Selecting models of nucleotide sequence evolution

Comparative Genomic Analyses

Rui Borges Vetmeduni Vienna

Selecting models of nucleotide evolution

An attractive procedure to select a model of evolution would be the use of the most complex parameter-rich one: e.g., GTR or UNREST.

- ▶ a large number of parameters need to be estimated
- ▶ the analyses become computationally difficult
- more error is introduced in each estimate

The fit of a model can be measured through the **likelihood function**.

▶ the likelihood expresses the probability of observing the data D, given a model of evolution M and a phylogeny T:

$$L = p(D|M, T)$$

Some models of evolution, or some phylogenies, will be more likely than others in explaining the data: **maximum likelihood estimates** (MLE)

► the values of the model parameters, the topology and branch lengths that make the likelihood functions as large as possible

$$\hat{M}, \hat{T} = \max p(D|M, T)$$

for computational reasons we work with the maximized log likelihood

$$\ell = \log p(D|\hat{M}, \hat{T})$$

A standard way of comparing the fit to two models of evolution is to contrast their log likelihoods using the **likelihood ratio test** (LRT) statistic:

$$LRT = 2(\ell_1 - \ell_0)$$

- $m \ell_1$: the maximum likelihood under the parameter richer complex model
- $ightharpoonup \ell_0$ the maximum likelihood under the simpler model

When the models compared are nested (i.e., the simple model is a special case of the complex model):

$$LRT \xrightarrow{D} \chi^2(\nu)$$

- \triangleright $\chi^2(\nu)$ distribution with ν degrees of freedom
- ightharpoonup
 u equal to the difference in the number of free parameters between the two models

The χ^2 distribution can be used to select the model of evolution that best fits the data

- ► LRT > critical value the inclusion of additional parameters increases the likelihood of observing the data significantly: the most complex model is favored
- ► LRT < critical value the simpler model is favored

	probability of the upper tail					
df	0.1	0.05	0.025	0.01	0.005	0.001
1	2.7055	3.8415	5.0239	6.6349	7.8794	10.827
2	4.6052	5.9915	7.3778	9.2103	10.596	13.815
3	6.2514	7.8147	9.3484	11.344	12.838	16.266
4	7.7794	9.4877	11.143	13.276	14.860	18.466
5	9.2364	11.070	12.832	15.086	16.749	20.515

Exercise

The likelihood of two models of evolution (JC69 and K80) was calculated for a multiple sequence alignment.

Model	lnL	np
JC69	-18 562.85	44
K80	-18 551.66	45

Perform an LRT and determine the model of evolution that best describes these sequences?

It is typically the case that models of sequence evolution are **nested**:

one model being equivalent to a restriction of one or more parameter values of a more complex model

K80 HKY85
$$\begin{bmatrix} . & \alpha & \beta & \alpha \\ \alpha & . & \alpha & \beta \\ \beta & \alpha & . & \alpha \\ \alpha & \beta & \alpha & . \end{bmatrix} \begin{bmatrix} . & \alpha \pi_{C} & \beta \pi_{G} & \alpha \pi_{T} \\ \alpha \pi_{A} & . & \alpha \pi_{G} & \beta \pi_{T} \\ \beta \pi_{A} & \alpha \pi_{C} & . & \alpha \pi_{T} \\ \alpha \pi_{A} & \beta \pi_{C} & \alpha \pi_{G} & . \end{bmatrix}$$

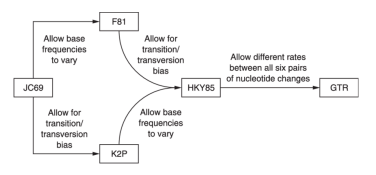
Exercise

Are the K80 and the F81 models of evolution nested? Justify

Several hypotheses can be tested hierarchically to select the best fitting model of evolution:

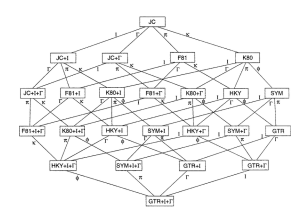
- ▶ are the base frequencies equal?
- ▶ is there a transition/transversion bias
- are the transition rates equal?
- ▶ are there invariable sites?
- ▶ is there rate homogeneity among sites

When comparing two nested models through an LRT, we test the different assumptions made by these models about our sequences.



There are some potential problems with the use of pairwise LRTs in a hierarchical LRT approach.

many hierarchies of LRTs are possible, and they can result in different models being selected



Literature

The Phylogenetic Handbook by Lemey, Salemi and Vandamme (2009) Cambridge University Press

► Chapter 10: sections 10.1, 10.2 and 10.3