Clustering

AB332 - Autumn 2021

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Illumina seq. and amplicon data

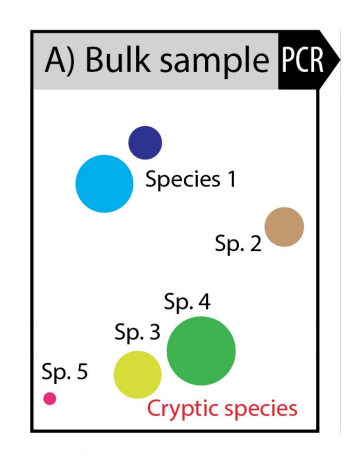
- Illumina HiSeq and MiSeq sequencing has been used extensively for amplicons sequencing in the last years
 - High yields with relative low cost
 - HiSeq (and NovaSeq) short fragments 2*150bp
 - MiSeq a bit longer reads 2*300bp
- Low error rate, but due to the high number of sequences generated errors are bound to occur in the library
 - (About 1% pr 1000 nt, not adjusted for improvements due to overlapping reads.)
- How to deal with errors, has been an ongoing discussion for high-throughput data for a quite some time



- Reasons for clustering of metabarcoding data:
 - Reduce effect of sequencing error
 - Reduce "noise"

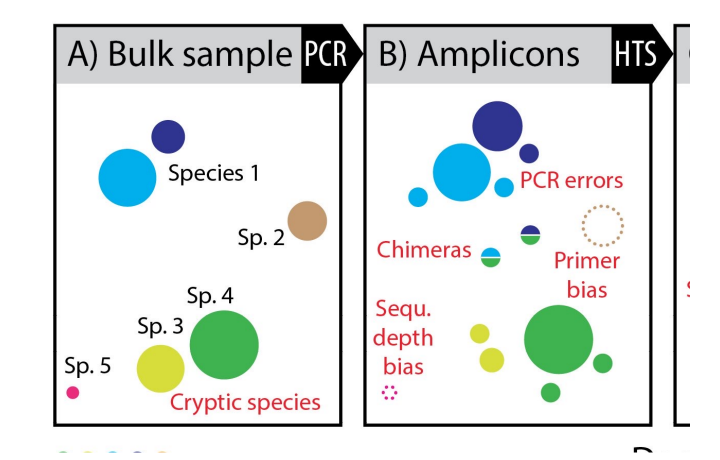


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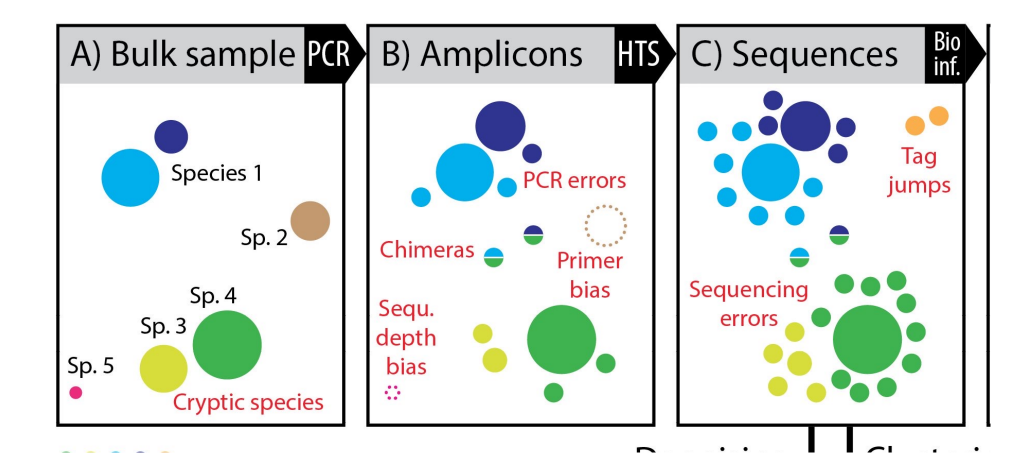


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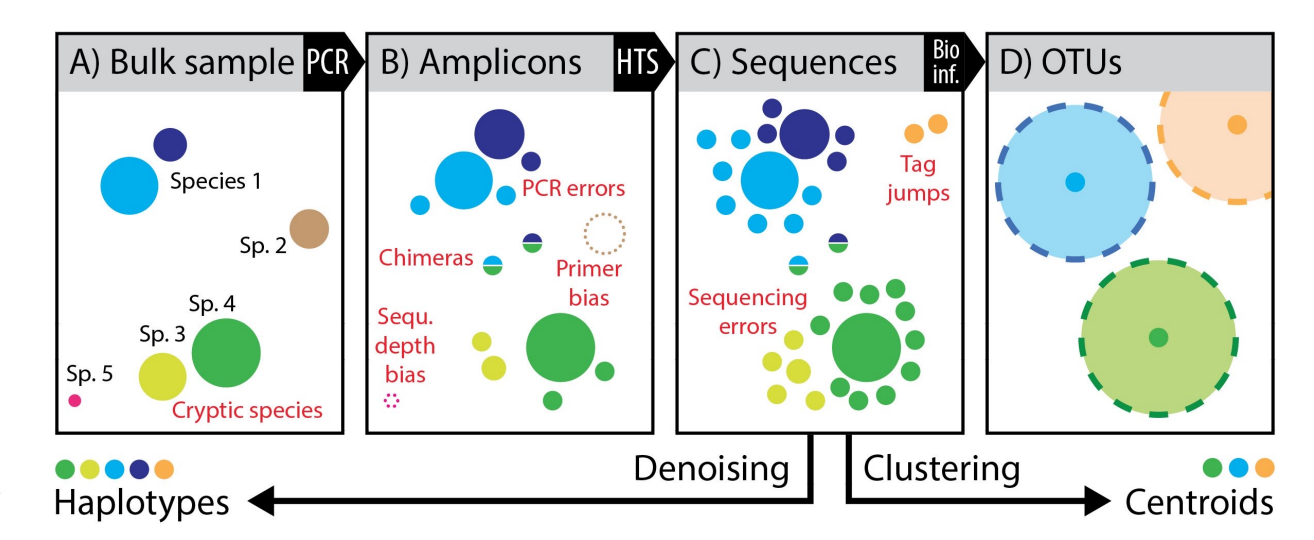


