Taxonomic Assignment and DNA Metabarcoding

Dr. Marie Louise Davey



Why assign taxonomy at all?

Not strictly necessary to answer alpha and betadiversity questions

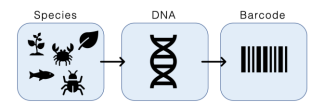
 Detecting shifts in community composition and genetic diversity doesn't require taxonomic assignments

Assigning taxonomy links sequences to a wealth of pre-existing information

 Linking sequences to species improves interpretation and explanation of patterns in alpha and beta diversity



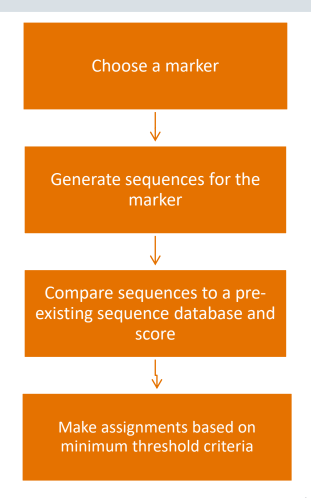
Taxonomic Assignment









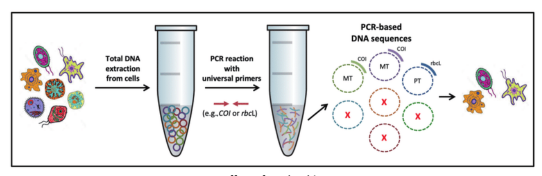




Markers

Marker choice impacts taxonomic assignment

- No marker is perfect
 - Markers have taxonomic bias
 - Discriminating power varies between markers and taxonomic groups
 - Database quality, availability, and completeness varies between markers



Effect of marker bias

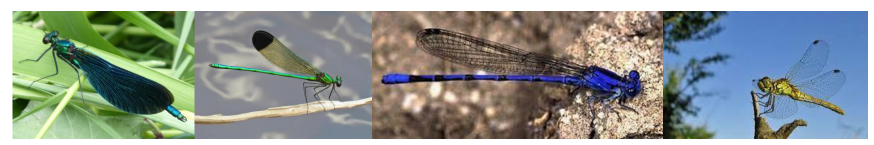


Marker Choice



Coverage and quality of DNA barcode references for Central and Northern European Odonata

Matthias Geiger¹, Stephan Koblmüller², Giacomo Assandri³, Andreas Chovanec⁴, Torbjørn Ekrem⁶, Iris Fischer^{5,7,8}, Andrea Galimberti⁹, Michał Grabowski¹⁰, Elisabeth Haring^{5,7,8}, Axel Hausmann¹¹, Lars Hendrich¹¹, Stefan Koch¹², Tomasz Mamos¹⁰, Udo Rothe¹³, Björn Rulik¹, Tomasz Rewicz¹⁰, Marcia Sittenthaler⁷, Elisabeth Stur⁶, Grzegorz Tończyk¹⁰, Lukas Zangl^{2,14,15} and Jerome Moriniere¹⁶



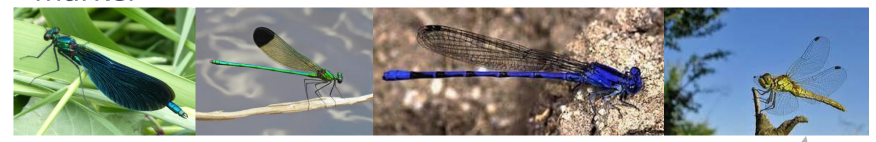


Marker Choice



Coverage and quality of DNA barcode references for Central and Northern European Odonata

- COI marker was selected for barcoding
- >80% of European species have a pre-existing publicly available barcode
- 88% of species tested could be resolved using the marker



Databases

Taxonomic assignment quality is highly dependent on database accuracy and completeness

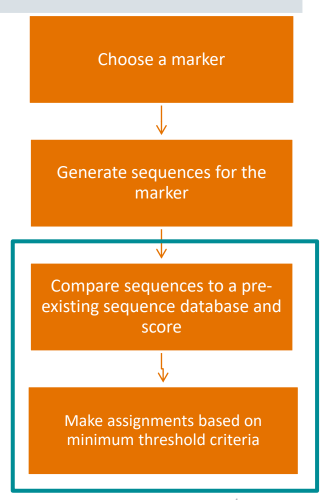
- Misidentified sequences create identification errors and low-quality assignments
- Missing reference sequences reduce resolution of taxonomic assignments, or results in misidentifications

Quality of taxonomic assignments can often be improved by creating custom-curated reference databases



Methods for Taxonomic Assignment

- Diverse algorithms are used for comparing sequences to databases and scoring the results
 - Alignment based BLAST, vsearch, OBItools
 - Phylogenetic based HmmUFOtu, TIPP, DECARD, SAP
 - Kmer-based machine learning approaches RDP, UTAX, SINTAX





Alignment based taxonomic assignment

Alignment strategy can be local (BLAST) or global (vsearch)

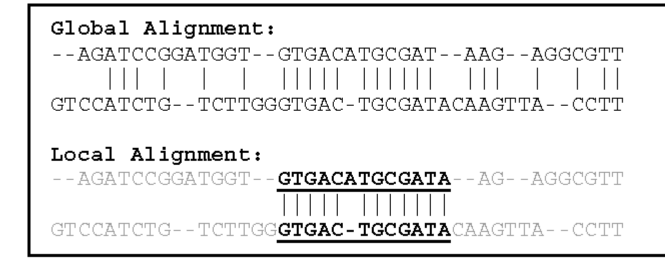
- Local alignments begin by checking a small piece of the query sequence against the reference database and then expanding the match to find areas of high similarity
- Global alignments find the best match in the reference database across the entire length of the query sequence

