

DTU Health Technology Bioinformatics

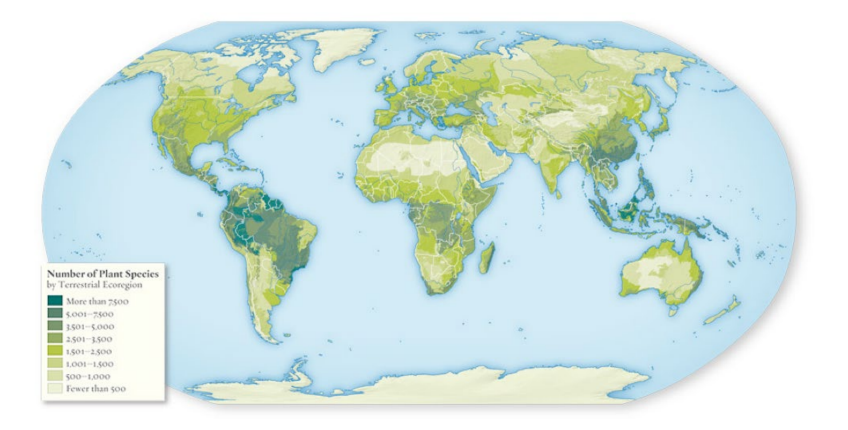
Quantitative metagenomics

Gisle Vestergaard Associate Professor Section of Bioinformatics Technical University of Denmark gisves@dtu.dk



Classical measures

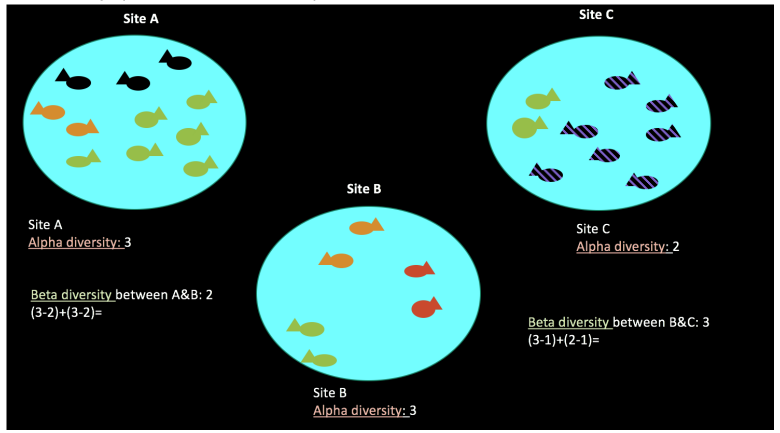
- Abundance
- Richness
- Rarefaction
- Diversity
 - Alpha
 - Beta





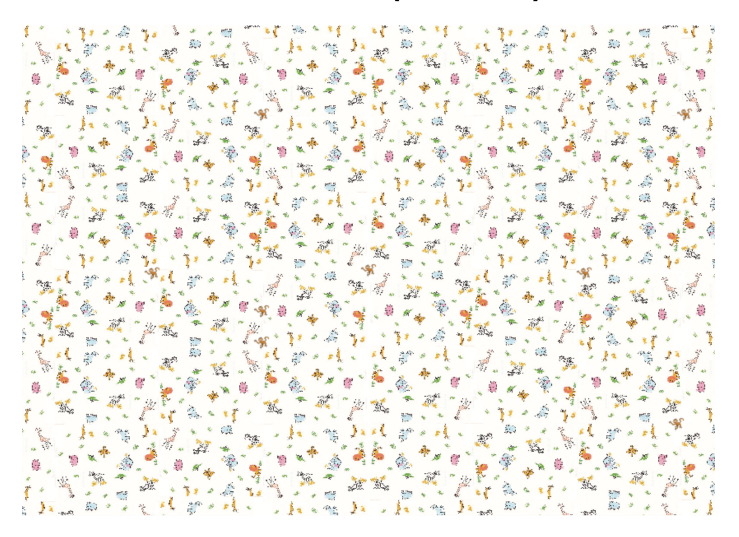
Describing the spatial component of biodiversity

