# Species Tree Inference using SVDquartets

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January 31, 2015

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## **SVD**quartets

• In this tutorial, we'll discuss several different data types:

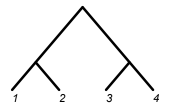
- Multi-locus data aligned DNA sequence data for many genes
- SNP data large number of SNPs sampled throughout the genome
- Single-locus data aligned DNA sequence data for a single gene
- In the first two cases, we'll assume that incongruence between gene trees and the species trees arises solely from the coalescent process
- In the third case, we assume that the locus under consideration is a single non-recombining unit

**Goal:** Estimate the underlying phylogenetic tree (species tree or gene tree)

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#### Definition: splits

 Definition: A split of a set of taxa L is a bipartition of L into two non-overlapping subsets A and B, denoted A|B. A split A|B is valid for tree T if the subtrees containing the taxa in A and in B do not intersect.

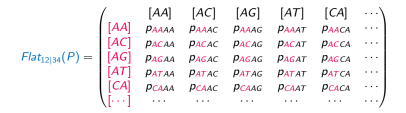


- Valid: 12|34
- Not valid: 13|24 14|23

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Definition: flattenings

$$p_{ijkl} = P(X_1 = i, X_2 = j, X_3 = k, X_4 = l)$$



**Theorem** (Chifman and Kubatko 2014): Under the coalescent model and the  $GTR+I+\Gamma$  model and its sub models, we have the following:

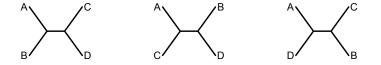
- If A|B is a valid split for a tree T, then  $rank(Flat_{A|B}(P)) \leq 10$ .
- If C|D is not a valid split for a tree T, then  $rank(Flat_{C|D}(P)) > 10$ .
- The species tree is completely determined by knowledge of valid splits on all quartets.

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• Arbitrary number of states,  $\kappa$ , under the coalescent model:

- If A|B is a valid split for a tree T, then  $rank(Flat_{A|B}(P)) \leq {\binom{\kappa+1}{2}}$ .
- If C|D is not a valid split for a tree T, then  $rank(Flat_{C|D}(P)) > \binom{\kappa+1}{2}$ .
- The species tree is completely determined by knowledge of valid splits on all quartets.
- Single underlying gene tree (no coalescent assumption):
  - If A|B is a valid split for a tree T, then  $rank(Flat_{A|B}(P)) \leq 4$ .
  - If C|D is not a valid split for a tree T, then  $rank(Flat_{C|D}(P)) = 16$ .
  - The species tree is completely determined by knowledge of valid splits on all quartets.

Main idea: use the observed site pattern distribution to provide information about which of the three possible splits for a set of four taxa is the true split.



The program SVDquartets computes a score for each split in a given quartet of taxa and chooses the split with the best (lowest) score.

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Species tree estimation using SVDquartets

# Algorithm

- Generate all quartets (small problems) or sample quartets (large problems)
- Stimate the correct quartet relationship for each sampled quartet
- Output to build the tree
  - PAUP\* uses the method of Reaz-Bayzid-Rahman (2014), called QFM, to build the tree.

Species tree estimation using SVDquartets

- Variability in the estimated tree is assessed using nonparametric bootstrapping
- Multiple lineages are handled as follows:
  - Sample four species
  - Select one lineage at random from each species
  - **③** Estimate the quartet relationships among the four sampled lineages
  - Restore the species labels (but lineage quartets are saved, too)

# Species tree estimation using SVDquartets

- Advantages:
  - Fast! How fast?
    - ★ Rattlesnakes: j 1 hour (~ 8500bp, 52 tips)
    - \* Soybeans: j 1 day (6 million SNPs, 62 tips)
  - Scales well:
    - Number of quartets needed increases as number of species increases (but can be done in parallel)
    - \* Linear in number of sites (but this is just counting)
  - Potential for application to other data types
  - Natural way to handle missing data
- Disadvantages:
  - Only the (unrooted) topology is estimated no parameters

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Described in the paper

Chifman, J. and L. Kubatko. 2014. Quartet inference from SNP data under the coalescent model, *Bioinformatics* 

- Implemented in PAUP\* thanks, Dave!
- Now on to the tutorial!

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