequence data analysis in Claident

- 1. Demultiplexing by clsplitseq
- 2. Concatenating pairs by VSEARCH via clconcatpairv
- 3. Quality-filtering by VSEARCH via clfilterseqv
- 4. Denoising by DADA2 via cldenoiseseqd
- 5. Removing chimeras by UCHIME3 via clremovechimev
- 6. Removing contaminants by clremovecontam
- 7. Additional clustering by VSEARCH via clclassseqv (Optional)
- 8. Assigning taxonomy by clmakecachedb, clidentseq, classigntax
- 9. Additional taxonomy processing by clmergeassign, clfillassign
- 10.Summarizing results by clsumclass, clsumtaxa

#### Non-overlapped paired-end sequence data analysis in Claident

- 1. Demultiplexing by clsplitseq
- 2. Evaluate sequence quality by VSEARCH via clcalcfastqstatv x2
- 3. Quality-trimming by VSEARCH via clfilterseqv x2
- 4. Joining pairs by VSEARCH via clconcatpairv
- 5. Quality-filtering by VSEARCH via clfilterseqv
- 6. Denoising by DADA2 via cldenoiseseqd
- 7. Removing chimeras by UCHIME3 via clremovechimev
- 8. Removing contaminants by clremovecontam
- 9. Additional clustering by VSEARCH via clclassseqv (Optional)
- 10.Dividing pairs by cldivseq
- 11.Assigning taxonomy by clmakecachedb,clidentseq,classigntax x2
- 12.Additional taxonomy processing by clmergeassign, clfillassign
- 13.Summarizing results by clsumclass, clsumtaxa

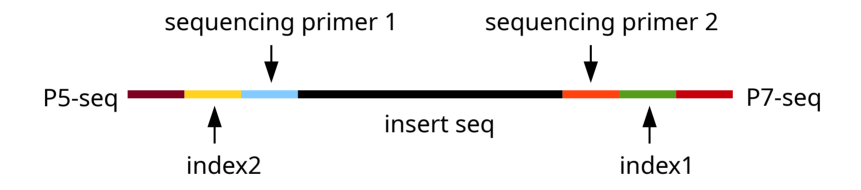
### Analysis demonstration of overlapped paired-end data using Claident

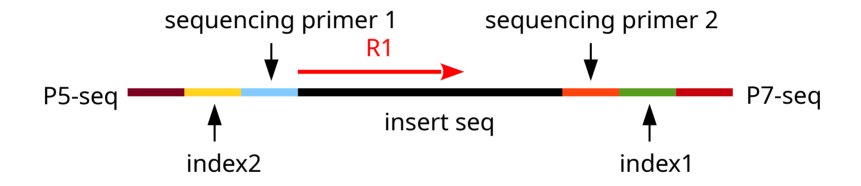
- Prerequisites to run Claident
  - Debian/Ubuntu/Linux Mint, RedHat/CentOS
  - Claident+BLASTDB+TaxonomyDB+UCHIMEDB
  - Code from https://github.com/astanabe/ClaidentTutorial
- Prerequisites to learn about analyses using Claident and R
  - Code from https://github.com/astanabe/ClaidentTutorial
    - This includes simulated data and all results

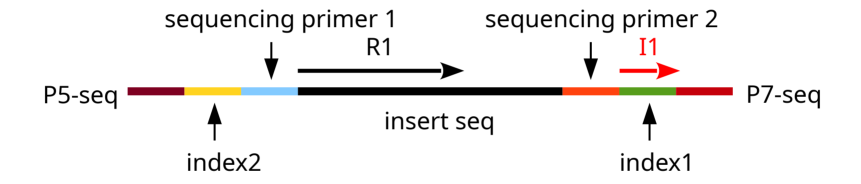
#### **Chapter 0: Simulated data creation**

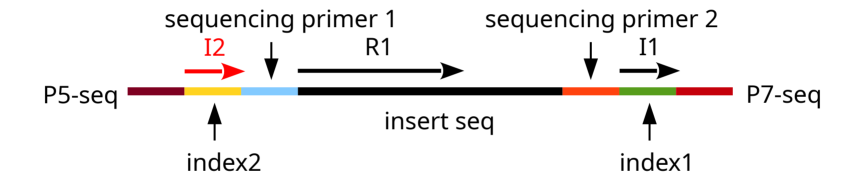
- 1. Download complete mitogenome seq data of fishes
- 2. Extract 12S rRNA region
- 3. Run in silico PCR using MiFish-U primer by ecoPCR and obtain amplicons
- 4. Cluster amplicon seqs and pick representative seqs
- 5. Randomly pick 50 seqs from all repseqs (1st sample)
- 6. Randomly pick 40 seqs from previous sample and randomly pick 10 seqs from all repseqs except for previous sample seqs (2nd-20th sample)
- 7. Pick all sequences from all 1st-20th samples for blank (1st-4th blank)
- 8. Generate 500 paired-end seqs for each picked seqs by ART for samples
- 9. Generate 50 paired-end seqs for each picked seqs by ART for blanks
- 10.Generate dual index seqs based on given fasta files

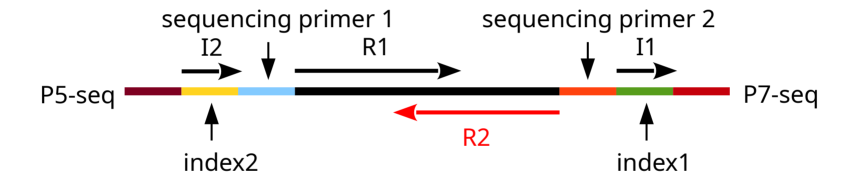
#### Interlude: The structure of Illumina dual-index library



