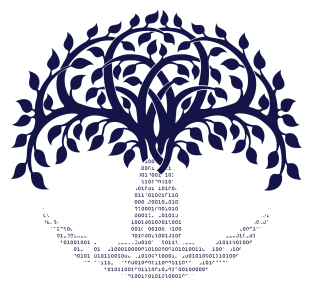
# 



# DTU Health Technology Bioinformatics

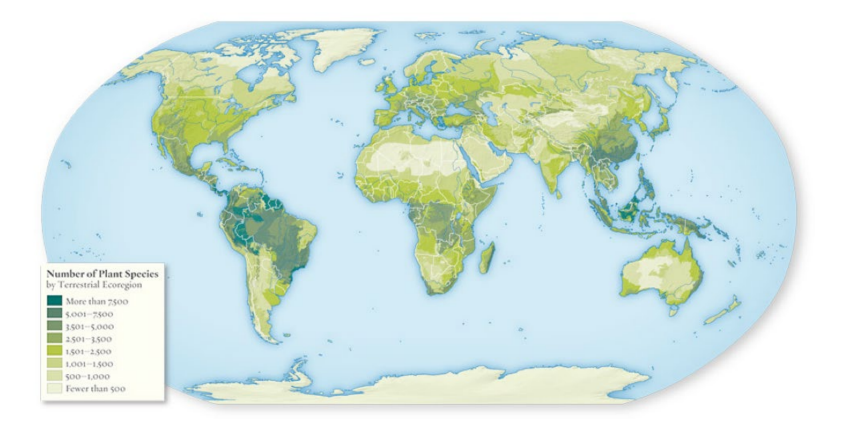
### Quantitative metagenomics

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# **Classical measures**

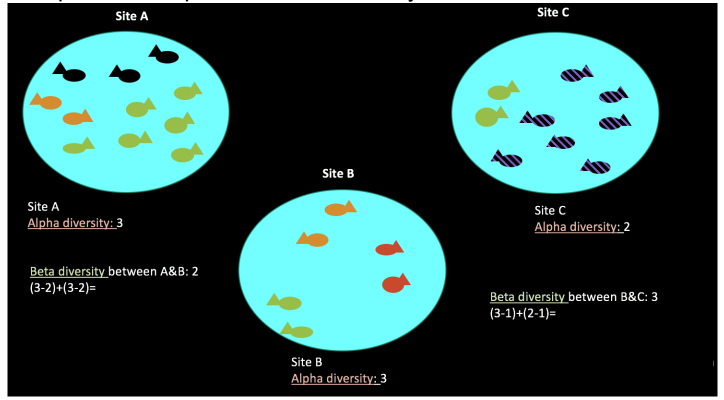
- Abundance
- Richness
- Rarefaction
- Diversity
  - Alpha
  - Beta





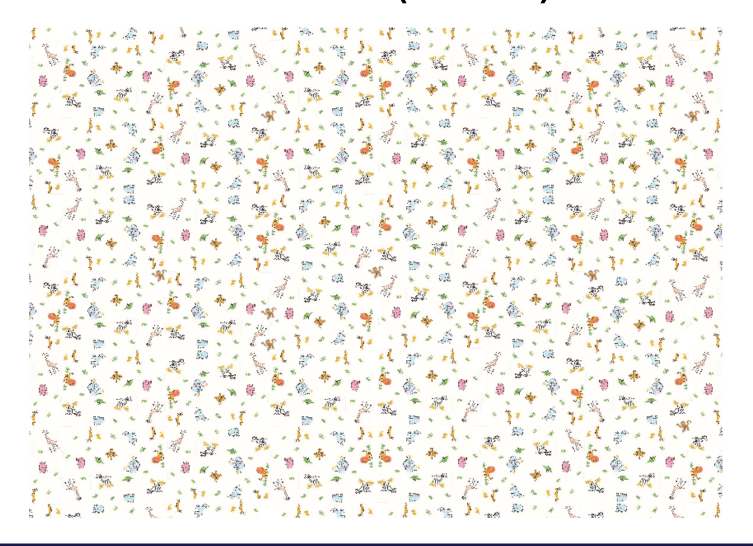
# Describing the spatial component of biodiversity

- Alpha diversity (within sample)
- Beta diversity (between samples)
- We can compare both alpha and beta diversity





# **Abundance (counts)**

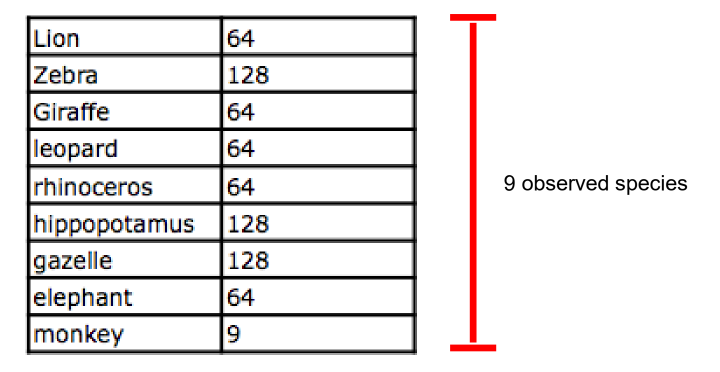


Lion	64
Zebra	128
Giraffe	64
leopard	64
rhinoceros	64
hippopotamus	128
gazelle	128
elephant	64
monkey	9