Output is typically an alignment score, percent identity, and coverage score

These are used as criteria for assigning taxonomy



Alignment based taxonomic assignment



62% similarity

93% similarity 32% coverage



Alignment based taxonomic assignment

Pros	Cons
 Extremely well developed infrastructure Easily applied to custom databases Computationally inexpensive, particularly for large databases High assignment rate 	 Typically lower taxonomic accuracy, lower precision, and lower taxonomic sensitivity than other methods Thresholds for successful assignment must be set by the user and require a priori knowledge No systematic assignment at higher taxonomic levels (unless paired with LCA)



Criteria for taxonomic assignments

Threshold Cutoffs

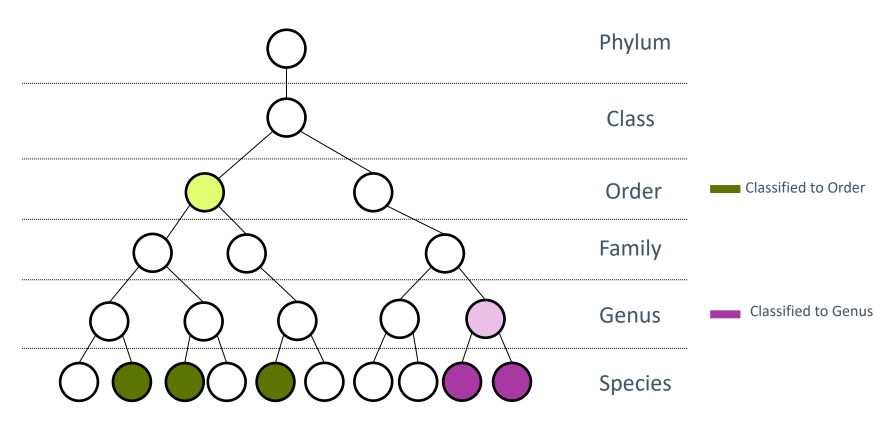
- A minimum 'score' must be met for a successful assignment
- 1. Identity based
- •reference sequence taxonomy is assigned for the best match exceeding the threshold value (often used in BLAST)
- 2. Confidence intervals
- •Proportion of kmers matching a given taxonomic assignment at a given taxonomic rank (ex/ SINTAX, RDP)

Lowest Common Ancestor (LCA) consensus

- Assigns consensus taxonomy based on a group of pre-defined 'best hits' for each sequence
- •Typically requires a separate additional step to calculate the LCA (ex/ LCA*, BASTA, MEGAN)



Lowest Common Ancestor Calculations









Phylogenetic based taxonomic assignment

Assignment of reads on a known reference tree, often followed by clustering or LCA analysis

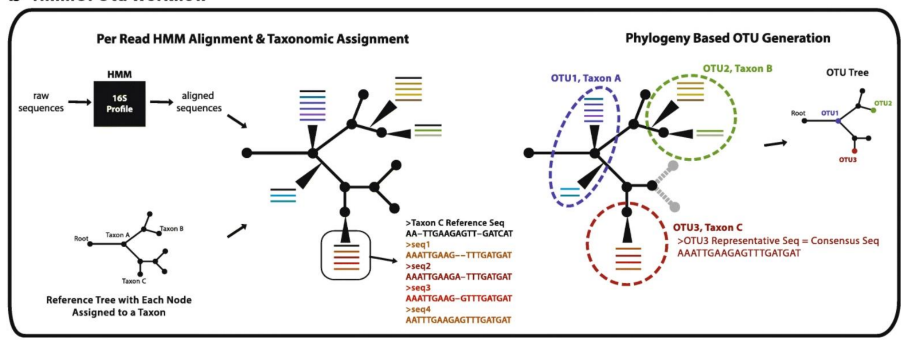
- 1.Placement on the reference tree is typically done using the pplacer algorithm, which is based on HMMER alignments of the query sequences with the reference database. This is often followed by clustering or LCA analysis
- 2.Alignment based searches (BLAST) are used to identify relevant reference sequences, and MCMC or bootstrapping are used to evaluate a series of phylogenetic trees generated from the query + references

Output is typically a taxonomy string with accompanying support statistics



Phylogenetic based taxonomic assignment

b HmmUFOtu Workflow





Phylogenetic based taxonomic assignment

Pros	Cons
 Assignment thresholds are flexible, and can vary between taxonomic groups Provides phylogenetic data about sequences Good taxonomic precision and sensitivity 	 Requires a known reference phylogeny Many algorithms require that the target gene region is alignable across the entire target group Computationally expensive Taxonomic accuracy can be compromised by an incomplete or poor reference phylogeny



Machine-learning based taxonomic assignment

Sequences are given a best classification based on a training set

- 1.Reads are broken into k-mers (sequence fragments of fixed length)
- 2.K-mers are compared to a training set of reference sequences with known taxonomies
- 3. Proportion of k-mers assigned to a given taxonomic level are used as a confidence score

Output is typically a taxonomy string with accompanying support statistics for each taxonomic level



Machine-learning based taxonomic assignment

Pros	Cons
 Provides confidence intervals for all taxonomic levels Assignment thresholds are flexible, and can vary between taxonomic groups High taxonomic precision and sensitivity 	 Can be computationally intensive Often requires establishment and benchmarking of a custom training set for the classification algorithm Database insufficiencies can create inaccuracy



How to decide?

- Find a method that balances your community's diversity with computational time
- Look for existing database resources, particularly for machine learning approaches
- Evaluate the importance of taxonomy to your conclusions