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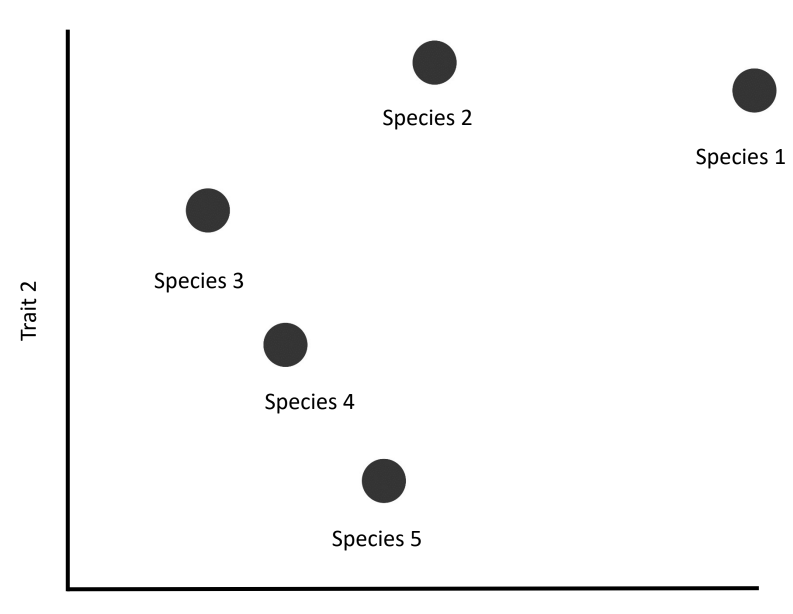


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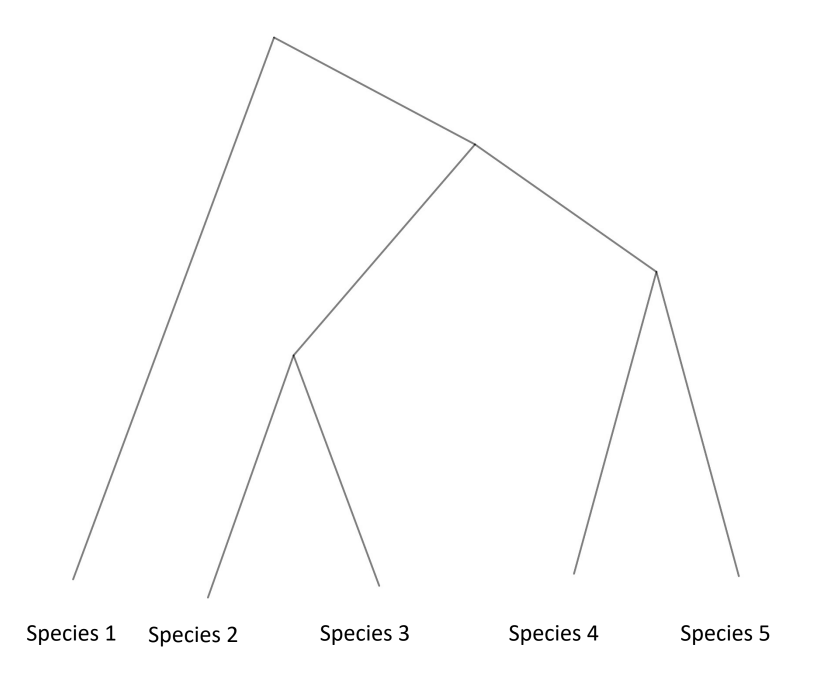


