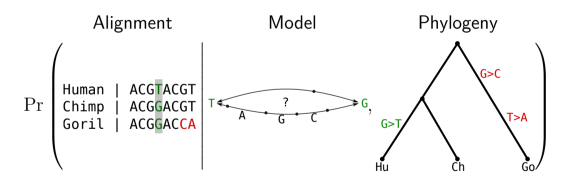
Polymorphism-aware phylogenetic models MIC-Phy 2021

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February 16, 2021

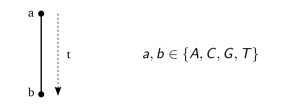


Comparative genomics and phylogenetics

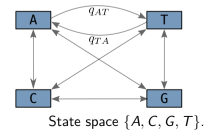


DNA substitution models

Evolution as a series of substitutions



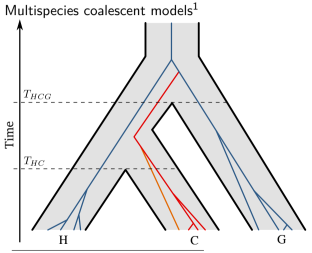
$$\Pr(X_t = b | X_0 = a) = \left(e^{tQ}\right)_{ab}$$

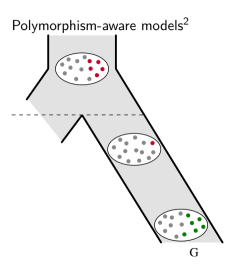


$$\boldsymbol{Q} = \begin{pmatrix} \cdot & q_{AC} & q_{AG} & q_{AT} \\ q_{CA} & \cdot & q_{CG} & q_{CT} \\ q_{GA} & q_{GC} & \cdot & q_{GT} \\ q_{TA} & q_{TC} & q_{TG} & \cdot \end{pmatrix}$$

Transition rate matrix.

Species are populations and recombination separates histories of genes Incomplete lineage sorting





¹Rannala and Yang (2003).

²De Maio et al. (2015) and Schrempf et al. (2016).

Neutral, K-allelic Wright-Fisher³ model

Discrete-time, discrete-state Markov process

N constant haploid population size.

K alleles.

 $m{z}_{ au}$ state of population $(z_{lpha},\ldots,z_{\kappa})$ in generation au; $\|m{z}_{ au}\|_1=N$; the total number of states is $\binom{N+\kappa-1}{\kappa-1}$.

 $m{U}$ $K \times K$ mutation probability matrix; the elements describe the probability to mutate from one state to another.

The distribution of alleles in the next generation $\tau+1$ is derived by sampling with replacement from the alleles of generation τ

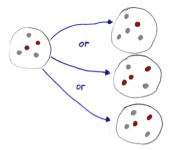
$$oldsymbol{z}_{ au+1} | oldsymbol{z}_{ au} \sim \mathsf{Mult}(oldsymbol{N}, rac{oldsymbol{z}_{ au}}{oldsymbol{N}} oldsymbol{U}).$$

For K = 4, and N = 10, we have 286 states.

³Wright (1931) and Fisher (1930).

Neutral, K-allelic Moran⁴ model with mutation

Continuous-time, discrete-state Markov process



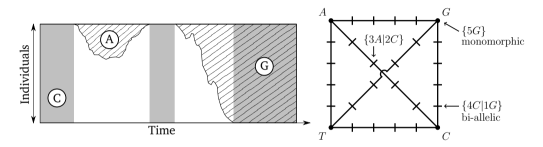
Individuals are randomly chosen to reproduce. The offspring is of the same type as the parent and replaces another randomly chosen individual from the population.

For
$$a,b\in\{\alpha,\ldots,\kappa\}$$
, $a\neq b$, and with mutation rates q_{ab} , we have
$$(\ldots,z_a,\ldots,z_b,\ldots)\to(\ldots,z_a-1,\ldots,z_b+1,\ldots)$$
 with rate
$$N\left(\frac{z_b}{N}\frac{z_a}{N}+\frac{z_a}{N}\frac{q_{ab}}{N}\right).$$

⁴Moran (1958).

Approximation for low mutation rates

Drift removes variation fast; disallow mutations when the population is polymorphic



Examples using nucleotides. Cartoon of evolving population with large size and state space for N=5.

Population can only be

Monomorphic
$$(..., z_a = N, ...) \equiv \{Na\}$$
; K states.

Bi-allelic
$$(\ldots, z_a = i, \ldots, z_b = N - i, \ldots) \equiv \{ia | (N - i)b\}; {K \choose 2} (N - 1) \text{ states.}$$

For K = 4, and N = 10, we have 4+54 states.

Discrete multivariate boundary mutation model⁵

From the Moran model with mutation, we have

$$(\ldots, z_a, \ldots, z_b, \ldots) \to (\ldots, z_a - 1, \ldots, z_b + 1, \ldots)$$
 at rate $\frac{z_a z_b}{N} + z_a \frac{q_{ab}}{N}$.

Transition rate matrix **M**

Boundary mutation leads to

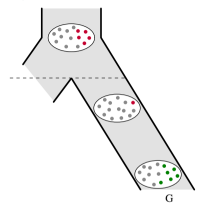
$$m_{\{Na\} \to \{(N-1)a|1b\}} = q_{ab}, \ m_{\{ia|(N-i)b\} \to \{(i\pm 1)a|(N-i\mp 1)b\}} = rac{i(N-i)}{N}.$$

⁵Schrempf and Hobolth (2017).

Polymorphism-aware phylogenetic Model (PoMo)

Use discrete multivariate boundary mutation model with

- K = 4 nucleotides;
- virtual population size N;
- transition rate matrix M.