# **Species richness**

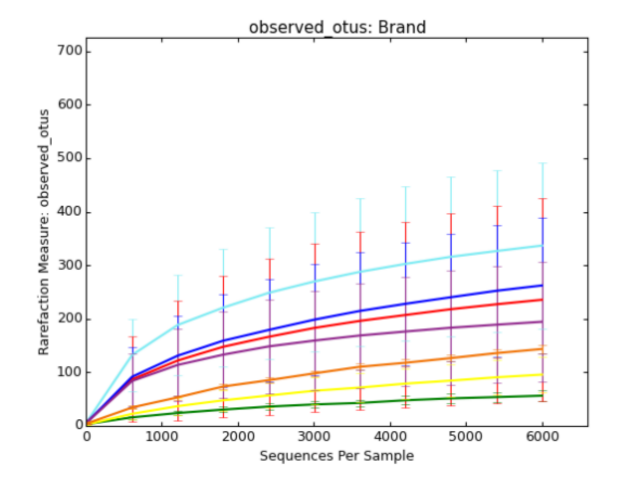
• The number of different species in a system





#### Rarefaction

- Species richness is a function of our no. observations
- When have we sampled enough?
- Used for 16s rRNA amplicons and not shotgun metagenomics...why?





#### **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

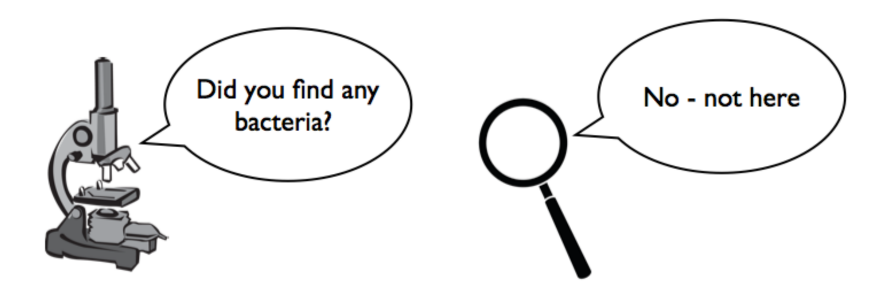
$$B_{s1s2} = 1 - 2*(2+1) / (9 + 13) = 0.73$$

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

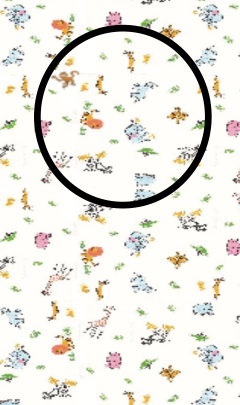


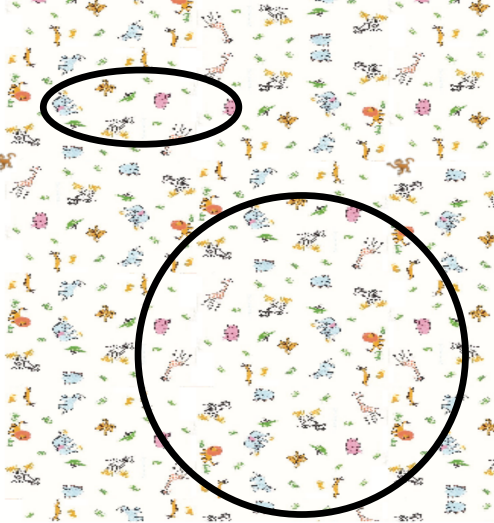
# Sampling effect

• 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%
- Absolute abundances vs. the relative abundances
  - Species can co-exist without abundance inter-influences
  - Independence between abundance is affected by the capacity of the sequencing instrument

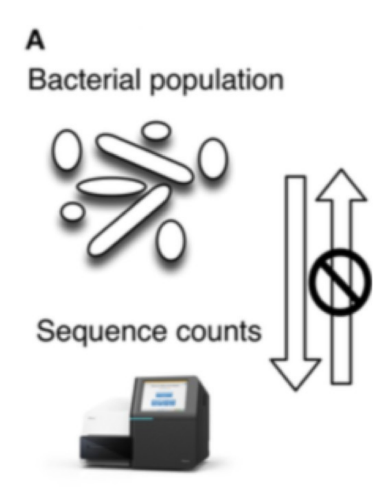
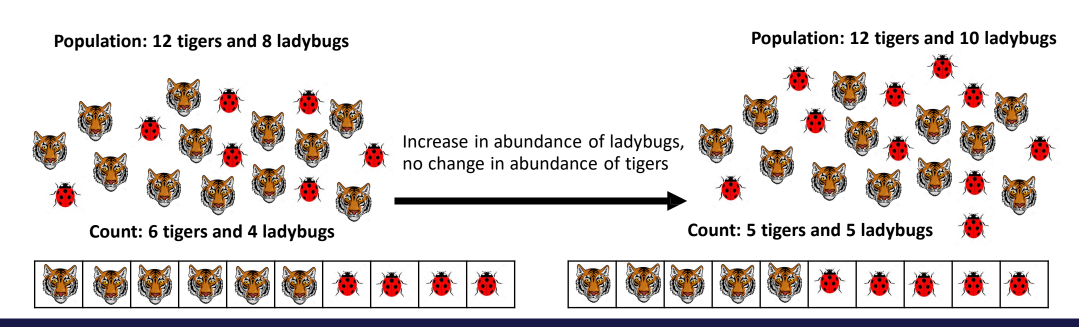


Figure from: Gloor, Gregory B. et al., Microbiome Datasets Are Compositional: And This Is Not Optional. Frontiers in Microbiology 8 (2017)



# **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





# 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<sup>1,2,3</sup>, Wolfgang Huber<sup>2</sup> and Simon Anders<sup>2\*</sup>

- Statistically model the variance & heteroscedasticity
- Use packages developed for RNA-seq such as DESeq2 and edgeR (negative binomial)
- Several alternatives which we will also try!

# 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