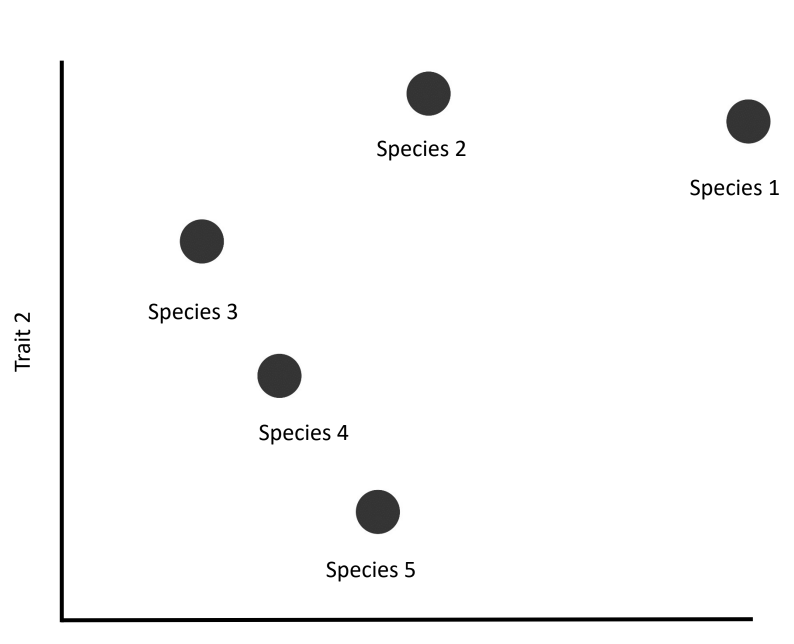
The ideal picture



Trait 1

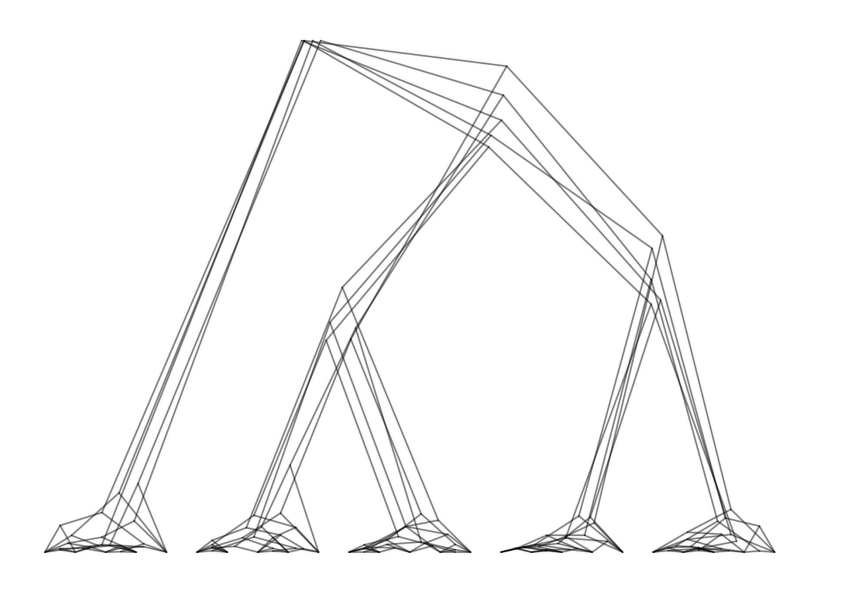
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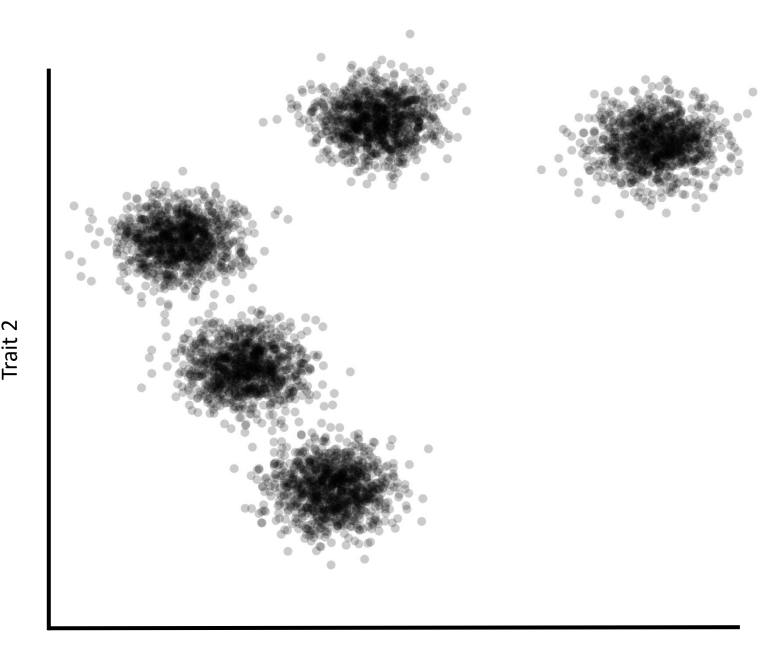




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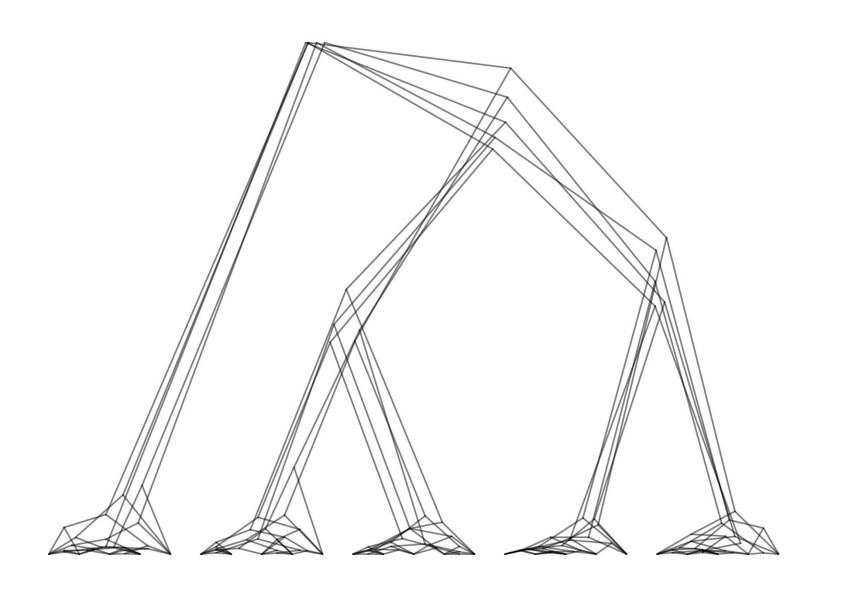
The reality