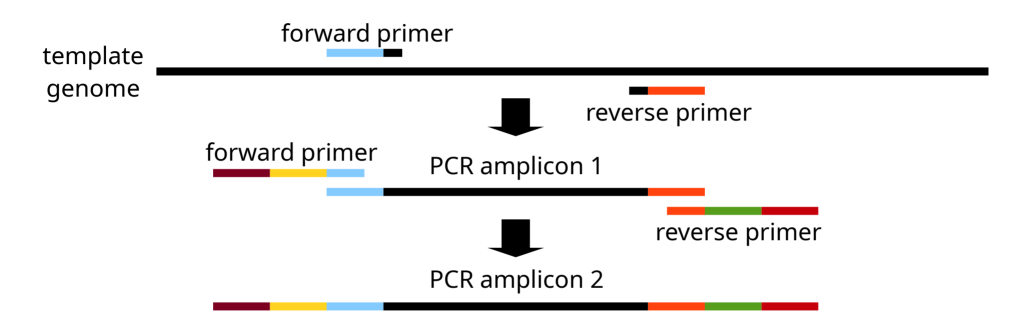








#### Interlude: Preparation of Illumina dual-index library



By 8 forward index primers and 12 reverse index primers, 96 samples can be distinguished (combinatorial dual-indexing).

### Interlude: Dual-index design of simulated data

reverse index (index1)	TTGCAGGT	Sample01	Sample07	not used	not used
	CAAGGAAC	Sample02	Sample08	not used	not used
	AGATCTGG	Sample03	Sample09	not used	not used
	TCACACTT	Sample04	Sample10	not used	not used
	GATCATGG	Sample05	Sample11	not used	not used
	AGACATGA	Sample06	Sample12	not used	not used
	GTGAGTTG	not used	not used	Sample13	Sample19
	AGTCTGTT	not used	not used	Sample14	Sample20
	AACCAACC	not used	not used	Sample15	Blank01
	AGTGTGCA	not used	not used	Sample16	Blank02
	CATGTCGA	not used	not used	Sample17	Blank03
	CGAGACTT	not used	not used	Sample18	Blank04
		AACCTCTC	GTGACTCT	GATCACCA	CTTCACAT

forward index (index2)

#### **Chapter 1: Demultiplexing**

- Inputs
- Undemultiplexed\_R1\_001.fastq.xz
- Undemultiplexed\_I1\_001.fastq.xz
- Undemultiplexed\_I2\_001.fastq.xz
- Undemultiplexed\_R2\_001.fastq.xz

in 01\_RawSequences

in top directory

- index1.fasta
- index2.fasta
- forwardprimer.fasta
- reverseprimer.fasta

- Outputs
- ClaidentTutorial\_\_\*\_MiFish.forw ard.fastq.xz
- ClaidentTutorial\_\_\*\_MiFish.rever se.fastq.xz
  - Sample\*
  - Blank\*
  - NNNNNNN+NNNNNNN

in PairedEnd\_02a\_DemultiplexedSequences

Chapter 1: Demultiplexing

Launch Terminal

#### **Chapter 2: Concatenating pairs**

- Inputs
- ClaidentTutorial\_\_\*\_MiFish.forwa rd.fastq.xz
- ClaidentTutorial\_\_\*\_MiFish.revers e.fastq.xz
  - Sample\*
  - Blank\*
  - NNNNNNN+NNNNNN

in PairedEnd\_02a\_DemultiplexedSequences

- Outputs
- ClaidentTutorial\_\_\*\_MiFish.fastq.
   xz
  - Sample\*
  - Blank\*

in OverlappedPairedEnd\_03\_ConcatenatedSequences

Chapter 2: Concatenating pairs

Switch to Terminal

#### **Chapter 3: Quality-filtering**

- Inputs
- ClaidentTutorial\_\_\*\_MiFish.fastq.
   xz
  - Sample\*
  - Blank\*

- Outputs
- ClaidentTutorial\_\_\*\_MiFish.fastq.
   xz
  - Sample\*
  - Blank\*
  - NNNNNNN+NNNNNNN

in OverlappedPairedEnd\_04\_FilteredSequences

Switch to Terminal

**Chapter 3: Quality-filtering** 

#### **Chapter 4: Denoising**

- Inputs
- ClaidentTutorial\_\_\*\_MiFish.fastq.
   xz
  - Sample\*
  - Blank\*
  - NNNNNNNN+NNNNNNN

in OverlappedPairedEnd\_04\_FilteredSequences

- Outputs
- denoised.fasta
- denoised.otu.gz
- denoised.tsv
- plotErrors.pdf
- runDADA2.R