- Alpha diversity (within sample)
- Beta diversity (between samples)





Abundance (counts)

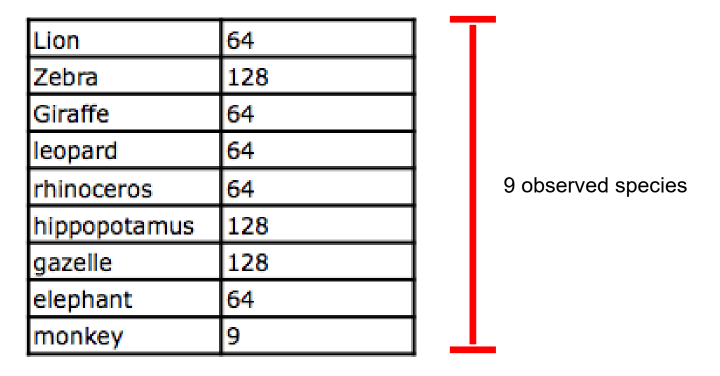


Lion	64
Zebra	128
Giraffe	64
leopard	64
rhinoceros	64
hippopotamus	128
gazelle	128
elephant	64
monkey	9



Species richness

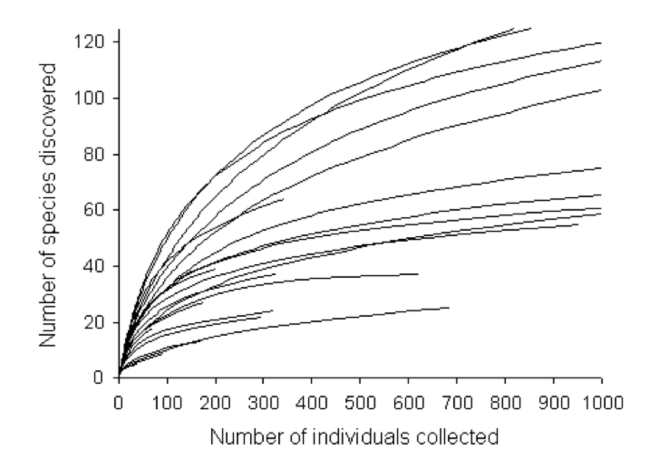
• The number of different species in a system





Rarefaction

- Species richness is a function of our no. observations
- When have we sampled enough?





Shannon index

- Incorporates species richness & eveness
- Quantify the entropy (information content)
- Quantifies the uncertainty (degree of surprise) associated with a prediction
- The Shannon index increases as both the richness and the evenness of the community increase
- Typical values are generally between 1.5 and 3.5 in most ecological studies, and the index is rarely greater than 4

$$H' = -\sum_{i=1}^{R} p_i \ln p_i \qquad H' = -(\ln p_1^{p_1} + \ln p_2^{p_2} + \ln p_3^{p_3} + \dots + \ln p_R^{p_R})$$

 P_i = species proportion

R = observed species



Alpha diversity



Lion	1
Zebra	2
Giraffe	1
Leopard	1
Rhinoceros	1
Hippopotamus	2
Gazelle	2
Elephant	1
Monkey	0

$$H' = -(\ln p_1^{p_1} + \ln p_2^{p_2} + \ln p_3^{p_3} + \dots + \ln p_R^{p_R})$$