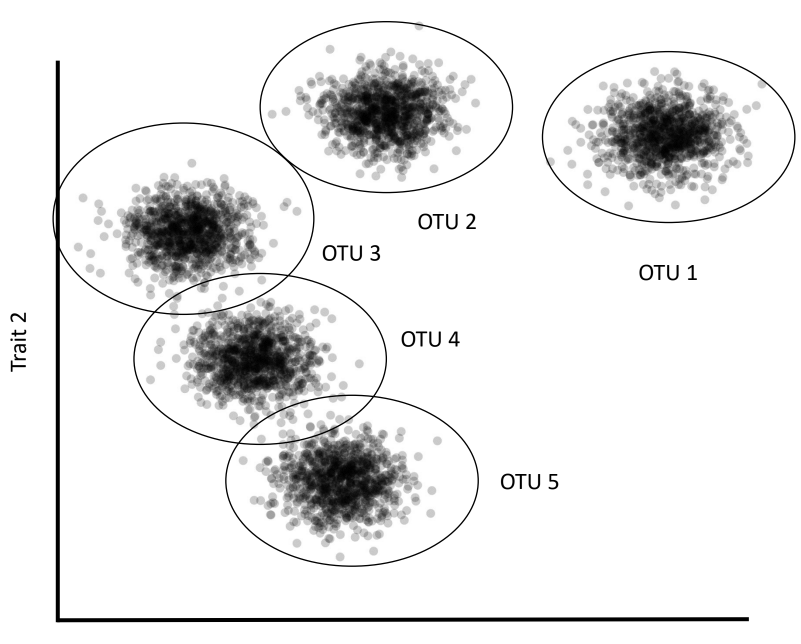




Trait 1

The reality





Trait 1

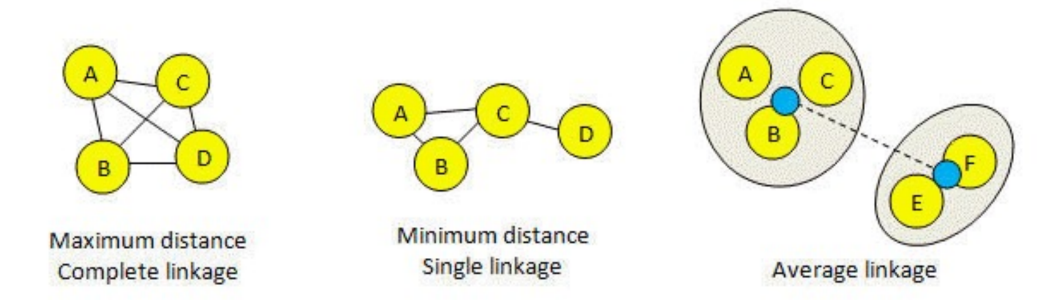
Amplicons, OTUs, and species

- Clusters of similar amplicons is called OTUs (operational taxonomic units)
 - Proxies for species, ecotypes, populations or another functional unit
 - Clustered based on often based on dissimilarity cut-off
 - But what cut-off to choose?
 - And how to calculate the similarity/dissimilarity?
- One solution is UPGMA-style clustering.
 - I.e. clustering at a fixed sequence distance
 - For instance 97%, which is often used for 18S V4



Clustering types

- Complete linkage means that all pairs of sequences in a cluster must be closer than the threshold.
- **Single linkage** means that a sequence should be included in a cluster if the distance to any other sequence is below the threshold.
- Average linkage (similar to UPGMA) distance between the "average sequence" for a cluster with the other cluster (the average is in fact calculated over all pairs).
- With average linkage, clusters tend to be larger than maximum linkage and smaller than minimum linkage.





UPARSE - (U/VSEARCH)

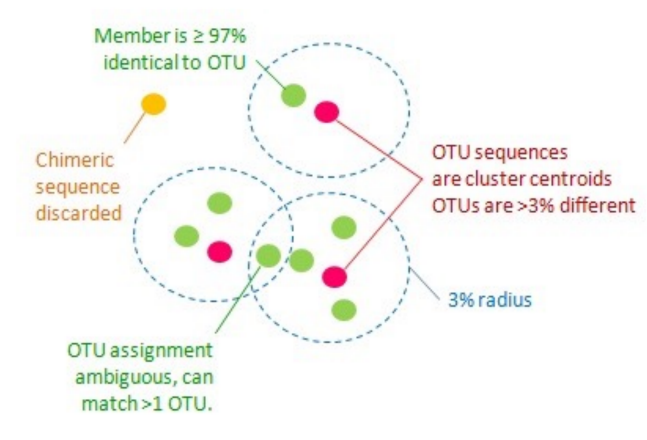
- UPARSE-OTU takes sequences in order of decreasing abundance as input. This means that OTU centroids tend to be selected from the more abundant reads, and hence are more likely to be correct biological sequences.
- 1. All pairs of OTU sequences should have pair-wise sequence identity for a set value (typically 97%)
- 2. An OTU sequence should be the most abundant within a 97% neighbourhood.
- 3. Chimeric sequences should be discarded.
- 4. All non-chimeric input sequences should match at least one OTU with >= 97% identity.