in OverlappedPairedEnd\_05\_DenoisedSequences

**Chapter 4: Denoising** 

Switch to Terminal

**Interlude: Methods in DADA2** 

observed number

ACCTCTCGATATCGAGATGAGGCT 10000

ACCTCTTGATATCGAGATGAGGCT 10

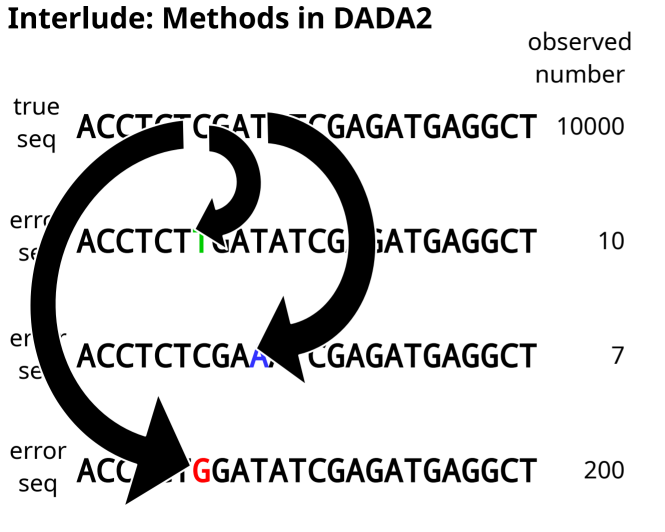
ACCTCTCGAAATCGAGATGAGGCT 7

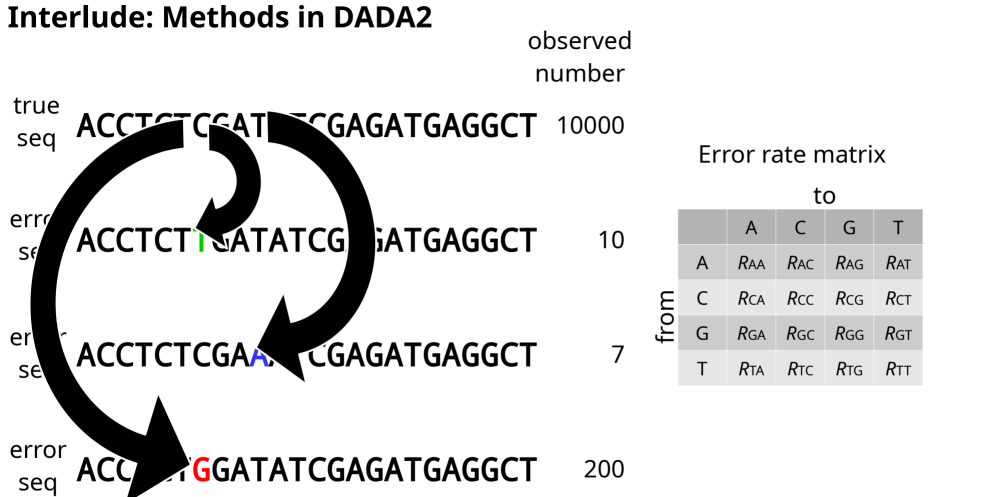
ACCTCTGGATATCGAGATGAGGCT 200

**Interlude: Methods in DADA2** observed number true ACCTCTCGATATCGAGATGAGGCT 10000 **ACCTCTTGATATCGAGATGAGGCT** 10

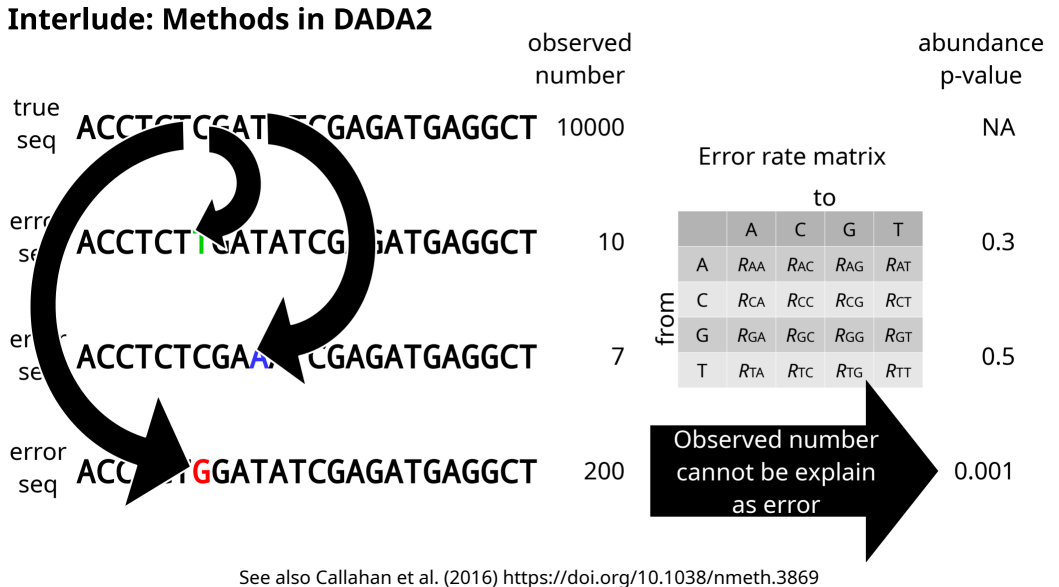
**ACCTCTCGAAATCGAGATGAGGCT ACCTCTGGATATCGAGATGAGGCT** 

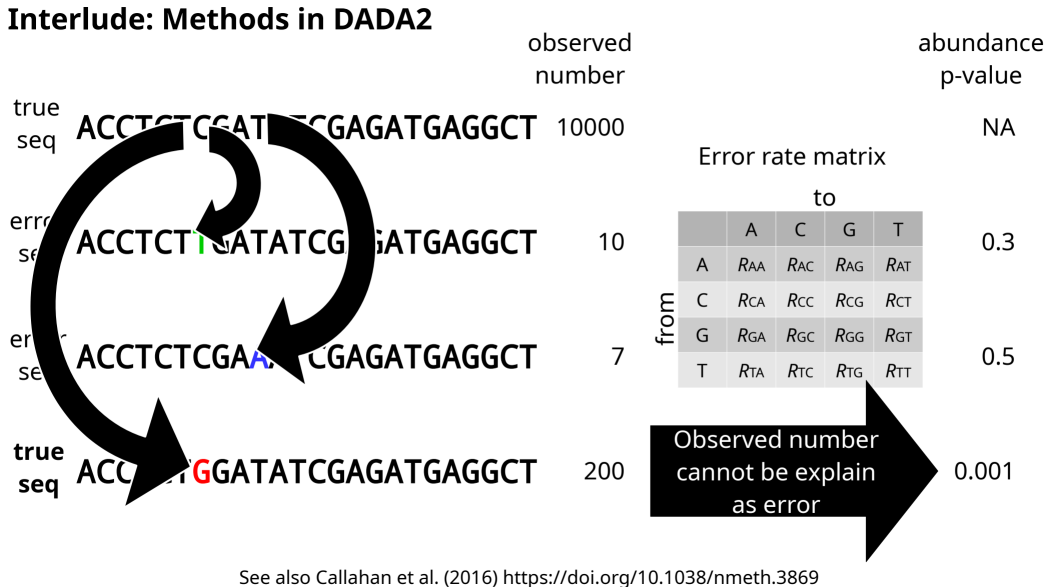
200











#### **Chapter 5: Chimera removal**

- Inputs
- denoised.fasta
- denoised.otu.gz

in OverlappedPairedEnd\_05\_DenoisedSequences

- Outputs
- nonchimeras.fasta
- nonchimeras.otu.gz
- nonchimeras.tsv
- \* borderline.fasta
- \*\_chimeras.fasta
- \*\_nonchimeras.fasta
- \*\_uchimealns.txt
- \*\_uchimeout.txt