11 animals (NOT species) meaning each animal is 0.09 of the total abundance

$$H' = -(In(0.09^{0.09}) + In(0.18^{0.18}) + ... = 2.0$$



Bray-curtis dissimilarity

$$0 \le B \le 1$$

$$B_{ij} = 1 - 2C_{ij} / (S_i + S_j)$$

C = sum of the lowest count of all common species

S = total count of the sample

1 means that they do not share anything

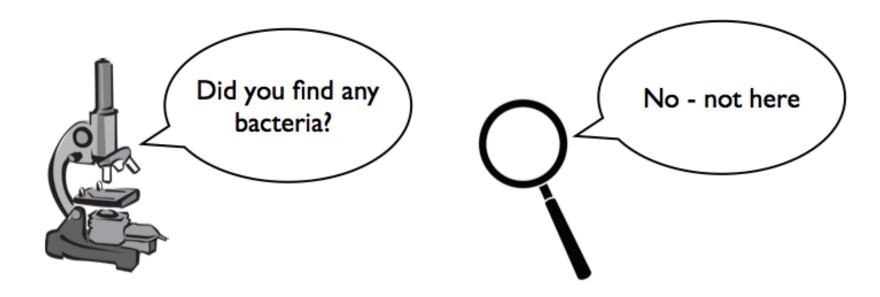
$$B_{s1s2} = 1 - 2*(2+1) / (9 + 13) = 0.73$$

Lion	0	2
Zebra	3	2
Giraffe	0	4
Leopard	0	2
Rhinoceros	1	2
Hippodrome	4	0
Gazelle	0	1
Elephant	1	0
Total	9	13

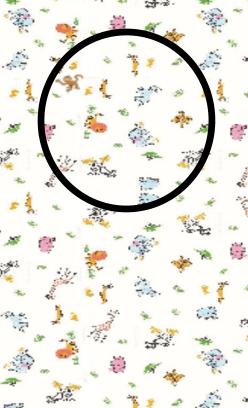


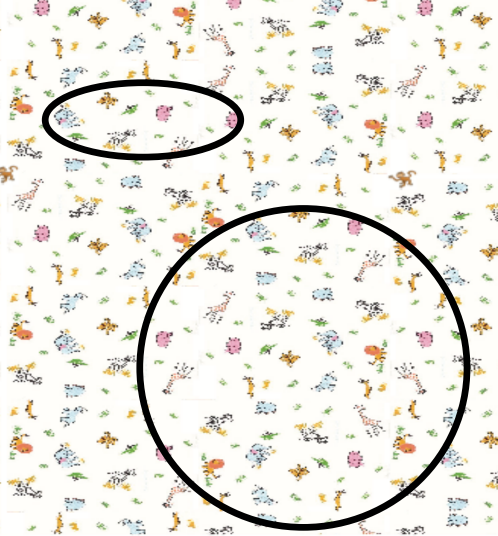
Sampling effect

• To be fair we should sample equally in the systems we investigate



Sample sizes







Sample sizes

- Accounting for different sample sizes:
 - –Normalise to sample size
 - -Rarefy (downsize) samples
 - -Statistically model the variance



Normalising

$$N = n_i/n_{tot}$$

Lion	64	1
Zebra	128	2
Giraffe	64	1
Leopard	64	1
Rhinoceros	64	1
Hippopotamus	128	2
Gazelle	128	2
Elephant	64	1
Monkey	9	0
Total	713	11

Lion	8.98	9.09
Zebra	17.95	18.18
Giraffe	8.98	9.09
Leopard	8.98	9.09
Rhinoceros	8.98	9.09
Hippopotamus	17.95	18.18
Gazelle	17.95	18.18
Elephant	8.98	9.09
Monkey	1.26	0
Total	100	100

Issue with different sampling power (higher chance of observing rare species)



Downsize / rarefy

- Resample an equal number of observations (reads) from each sample
- Select the target depth carefully
- The more reads we keep the more sensitive
- We may have to remove samples with few counts
- We might throw away a lot of data



Downsize / rarefy

Resample x amount of observations

Lion	64	1
Zebra	128	2
Giraffe	64	1
Leopard	64	1
Rhinoceros	64	1
Hippopotamus	128	2
Gazelle	128	2
Elephant	64	1
Monkey	9	0
Total	713	11