Published: 18 August 2013

UPARSE: highly accurate OTU sequences from microbial amplicon reads

Robert C Edgar

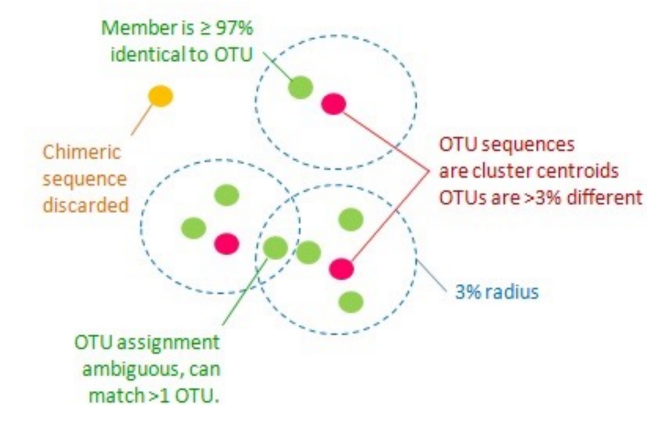
Nature Methods 10, 996–998(2013) | Cite this article
6924 Accesses | 5903 Citations | 113 Altmetric | Metrics





UPARSE

- UPARSE implements a greedy algorithm that performs
 OTU clustering
- Advantages
 - Fast and greedy
 - Similarity threshold is relatively easy to interpret (although not necessarily easy to decide...)
- Disadvantages
 - Arbitrary fixed global clustering threshold. But different lineages evolve at different rate, which means that there is no single cut-off value for the entire tree of life
 - The input order of the amplicons strongly influences the clustering results. Centroid selections are not re-evaluated during the clustering process, this might lead to inaccurately formed OTUs.







Swarm: robust and fast clustering method for amplicon-based studies

Frédéric Mahé^{1,2,3}, Torbjørn Rognes^{4,5}, Christopher Quince⁶, Colomban de Vargas^{1,2} and Micah Dunthorn³

- Fast and exact, two-phased, agglomerative, unsupervised (de novo) single-linkage-clustering algorithm.
- Advantage
 - No global (and arbitrary) clustering threshold
 - The result is not dependent on the input sequence order
- SWARM builds OTUs in two steps
 - An initial set of OTUs is constructed by iteratively agglomerating similar amplicons
 - Amplicon aundances are used to break clusters into sub-OTUs



Greedy cluster vs. SWARM

PeerJ

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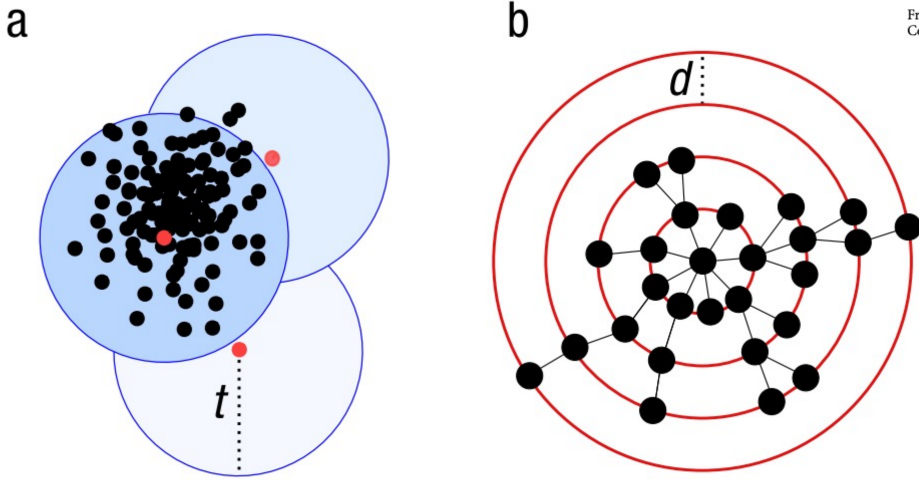
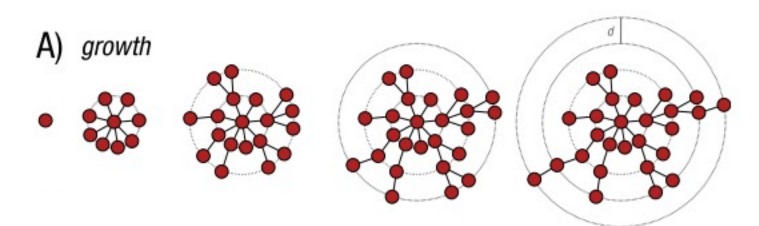


Figure 1 Schematic view of the greedy clustering approach and comparison with swarm. (A) Visualization of the widely used greedy clustering approach based on centroid selection and a global clustering threshold, *t*, where closely related amplicons can be placed into different OTUs. (B) By contrast, Swarm clusters iteratively by using a small user-chosen local clustering threshold, *d*, allowing OTUs to reach their natural limits.

