Reconstructing ancestral character states

The pathways and patterns shown by character evolution are important to our understanding of evolutionary processes and predicting the future. Could we see into the past, we would like to know the history of changes both in time and in relationship to each other. This would allow us to learn how many times traits evolved, the conditions that favored their evolution, and the lineages in which changes occurred. Of course, in most cases we don’t know the past so we have to be satisfied with inferring the history of character evolution indirectly (“statistical paleontology”). A number of methods have been developed to do this.

Some examples of ancestral character state reconstruction

- Michael Ryan and Stanley Rand (1995) reconstructed ancestral advertisement calls in Tungara Frogs and observed reactions of female frogs to them.
- Gaucher et al. (2003) explored the palaeoenvironment of ancient bacteria on the basis of resurrected proteins.
- Bridgham et al. (2009) investigated the importance of epistasis in protein evolution.

Do early branching lineages signify ancestral traits (after Crisp and Cook, 2005)?

Failure to distinguish between present-day descendants and long-dead ancestors has led to incorrect interpretation of phylogenetic trees. Misinterpretation becomes evident when authors use the terms “basal” or “early diverging” to refer to extant taxa.

For example are monotremes “ancient/ancestral mammals”? Need to remember that whereas some character states can be relatively ancestral (plesiomorphic) or derived (apomorphic), these concepts should not be applied to whole organisms. Only nodes on a tree can be referred to as “basal” or “derived” relative to each other.

A common misinterpretation occurs when a species-poor/less diverse sister group is labeled as “basal” or “early-diverging” with respect to its species-rich sister. This misreading of phylogenies is encouraged by asymmetrical trees. Simplistic interpretation of phylogenies is also common in biogeography, where, for example, researchers may attempt to trace the history of a lineage back to its ancestral area.

Methods and models for inferring ancestral states

In reconstructing character evolution, our goal is to reconstruct the character states of hypothetical ancestors throughout the phylogenetic tree. The usual practice is to estimate the states of the hypothetical ancestors at the nodes or branch points of the tree, because these provide convenient reference points. Many methods are available (e.g., table 1 from Crisp and Cook below)

Parsimony methods

Parsimony methods seek to reconstruct ancestral states by choosing the states at the branch points that require the minimal number of steps on the tree.
A state can be placed at the node parsimoniously if it allows the observed states in the taxa to be evolved in as few as possible evolutionary steps. If more than one state can be placed parsimoniously at a node, the node’s assignment is said to be equivocal. The set of all the states that can be parsimoniously assigned to a node is called the node’s most parsimonious reconstructions (MPR) set.

A reconstruction is a set of assignments to the nodes such that each node is assigned only one state. You can think of it as one particular scenario for the evolution of the trait in question. The meaning of "reconstruction" given above is a very specific one — a fully resolved (lacking ambiguity) scenario for the evolution of the character, node by node, on the tree.

**Finding the most parsimonious ancestral states**

Ancestral state reconstruction using parsimony is done in two or three passes up and down the tree. The details of the process may differ with the different assumptions used; the algorithm for unordered characters is presented below. It creates three sets of character states. The first set is called the downpass set of the node, because this is the set of states preferred by that part of the tree above the node. The second set is called uppass set and is created by uppass optimization. The uppass state set of a node is calculated from the uppass set of the node below it (its ancestor)
Example from the class:

with the downpass set of the node beside it (its sister node), using the same calculation for combining state sets as for the downpass. The final set combines the first two and is called the MPR set of the node.

Reconstruction uncertainty.

"Character state reconstructions can provide a powerful mechanism for studying many facets of the evolutionary process. However, the zeal with which these techniques are sometimes advocated belies the complexity of the problem"

The first expression of uncertainty is ambiguity in the reconstruction or the existence of multiple equally parsimonious reconstructions. Ideally, when faced with multiple reconstructions, we should examine all of them. In reality, the number of possible reconstruction is often prohibitory large and only a few of them are examined/presented.

Two popular methods of choice are ACCTRAN and DELTRAN, which yield extremes of reversals versus parallelisms in the reconstruction.

ACCTRAN and DELTRAN
For characters of unordered and ordered type, ambiguities in character tracings can be resolved so as to choose the assignments that delay or accelerate transformations (look at the figure above and think what will happen to our reconstruction if we assign white/black color to node G). The DELTRAN option prefers most parsimonious assignments that delay changes away from the root; this maximizes parallel changes. The ACCTRAN tracing shows those assignments that accelerate changes toward the root; this procedure maximizes early gains and thus forces subsequent reversals. ACCTRAN and DELTRAN are but two of various methods to select from among the most parsimonious reconstructions.
Uncertainty does not end with ambiguity in the reconstruction. Even unequivocal assignment may be incorrect. Reconstructions of ancestral states are subject to error, as are all estimates of history. Studies of the reliability of ancestral state reconstructions have yielded mixed results, although it is clear that when rates of evolution are high over the time scale of the tree, error rates can be high. Also, the rate of change from one character state to another character state can be different (e.g., losses can be much easier than gains).

The following notes is a synopsis of a recent ms by Joy et al. 2016: (PLoS Comput Biol 12(7): e1004763. doi:10.1371/journal.pcbi.1004763)

Probabilistic methods (ML and Bayesian)

Hence the use of probabilistic methods (ML) provides several advantages:
   1) Use an explicit model of character evolution
   2) Consider branch length
   3) Can estimate the relative probability of each character state at every node.