in OverlappedPairedEnd\_06\_NonchimericSequences

Chapter 5: Chimera removal

Switch to Terminal

#### **Chapter 6: Removing index-hopped sequences**

- Inputs
- nonchimeras.fasta
- nonchimeras.otu.gz
   in OverlappedPairedEnd\_06\_NonchimericSequences
- index1.fasta
- index2.fasta

in top directory

- Outputs
- decontaminated.fasta
- decontaminated.otu.gz
- decontaminated.tsv

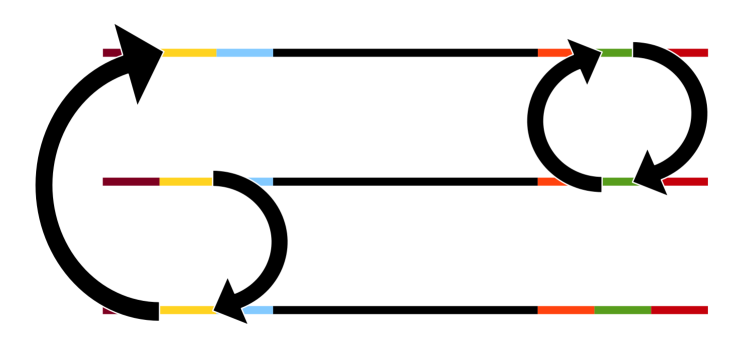
in OverlappedPairedEnd\_07\_NonhoppedSequences

Chapter 6: Removing index-hopped sequences

Switch to Terminal

Interlude: Index can hop into another amplicon within a flowcell!

Index-hopping potentially causes sequence misassignments! Especially in newer models! OMG!



#### Interlude: Detecting index-hopping using unused index combinations

	AACCTCTC	GTGACTCT	GATCACCA	CTTCACAT
CGAGACTT	not used	not used	Sample18	Blank04
CATGTCGA	not used	not used	Sample17	Blank03
AGTGTGCA	not used	not used	Sample16	Blank02
AACCAACC	not used	not used	Sample15	Blank01
AGTCTGTT	not used	not used	Sample14	Sample20
GTGAGTTG	not used	not used	Sample13	Sample19
AGACATGA	Sample06	Sample12	not used	not used
GATCATGG	Sample05	Sample11	not used	not used
TCACACTT	Sample04	Sample10	not used	not used
AGATCTGG	Sample03	Sample09	not used	not used
CAAGGAAC	Sample02	Sample08	not used	not used
TTGCAGGT	Sample01	Sample07	not used	not used

reverse index (index1)

- 1. Count abundances
- 2. Collect abundances of a sample + "not used"
- 3. Test whether sample abundance is outlier or not
- 4. If it's not outlier, it's determined as hopped

forward index (index2)

See also Esling et al. (2015) https://doi.org/10.1093/nar/gkv107

### **Chapter 7: Removing contaminant sequences**

- Inputs
- decontaminated.fasta
- decontaminated.otu.gz
   in OverlappedPairedEnd\_07\_NonhoppedSequences
- blanklist.txt

in top directory

- Outputs
- decontaminated.fasta
- decontaminated.otu.gz
- decontaminated.tsv

in OverlappedPairedEnd\_08\_DecontaminatedSequences

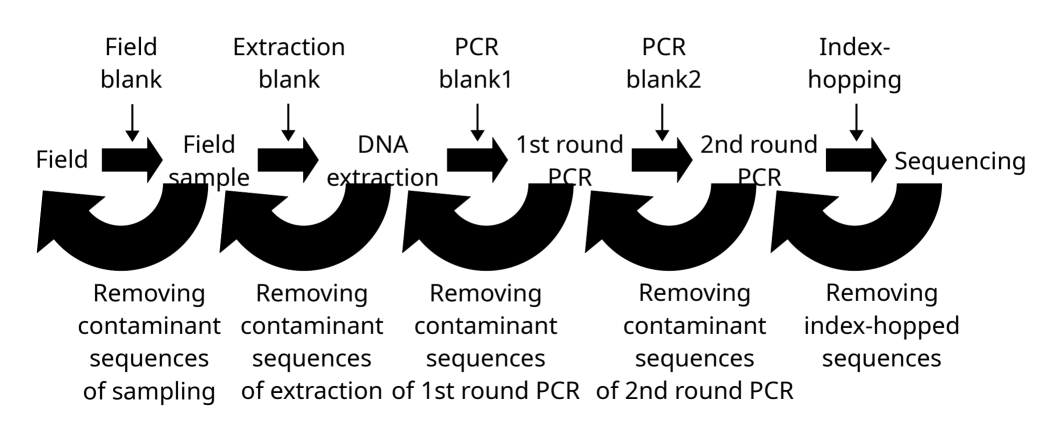
Chapter 7: Removing contaminant sequences

#### Interlude: Detecting contaminants using blank samples

- 1. Count abundances
- 2. Collect abundances of a sample + associated blanks
- 3. Test whether sample abundance is outlier or not
- 4. If it's not outlier, it's determined as contaminant

#### Interlude: Multistep contamination and multistep decontamination

My recommendation is index-hopping removal + the other contaminant removal. However, the best practice has been still unknown.