• (A) Swarm clusters amplicons iteratively by using a small user-chosen local threshold, d, allowing OTUs to grow to their natural limits, where no other amplicons can be added.





Swarm v2: highly-scalable and high-resolution amplicon clustering

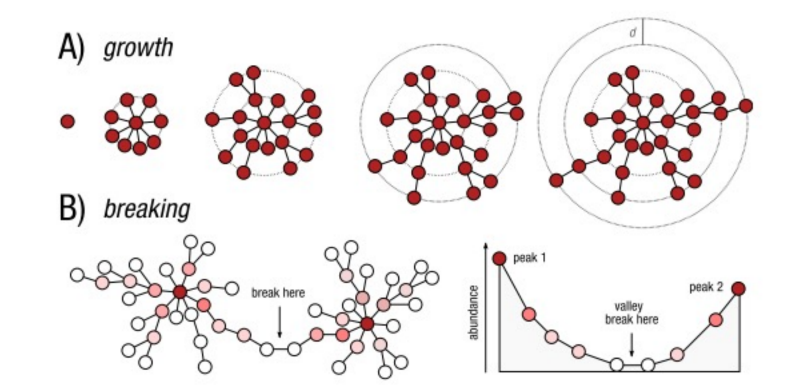
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- ⁶ UMR7144 Station Biologique de Roscoff, Sorbonne Universités, UPMC Univ Paris 06, Roscoff,



- (A) Swarm clusters amplicons iteratively by using a small user-chosen local threshold, d, allowing OTUs to grow to their natural limits, where no other amplicons can be added.
- (B) Swarm takes into account the abundance of each amplicon to produce higher resolution clusters, by not allowing the formation of amplicon chains. The darker the red, the higher the abundance.





Swarm v2: highly-scalable and high-resolution amplicon clustering

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6 UMR7144 Station Biologique de Roscoff, Sorbonne Universités, UPMC Univ Paris 06, Roscoff



- (A) Swarm clusters amplicons iteratively by using a small user-chosen local threshold, d, allowing OTUs to grow to their natural limits, where no other amplicons can be added.
- (B) Swarm takes into account the abundance of each amplicon to produce higher resolution clusters, by not allowing the formation of amplicon chains. The darker the red, the higher the abundance.
- (C) The fastidious option avoids under-grouping (e.g., the production of small OTUs such as singletons and doubletons) by postulating the existence of virtual linking amplicons to graft smaller OTUs onto larger ones.



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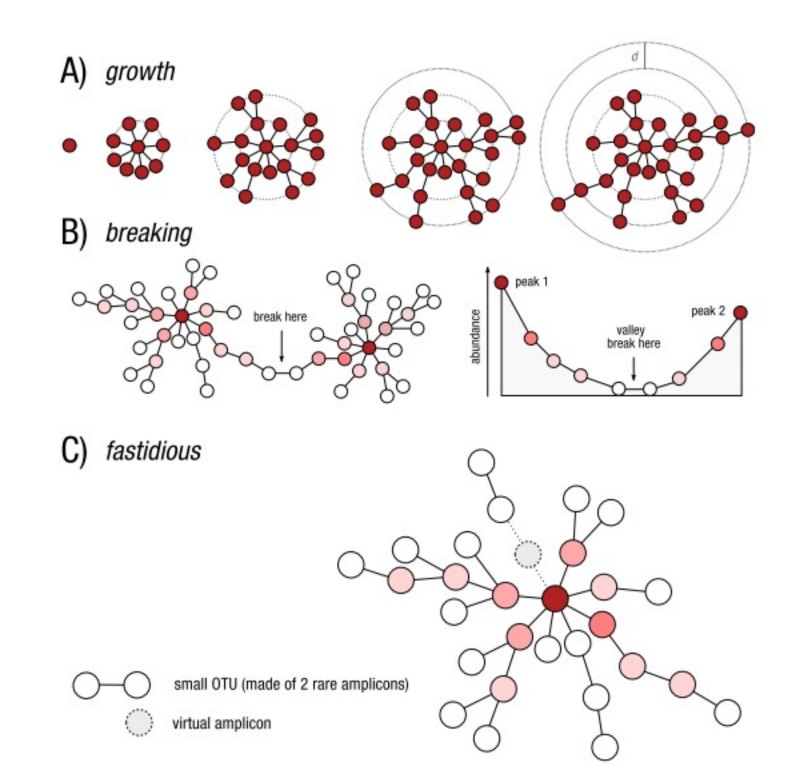
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Erance





- An update of DADA (Rosen et al., 2012)
- Divisive Amplicon Denoising Algorithm
 - Originally made for 454 data.
- **DADA2** Infers "amplicon sequence variants from Illumina-scale amplicon data data without imposing the arbitrary dissimilarity thresholds that define molecular OTUs"

DADA2: High-resolution sample inference from Illumina amplicon data

Benjamin J Callahan¹, Paul J McMurdie², Michael J Rosen³, Andrew W Han², Amy Jo A Johnson² & Susan P Holmes¹



- Advantages (according to themselves, aka the selling point...)
 - **Resolution**: DADA2 infers exact amplicon sequence variants (ASVs) from amplicon data, resolving biological differences of even 1 or 2 nucleotides.
 - Accuracy: DADA2 reports fewer false positive sequence variants than other methods report false OTUs.
 - Comparability: The ASVs output by DADA2 can be directly compared between studies, without the need to reprocess the pooled data.
 - **Computational Scaling**: The compute time of DADA2 scales linearly sample number, and memory requirements are essentially flat.