Lion	2	1
Zebra	3	2
Giraffe	0	1
Leopard	1	1
Rhinoceros	0	1
Hippopotamus	3	2
Gazelle	1	2
Elephant	0	0
Monkey	0	0
Total	10	10



Compositional analysis

- Arbitrary total
 - Sequencing depth never 100%
- Species can co-exist without abundance inter-influences
 - Independence between abundance is affected by the capacity of the sequencing instrument
 - Sequencing instrument has fixed number of slots

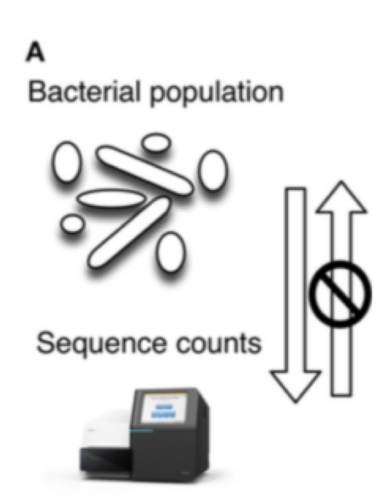
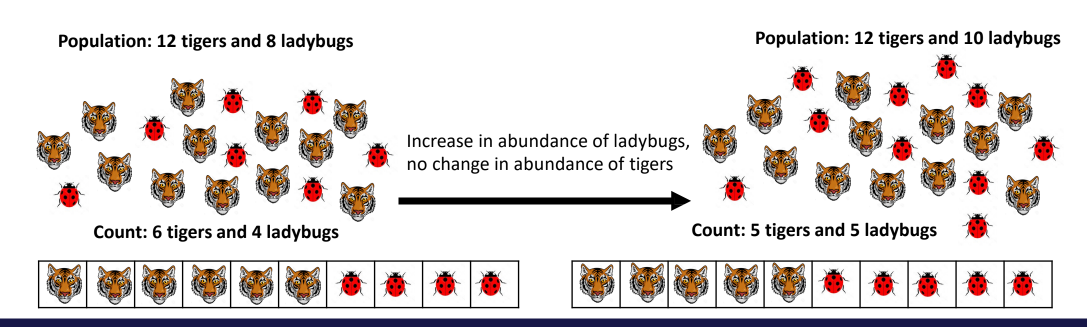


Figure from Cloor Crogory R. et al. Microbiome Datacets Are Compositional, And This Is Not Ontional, Frontiers in Microbiology 9, (2017)



Compositional analysis

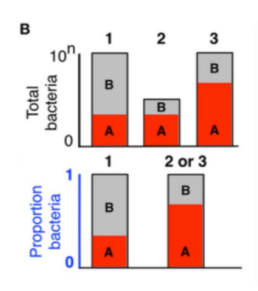
- **Example**: an environment containing both tigers and ladybugs
 - The abundances of the two are not affected by each other
 - If the abundance of the ladybugs increases some of the slots with tigers must instead be filled by ladybugs
 - i.e. the two environmentally independent species are affecting the read count of each other

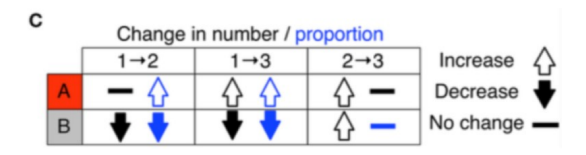




Compositional analysis

 Data can be differently interpreted when looking at absolute abundances vs. the relative abundances after sequences







DESeq2

Love et al. Genome Biology (2014) 15:550 DOI 10.1186/s13059-014-0550-8



METHOD Open Access

Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2

Michael I Love^{1,2,3}, Wolfgang Huber² and Simon Anders^{2*}

- Statistically model the variance & heteroscedasticity
- Use packages developed for RNA-seq such as DESeq2 and edgeR (negative binomial)
- Bayesian approach
 - PhILR

Lets try it

Take a sample and count the animals!

Determine Richness, Species abundance and calculate Shannon diversity index

Calculate Bray-Curtis dissimilarity with your neighbour