#### Interlude: Study purpose and decontamination

- Non-decontaminated metabarcoding results contain contaminants
- Decontamination should be applied?
  - If you want to maximize detection power, NO. Decontamination potentially misidentify true sequence as contaminant
  - If you want to minimize misdetection, YES. Lack of decontamination may cause many misdetection
  - If you want to analyse community composition, UNKNOWN. Because abundances of contaminants may be low in many cases, their effects to analysis may be low. However, whether abundances of contaminants are really low or not IN YOUR DATA is unknown.

#### **Chapter 8: Additional clustering**

- Inputs
- decontaminated.fasta
- decontaminated.otu.gz

in OverlappedPairedEnd\_08\_DecontaminatedSequences

- Outputs
- clustered.fasta
- clustered.otu.gz
- clustered.tsv

in OverlappedPairedEnd\_09\_ClusteredSequences

Chapter 8: Additional clustering

#### **Chapter 9: Taxonomic assignment**

- Inputs
- clustered.fasta

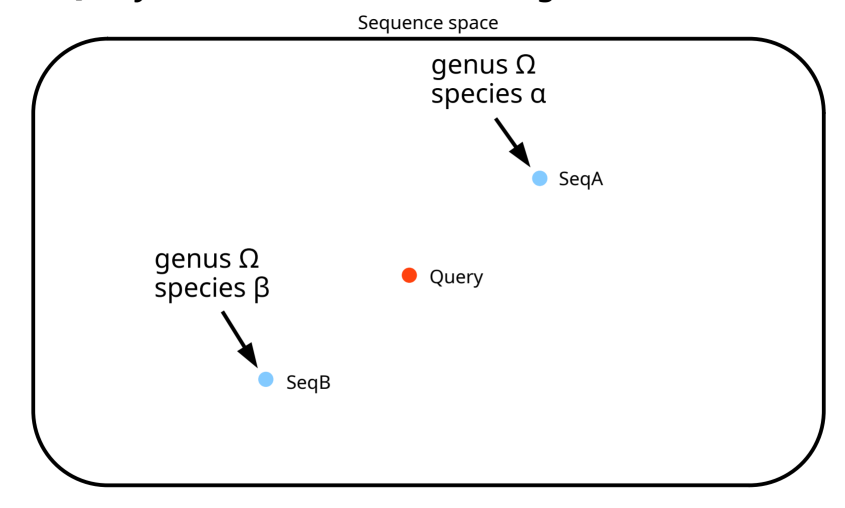
 $in\ Overlapped Paired End\_09\_Clustered Sequences$ 

- Outputs
- neighborhoods\_1nn\_\*.txt
- neighborhoods\_qc\_\*.txt
- taxonomy\_1nn\_\*.tsv
- taxonomy\_qc\_\*.tsv
- taxonomy\_merged.tsv
- taxonomy\_merged\_filled.tsv

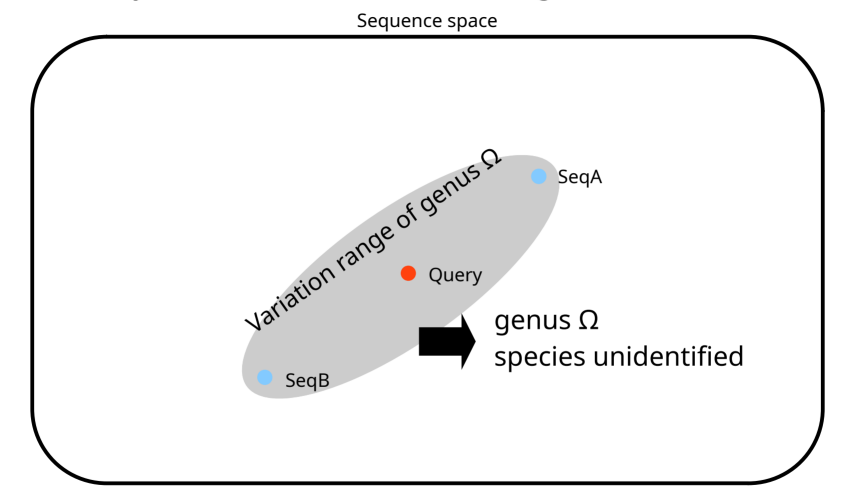
in OverlappedPairedEnd\_10\_ClaidentResults

Chapter 9: Taxonomic assignment

#### Interlude: Query-centric auto-*k*-nearest neighbor method



#### Interlude: Query-centric auto-*k*-nearest neighbor method



### Interlude: Which method should be used for taxonomic assignment?

- If reference database is imperfect (most cases), QCauto shows the best balance between less misidentification and less successful identification
- If reference database is perfect or nearly perfect, 1-NN is the best.
   However, whether the reference database is really perfect or not should not be known by anyone

#### **Interlude: Ready-made reference databases**

- Installed to INSTALLPATH/share/claident/blastdb
- overall\_class, overall\_order, overall\_family
  - Subset of NCBI nt including class, order or family level identified seqs
- \*\_genus
  - Subset of overall\_\* including genus level identified seqs
- \*\_species\_wsp
  - Subset of overall\_\* including species level identified seqs
- \*\_species
  - Subset of overall\_\* including species level identified seqs except for the seqs which have "sp." at the tail in species name
- \*\_species\_wosp
  - Subset of overall\_\* including species level identified seqs except for the seqs which have "sp." in species name

## Interlude: Taxonomic infomation reliability in reference databases

- \*\_species\_wosp>\*\_species>\*\_species\_wsp>\*\_genus>\*\_family>\*\_order>\*\_class
  - Because the seqs which only have higher level taxonomic info likely to be identified based on closest INSD seqs, such taxonomic info are less reliable
  - Because the seqs identified as "sp." is not strictly identified or such species are undescribed, such taxonomic info are less reliable

#### **Interlude: Which reference database should be used?**

- overall\_species\_wosp is recommended in most cases because the seqs lacking lower level taxonomic info likely to be less reliable
- The other overall\_\* are recommended if you want to minimize
   "unidentified" in \* level and can rolerate misidentification in lower level
- The others are recommended for screening or PCs lacking enough amount of memory

