Maximum Likelihood and the Bootstrap

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September 29, 2011

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Principle of Maximum Likelihood

- Given parameters θ and data X
- The function $f(X | \theta)$ is the probability of observing data X given parameter θ . (Both X and θ can be multi-dimensional.)
- Keeping θ fixed, and treating f as a function of X, the total probability is one.

Principle of Maximum Likelihood

- The function $L(\theta) = f(X | \theta)$ with X fixed and θ unknown is called the *likelihood function*.
- The *principle of maximum likelihood* is to estimate θ with the value $\hat{\theta}$ that maximizes $L(\theta)$.
- In practice, it is common to maximize the log-likelihood, $\ell(\theta) = \ln L(\theta)$.
- This is because X often takes the form of an independent sample so that

$$L(X) = \prod_{i=1}^{n} f(X_i \mid \theta), \qquad \ell(\theta) = \sum_{i=1}^{n} \ln f(X_i \mid \theta)$$

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Coin-tossing Example

- A coin has a probability θ of being a head.
- Consider tossing the coin 100 times. The probability of each single sequence with exactly x heads is $f(x | \theta) = p^x (1 p)^{100-x}$.
- Say we observe the sequence

where heads appear 57 times.

• The maximum likelihood estimate is the value $\hat{\theta}$ that maximizes the function

$$L(\theta) = \theta^{57} (1-\theta)^{43},$$

or, equivalently that maximizes

$$\ell(heta) = 57(\ln heta) + 43(\ln(1- heta)) \;.$$

Simple calculus and common sense lead to the estimate $\hat{\theta} = 0.57$.

Maximum-likelihood edge lengths

- For the Jukes-Cantor model, a pair of sequences have x sites with observed differences and n x sites with the same base.
- The probability of any given sequence pair is

$$L(d) = \left(\frac{1}{4}\right)^{n} \times \left(\frac{1}{4} - \frac{1}{4}e^{-\frac{4}{3}d}\right)^{x} \times \left(\frac{1}{4} + \frac{3}{4}e^{-\frac{4}{3}d}\right)^{n-x}$$

which has the form

$$L(\theta) = C \times \theta^{x} (1 - 3\theta)^{n-x}$$

where

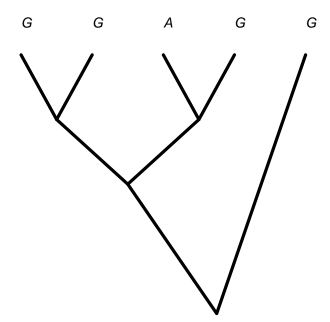
$$\theta = \frac{1}{4} - \frac{1}{4} \mathrm{e}^{-\frac{4}{3}d}$$

- Solving the calculus problem yields $\hat{\theta} = \frac{x}{3n}$.
- Plugging in and solving for *d* gives

$$\hat{d} = -\frac{3}{4}\ln\left(1 - \frac{4}{3}\frac{x}{n}\right)$$

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Computing Likelihood on a Tree



Transition Probabilities

$$P(0.1) = \begin{bmatrix} 0.90 & 0.04 & 0.04 & 0.03 \\ 0.03 & 0.91 & 0.04 & 0.03 \\ 0.03 & 0.04 & 0.91 & 0.03 \\ 0.03 & 0.04 & 0.04 & 0.90 \end{bmatrix} P(0.2) = \begin{bmatrix} 0.81 & 0.07 & 0.07 & 0.05 \\ 0.05 & 0.83 & 0.07 & 0.05 \\ 0.05 & 0.07 & 0.83 & 0.05 \\ 0.05 & 0.07 & 0.07 & 0.81 \end{bmatrix}$$
$$P(0.4) = \begin{bmatrix} 0.67 & 0.13 & 0.13 & 0.08 \\ 0.08 & 0.71 & 0.13 & 0.08 \\ 0.08 & 0.13 & 0.71 & 0.08 \\ 0.08 & 0.13 & 0.13 & 0.67 \end{bmatrix}$$

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Model Selection

12 rbcL genes from 12 plant species

Model	р	ℓ
JC69	21	-6262.01
K80	22	-6113.86
HKY85	25	-6101.76
HKY85 + Γ_5	26	-5764.26
HKY85 + C	35	-5624.70

• The AIC criterion is to select the model with the lowest AIC score, which is

 $AIC = -2 \ln(likelihood) + 2 \times (\# \text{ of parameters})$

- AIC balances the competing goals to fit the data well (likelihood high) and keep the model simple (few parameters).
- For this data, the HKY85+C model is the best among those compared; using 9 more parameters yielded an improvement in loglikelihood of over 139, which lowered the AIC by about 130.