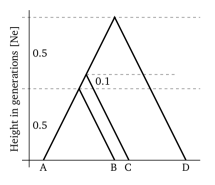
Likelihood calculation similar to DNA substitution models, for example,

$$\Pr\left(X_t = \{(N-1)a|1b\} \,|\, X_0 = \{\mathit{Na}\}\right) = \left(e^{t\textit{\textbf{M}}}\right)_{\{\mathit{Na}\}\{(N-1)a|1b\}}.$$

Q

Assessment of tree estimation error

Incomplete lineage sorting



Simulate

- Up to 1000 gene trees with the multispecies coalescent model; 10 samples per species⁶.
- **2** Sequences with 1000 base pairs per gene (HKY⁷ model); $\theta = 0.025^8$.

Infer phylogeny from data.

Measure branch score distance between original and estimated species tree.

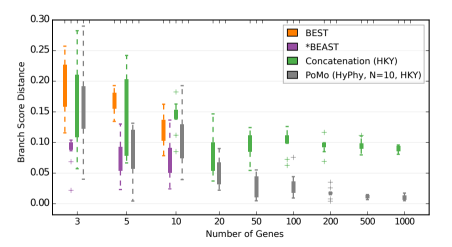
⁶MSMS, Ewing and Hermisson (2010).

⁷Hasegawa et al. (1985).

⁸SegGen, Rambaut and Grassly (1997).

Tree estimation error

Incomplete lineage sorting, $1N_e$ generations height, 10 samples per species



BEST (Liu 2008), *BEAST (Heled and Drummond 2010), and HyPhy (Pond et al. 2005).

Exchangeabilities, stationary distributions, and reversibility

Some mathematical prerequisites

The mutation rates can be separated into

$$q_{ab}=r_{ab}\pi_b,$$

where

 π_b is the stationary distribution of allele frequencies, and r_{ab} are the exchangeabilities.

If the mutation model is reversible, the exchangeabilities are symmetric $r_{ab} = r_{ba}$.

Theorem (Retention of reversibility of mutation model)

The discrete multivariate boundary mutation model is reversible if and only if the underlying mutation model is reversible.

Stationary distribution (reversible mutation model)

Theorem

For K,N>1 and reversible mutation models, the discrete multivariate boundary mutation model defined by the transition rate matrix ${\bf M}$ has a stationary distribution of

$$arphi_{\{\mathit{Na}\}} = rac{1}{Z}\pi_{\mathit{a}},$$

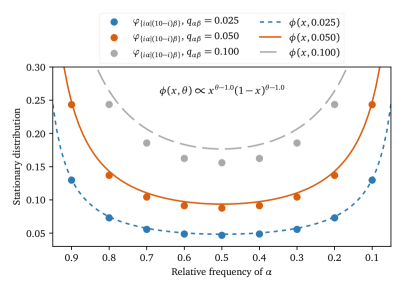
$$arphi_{\{\mathit{ia}\mid(\mathit{N}-\mathit{i})\mathit{b}\}} = rac{1}{Z}\pi_{\mathit{a}}\pi_{\mathit{b}}r_{\mathit{a}\mathit{b}}\left(rac{1}{\mathit{i}} + rac{1}{\mathit{N}-\mathit{i}}
ight)$$

with normalization constant

$$Z = 1 + \sum_{k=1}^{N-1} \frac{1}{k} \sum_{\substack{a,b \ a \neq b}} r_{ab} \pi_a \pi_b.$$

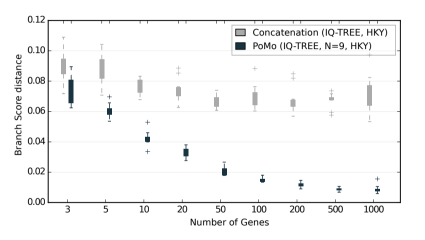
Stationary distribution

Alleles α and β (K=2), N=10, $q_{\alpha\beta}=q_{\beta\alpha}=\theta$; comparison to diffusion theory



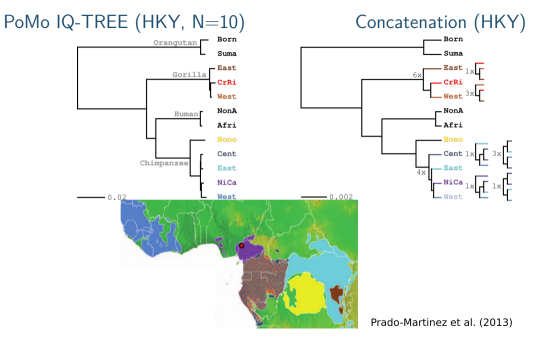
Tree estimation error

Yule 9 tree with 60 species, $3N_e$ generations height, 10 samples per species



IQ-TREE (Minh et al. 2020).

⁹Yule (1925).



Summary

Idea of PoMo

Model the evolution of populations and not of individuals in order to improve phylogenetic inferences.

Discrete multivariate boundary mutation model

- Moran model with mutations.
- 2 Approximation for low mutation rates.

Stationary distribution fits well if $\theta < 0.1$.

Suggestions for further reading

Non-reversible mutation models¹⁰

The stationary distribution is also known for non-reversible mutation models. Then, the mutation rate matrix Q can be separated into a reversible part, and a part describing circular probability flux.

Advanced mutation models¹¹



¹⁰Burden and Tang (2016) and Schrempf and Hobolth (2017).

¹¹Schrempf et al. (2019) and Borges et al. (2019).

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