#### **Interlude: Merging of taxonomy**

- More reliable taxonomy should be preferred but less reliable taxonomy which reached to lower taxonomic level could be tolerated
- The best balance between reliability and identifiability can be achieved by merging taxonomy from overall\_species\_wosp and the other overall \*

#### **Chapter 10: Making summary tables**

- Inputs
- clustered.tsv

 $in\ Overlapped Paired End\_09\_Clustered Sequences$ 

taxonomy\_merged\_filled.tsv

in OverlappedPairedEnd\_10\_ClaidentResults

- Outputs
- sample\_otu\_matrix\_fishes.tsv
- sample\_species\_matrix\_fishes.tsv
- sample\_top50species\_nreads\_fis hes.tsv
- sample\_top50family\_nreads\_fish es.tsv
- sample\_species\_nreads\_fishes.tsv
- sample\_family\_nreads\_fishes.tsv
   in OverlappedPairedEnd\_10\_ClaidentResults

**Chapter 10: Making summary tables** 

#### **Chapter 11: Plotting community structure**

- Inputs
- sample\_top50species\_nreads\_fish es.tsv
- sample\_top50family\_nreads\_fishe s.tsv
- sample\_species\_nreads\_fishes.tsv
- sample\_family\_nreads\_fishes.tsv

in OverlappedPairedEnd\_10\_ClaidentResults

- Outputs
- barplottop50species.pdf
- barplottop50family.pdf
- heatmapspecies.pdf
- heatmapfamily.pdf

in OverlappedPairedEnd\_11\_RAnalysisResults

Chapter 11: Plotting community structure

#### **Chapter 12: Plotting sampling/sequencing coverage**

- Inputs
- sample\_species\_matrix\_fishes.tsv

in OverlappedPairedEnd\_10\_ClaidentResults

- Outputs
- specaccum.pdf
- rarecurve.pdf

in OverlappedPairedEnd\_11\_RAnalysisResults

Community (data.frame)

in R workspace

**Chapter 12: Plotting sampling/sequencing coverage** 

#### **Chapter 12: Applying coverage-based rarefaction**

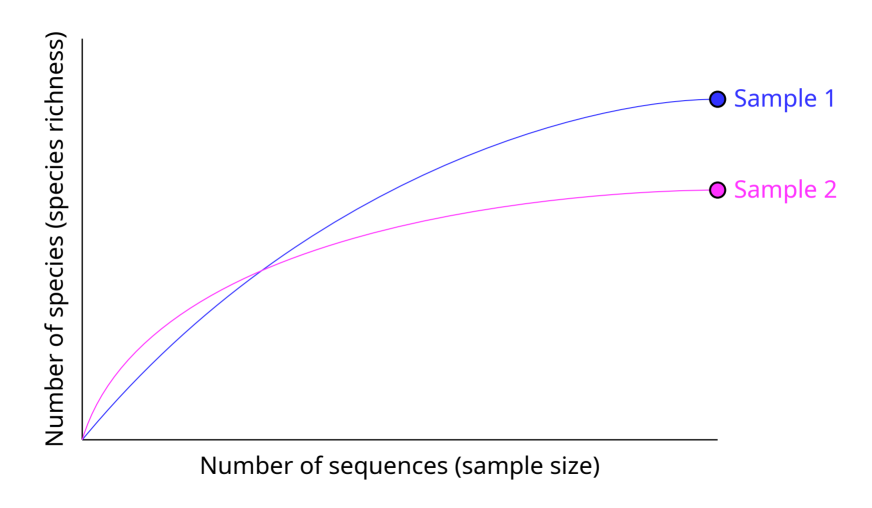
- Inputs
- Community (data.frame)
- in R workspace
- Outputs
- RarefiedCommunity (data.frame)
- BinaryRarefiedCommunity (d.f.)