The Bootstrap: A brief history

- The bootstrap was introduced to the world by Brad Efron, chair of the Department of Statistics at Stanford University, in 1979.
- The bootstrap is one of the most widely used new method in statistics that was invented within the past 50 years.
- In a special issue of *Statistical Science* that celebrates the 25th anniversary of the bootstrap, Brad Efron uses its application to phylogenetics as one of a small number of examples to illustrate its use and importance.

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The General Bootstrap Framework

- We have a sample x₁,..., x_n drawn from a distribution F from which we wish to estimate a parameter θ using a statistic θ̂ = T(x₁,..., x_n). (We might think of θ as being the median of the distribution, for example, and θ̂ = T(x₁,..., x_n) as the sample median.)
- If we wanted to compute the standard error of the estimate, we would ideally compute the standard deviation of T(X₁,...,X_n) where X_i ~ iid F.
- We could estimate this to any desired degree of accuracy by generating a large enough number (say B) of random samples X₁,..., X_n, computing \(\heta_i = T(X_1,...,X_n)\) for the *i*th such sample, and then computing the standard deviation of these estimates.

$$\sqrt{rac{\sum_{i=1}^{B}(\hat{ heta}_i- heta)^2}{B}}$$

The Key Idea

- Unfortunately, we cannot take multiple samples from F.
- However, our original sample x_1, \ldots, x_n is an estimate of the distribution *F*.
- Instead of taking samples from F, we could sample from the estimated distribution \hat{F} by sampling from our original sample with replacement.

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The Procedure

- We sample *n* values x_1^*, \ldots, x_n^* with replacement from x_1, \ldots, x_n .
- It is very likely that some of the original x values will be sampled multiple times and others will not be sampled at all.
- For each sample, compute the estimate of θ using the original statistic.
- The *i*th estimate is $\hat{\theta}_i^* = T(x_1^*, \dots, x_n^*)$.
- Repeat this *B* times and compute the standard deviation of the bootstrap estimates around the estimate from the original sample.

$$\sqrt{\frac{\sum_{i=1}^{B}(\hat{\theta}_{i}^{*}-\hat{\theta})^{2}}{B}}$$

Why it works

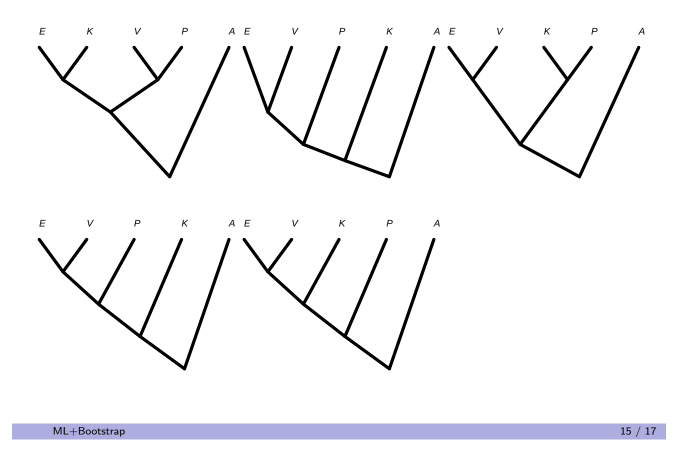
- If the sampling distribution of the bootstrap sample estimate θ^{*} around the estimate θ is similar to the sampling distribution of the estimate θ around the true value θ, then the boostrap standard error will be a good estimate of the real standard error.
- The bootstrap can be used to estimate bias, variance, for confidence intervals, and for hypothesis testing in many situations.
- It does depend critically on the assumption of independence of the original sample.

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Consensus Trees

- A *strict consensus tree* shows only those clades that appear in every sampled tree.
- A *majority rule consensus tree* shows all clades that appear in more than half the sample of trees.
- (Notice that two clades that each appear in more than half the sampled trees must appear in at least one tree together, implying that they are compatible with one another.)
- A *priority consensus tree* adds clades to the majority rule consensus tree in order of decreasing frequency in the sample provided that these clades do not conflict with a clade with higher frequency.

Example



Dynamic Exploration of Tree Samples

- Show off Mark Derthick's **Summary Tree Explorer**.
- Software is free and available at http://cityscape.inf.cs.cmu.edu/phylogeny/.

Interpretation of Bootstrap Proportions

What does a bootstrap porportion mean? Let me count the ways.

- *Confidence* that the clade is in the true tree.
- Bayesian posterior probability that the clade is in the true tree.
- One minus p-value for a formal hypothesis test that the clade is in the true tree.
- Rough measure of method robustness.
- Measure of repeatability of the inferences for the method at hand.
- Others?

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