- The error model incorporates <u>quality information</u>, which is usually ignored by other methods (after filtering).
- The error model incorporates <u>quantitative abundances</u>, whereas most other methods use abundance ranks (if they use abundance at all).
- The error model have different error rates for different substitutions,
 - eg. A->C, is different from A->G whereas other methods merely count the mismatches.
- DADA2 can <u>parameterize</u> its error model from the data itself, rather than relying on previous datasets that may or may not reflect the PCR and sequencing protocols used in your study.



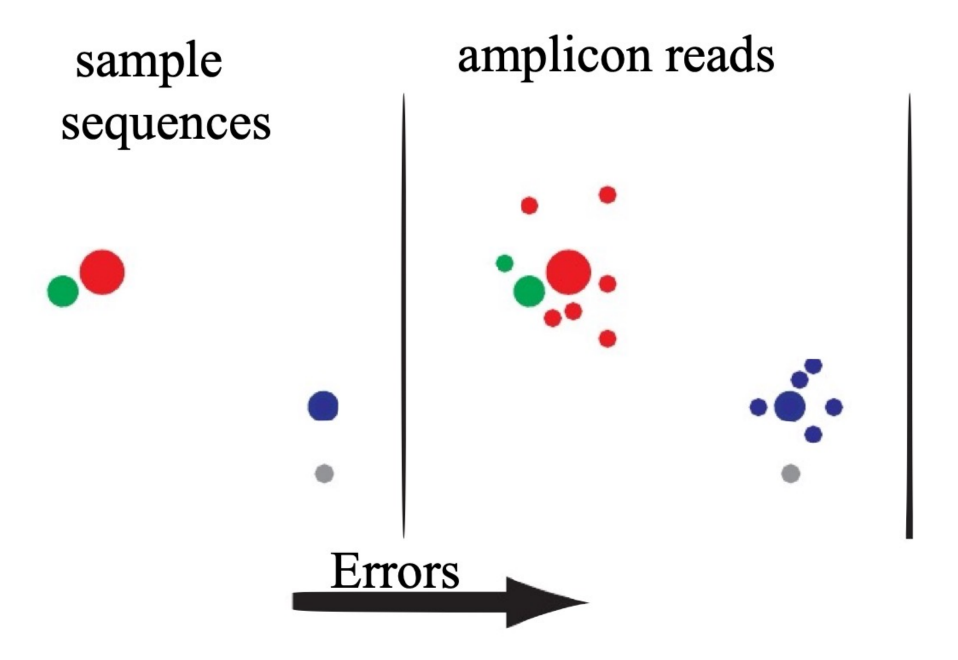
sample sequences



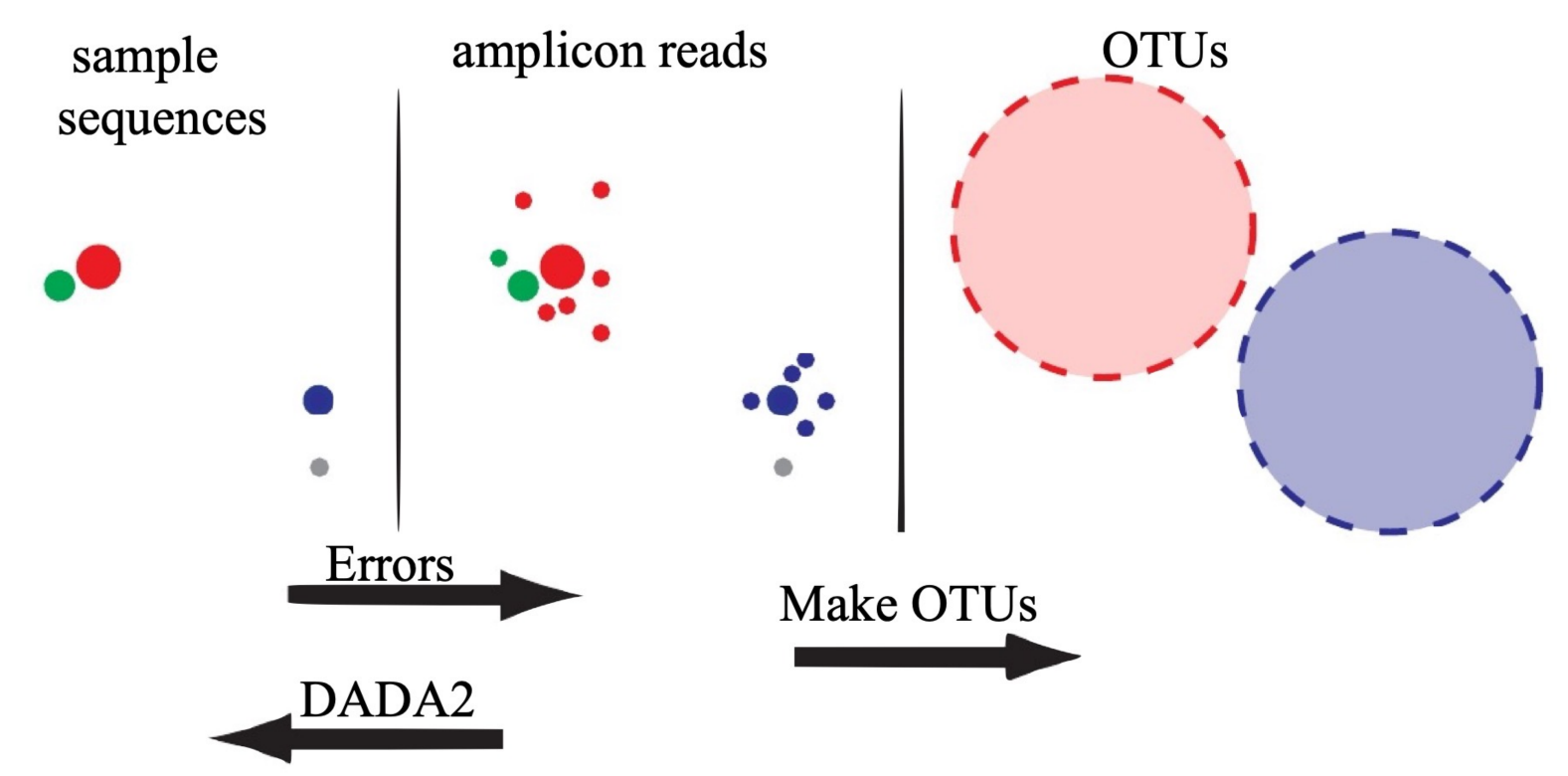












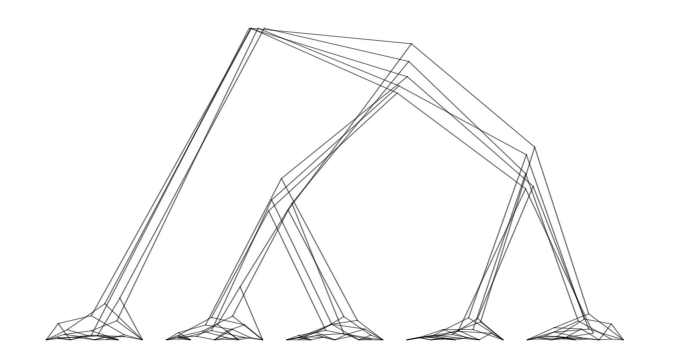


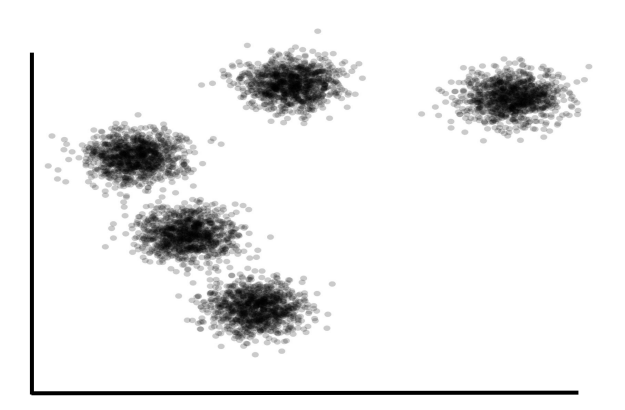
- The core denoising algorithm in the DADA2 is built on a model of the errors in Illumina-sequenced amplicon reads.
- **However:** Clustering might still be needed since there are other sources of noise.
- And depends on the marker sequenced
 - SSU (18S) has multiple copies in many eukaryotes
 - Intragenomic variation might be around the "usuall" cut-off used for vsearch/usearch type clustering (i.e. 2-3% difference).
 - ITS is highly variable



Do you still need to cluster?

- Clustering might still be needed since there are other sources of "noise".