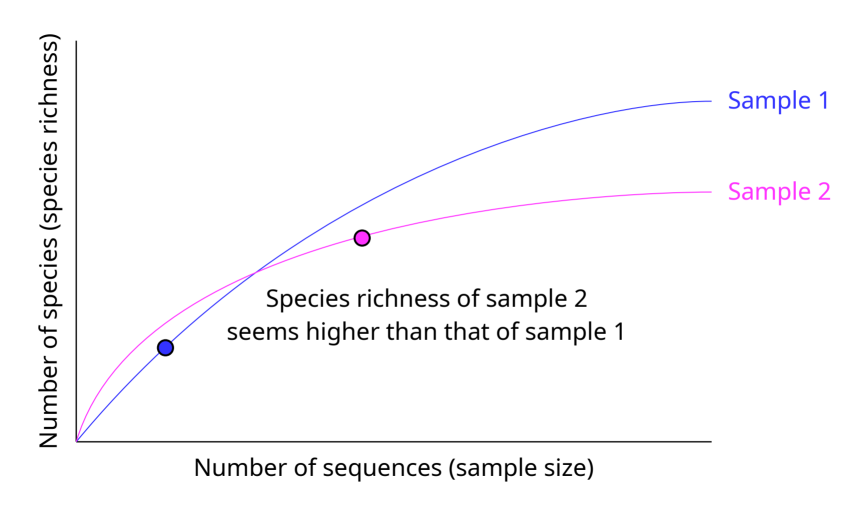
in R workspace

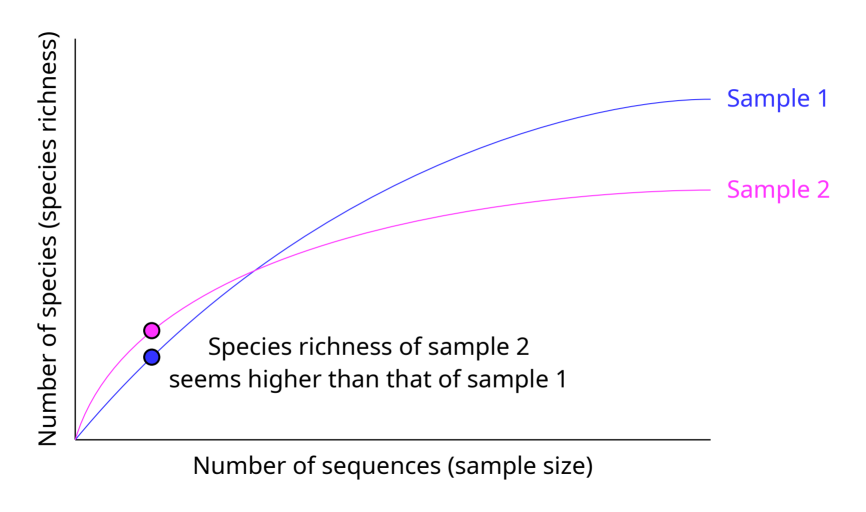
- RarefiedCommunity.tsv
- BinaryRarefiedCommunity.tsv

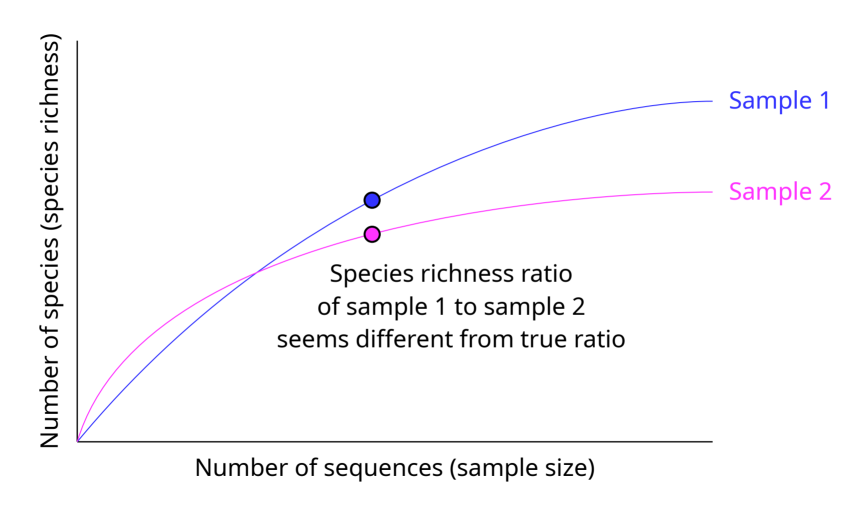
in OverlappedPairedEnd\_11\_RAnalysisResults

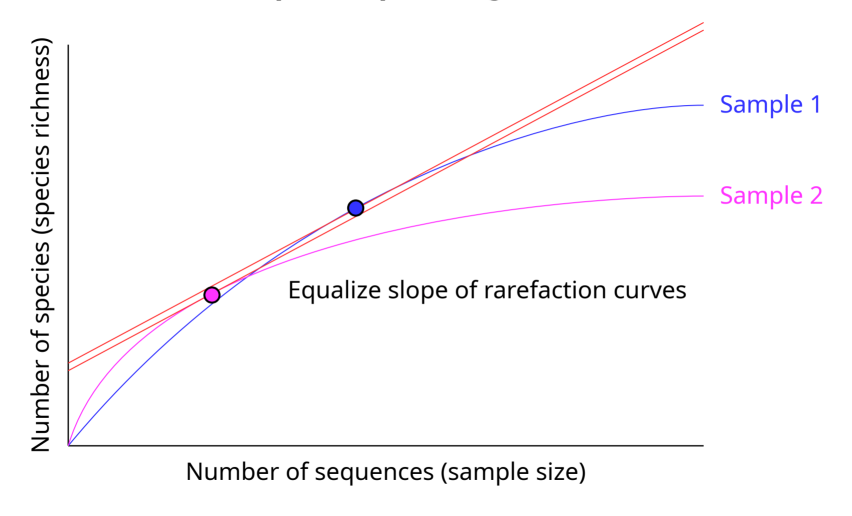
**Chapter 12: Applying coverage-based rarefaction** 