Many models have been developed to estimate ancestral states of discrete and continuous characters from phylogenetic data and character states of extant descendants. Such models use some well-understood random processes assume to model the evolution of a trait through time. For discrete-valued traits (such as "number of legs"), this random process is typically taken to be a Markov chain; for continuous-valued traits (such as "brain mass"), the process is frequently taken to be a Brownian motion or an Ornstein-Uhlenbeck process.

Discrete-state models
If one wishes to recover the state of a given ancestral node in the phylogeny (call this node \( \alpha \)) by maximum likelihood, the procedure is: find the maximum likelihood estimate of substitution rates; then compute the likelihood of each possible state for \( \alpha \) given these rates; finally, choose the ancestral state which maximizes this. One may also use this substitution model as the basis for a Bayesian inference procedure, which would consider the posterior belief in the state of an ancestral node given some user-chosen prior. Because such models may have as many as \( k(k-1) \) parameters, overfitting may be an issue.

Binary state speciation and extinction model
The binary state speciation and extinction model (BiSSE) is a discrete-space model that allows estimation of ancestral binary character states jointly with diversification rates associated with different character states; it may also be straightforwardly extended to a more general multiple discrete-state model. In its most basic form, this model involves 6 parameters: 2 speciation rates (one each for lineages in states 0 and 1); similarly, 2 extinction rates; and 2 rates of character change. This model allows for hypothesis testing on the rates of speciation/extinction/character change, at the cost of increasing the number of parameters.

Continuous-state models
Although continuous traits can be split into discrete categories, it is more appropriate to model their evolution as some continuous process. In this case the likelihoods of transitions in state between adjacent nodes will be given by a continuous probability distribution such as Brownian motion. In this case, if nodes \( \alpha \) and \( \beta \) are adjacent in the phylogeny (say \( \alpha \) is the ancestor of \( \beta \)) and separated by a branch of length \( t \), the likelihood of a transition from \( \alpha \) being in state \( x \) to \( \beta \) being in state \( y \) is given by a Gaussian density with mean \( 0 \) and variance \( \sigma^2 t \). In this case, there is
only one parameter ($\sigma^2$), and the model assumes that the trait evolves freely without a bias toward increase or decrease, and that the rate of change is constant throughout the branches of the phylogenetic tree.

**Ornstein-Uhlenbeck process**: in brief, an Ornstein-Uhlenbeck process is a continuous stochastic process that behaves like a Brownian motion, but attracted toward some central value, where the strength of the attraction increases with the distance from that value. This is useful for modelling scenarios where the trait is subject to stabilizing selection around a certain value (say 0). Under this model the above-described transition of $\alpha$ being in state $x$ to $\beta$ being in state $y$ would have likelihood defined by the transition density of an Ornstein-Uhlenbeck process with two parameters: $\sigma^2$, which describes the variance of the driving Brownian motion, and $\alpha$, which describes the strength of its attraction to 0. As $\alpha$ tends to 0, the process is less and less constrained by its attraction to 0 and the process becomes a Brownian motion. Because of this, the models may be nested, and log-likelihood ratio tests discerning which of the two models is appropriate may be carried out.

**Stable models of continuous character evolution**: though Brownian motion is appealing and tractable as a model of continuous evolution, it does not permit non-neutrality in its basic form, nor does it provide for any variation in the rate of evolution over time. Instead, one may use a stable process, one whose values at fixed times are distributed as stable distributions, to model the evolution of traits. Stable processes, roughly speaking, behave as Brownian motions that also incorporate discontinuous jumps. This allows one to appropriately model scenarios in which short bursts of fast trait evolution are expected. In this setting, maximum likelihood methods are poorly suited due to a rugged likelihood surface and because the likelihood may be made arbitrarily large, so Bayesian methods are more appropriate.

Bayesian methods provide additional advantages
  1) avoid potential errors in fixed parameter estimates
  2) address phylogenetic uncertainty

**Software:**
There are many software packages available, which can perform ancestral state reconstruction a selection is described below:

Within the **R statistical language**:
- **APE** implements a variety of ancestral state reconstruction methods for both discrete and continuous characters.
- **Diversitree** implements ancestral state reconstruction under Mk2 and BiSSE models.
- **HyPhy** is a modular software package for hypothesis testing using phylogenies in a maximum likelihood framework. HyPhy implements a fast joint likelihood method of ancestral sequence reconstruction that can be readily adapted to reconstructing discrete ancestral character states or geographic ranges.
- **Mesquite** implements parsimony, maximum likelihood, and Bayesian methods of ancestral state reconstruction for both discrete and continuous characters and has several display methods for resulting reconstructions.
- **Bayes Traits** is a computer package which performs analyses on discrete or continuous characters in a Bayesian framework and allows testing of hypotheses about models of evolution, ancestral states, and correlations among pairs of traits.
**BEAST** Also performs ancestral state and ancestral sequence reconstruction analyses.

**Lagrangle** is an application which allows analyses reconstruction of geographic range evolution on phylogenetic trees available from http://www.reelab.net/home/software/.

**Phylomapper** implements a likelihood-based statistical framework for estimating historical patterns of gene flow and ancestral geographic locations. Available from: http://www.evotutor.org/LemmonLab/PhyloMapper1.html