 - I.e. even though an inferred read is a "true sequence" with a discernible abundance it might not by itself represent a species.







The output

- For our purpose, the important part is that the output is a table with the *read* abundance for *OTUs* across the samples.
- Keep in mind that these are Operational Taxonomic Units not necessarily the same as species
- Read abundance does not equal organismal abundance

	Sample_1	Sample_2	Sample_3	Sample_4	Sample_5	Sample_6	etc
OTU_1	1	563	87	459	43	0	
OTU_2	213	7	5	3	68	8	
OTU_3	15	6	88	95	9	6	
OTU_4	762	5	43	1	0	0	
etc							



Long read amplicon sequencing

- Recent development in long read sequencing technologies:
 - PacBio HiFi reads are 99.9% accurate and up to 15 kbp long!
 - Still not as many reads as Illumina sequencing, but there is no need for denoise.
 - The much longer reads have a higher phylogenetic signal

MOLECULAR ECOLOGY RESOURCES

Long-read metabarcoding of the eukaryotic rDNA opero to phylogenetically and taxonomically resolve environmental diversity

Mahwash Jamy, Rachel Foster, Pierre Barbera, Lucas Czech, Alexey Kozlov, Alexandros Stamatakis, Gary Bending, Sally Hilton, David Bass ⋈, Fabien Burki ⋈,



- Resources:
- The developer has a nice tutorial
 - https://benjjneb.github.io/dada2/
 - https://benjjneb.github.io/dada2/tutorial.html
- DADA2 for ITS
 - https://benjjneb.github.io/dada2/ITS_workflow.html