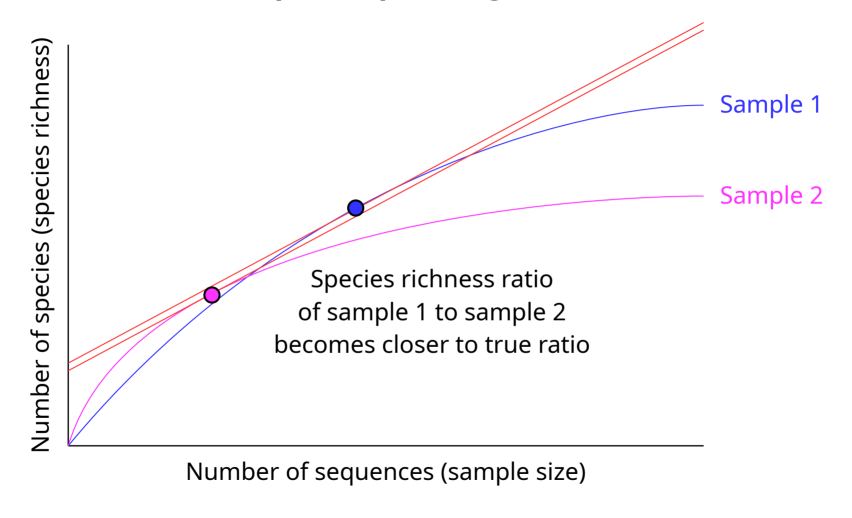


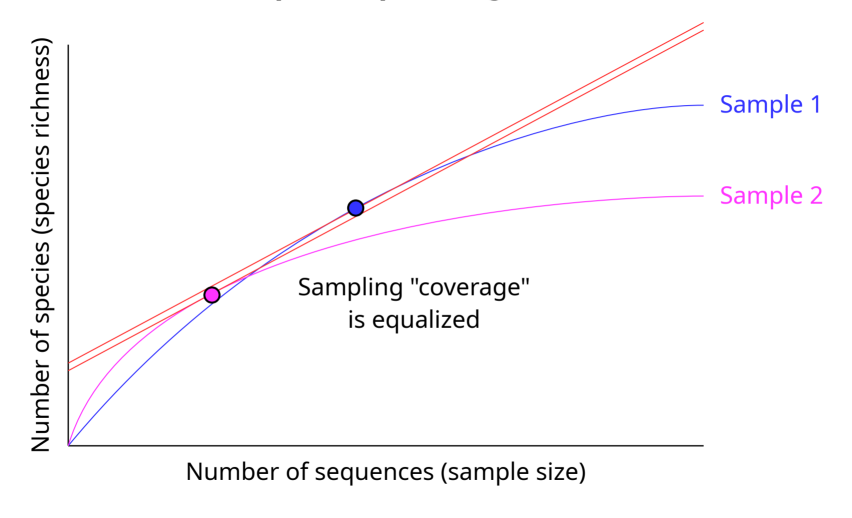


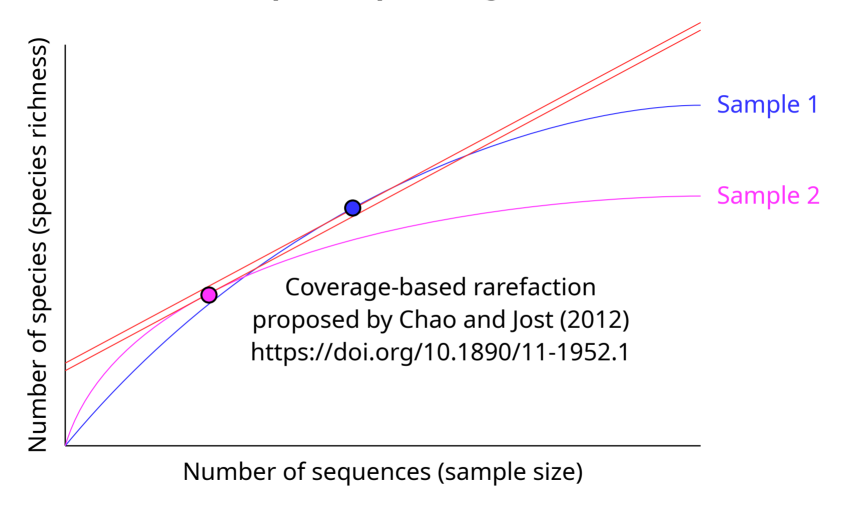












#### **Chapter 13: Calculating distance matrices**

- Inputs
- RarefiedCommunity (data.frame)
- BinaryRarefiedCommunity (d.f.)
  - in R workspace

- Outputs
- BrayCurtis (dist)
- Jaccard (dist)
- BinaryJaccard (dist)
- BinaryRaupCrick (dist)

in R workspace

Chapter 13: Calculating distance matrices

# Interlude: Community distance (β diversity) metrics, PERMANOVA and NMDS

#### See

- Anderson et al. (2010) https://doi.org/10.1111/j.1461-0248.2010.01552.x
- Anderson (2001) https://doi.org/10.1111/j.1442-9993.2001.01070.pp.x
- Anderson (2017) https://doi.org/10.1002/9781118445112.stat07841
- 土居・岡村 (2010) https://doi.org/10.18960/seitai.61.1\_3

#### **Chapter 14: Executing PERMANOVA**

- Inputs
- BrayCurtis (dist)
- Jaccard (dist)
- BinaryJaccard (dist)
- BinaryRaupCrick (dist)
- Metadata.tsv

in top directory

in R workspace

- Outputs
- BrayCurtisPERMANOVA.txt
- JaccardPERMANOVA.txt
- BinaryJaccardPERMANOVA.txt
- BinaryRaupCrickPERMANOVA.txt

in OverlappedPairedEnd\_11\_RAnalysisResults

**Chapter 14: Executing PERMANOVA** 

#### **Chapter 15: Executing NMDS**

- Inputs
- BrayCurtis (dist)
- Jaccard (dist)
- BinaryJaccard (dist)
- BinaryRaupCrick (dist)
- Metadata.tsv

- Outputs
- NMDS.pdf

in OverlappedPairedEnd\_11\_RAnalysisResults

in R workspace

in top directory

**Chapter 15: Executing NMDS** 

## Chapter 16: Executing Mantel correlogram analysis using geographical distance

- Inputs
- BrayCurtis (dist)
- Jaccard (dist)
- BinaryJaccard (dist)
- BinaryRaupCrick (dist)
- in R workspace

Metadata.tsv

in top directory

- Outputs
- GeoMCA.pdf

in OverlappedPairedEnd\_11\_RAnalysisResults

Chapter 16: Executing Mantel correlogram analysis using geographical distance

#### Chapter 17: Executing Mantel correlogram analysis using date interval

- **Inputs**
- BrayCurtis (dist)
- Jaccard (dist)
- BinaryJaccard (dist)
- BinaryRaupCrick (dist)
- Metadata.tsv

- **Outputs**
- DateMCA.pdf

in OverlappedPairedEnd\_11\_RAnalysisResults

in R workspace

in top directory

**Chapter 17: Executing Mantel correlogram analysis using date interval** 

#### Conclusion: Metabarcoding analysis using Claident and R

- Claident is integrated package for translation from high-throughput amplicon sequence data into ecological communities
- Claident can remove contaminants including index-hopped sequences using unused index combinations and blank samples (negative controls)
- Most studies lack decontamination and this might affect the conclusion of such studies
- Detection power of metabarcoding should re-evaluate using decontamination and our knowledge of that might need to be revised
- Claident provides multiple taxonomic assignment methods and can merge those results
- R can import tab-separated text made by Claident
- vegan is strongly recommended for community ecological analyses