Polytomies in Comparative Analyses of Continuous Characters

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Felsenstein's (1985) method of phylogenetically independent contrasts was the first statistically valid parametric method for correlation and regression analysis of continuous variables measured on multiple species. One requirement of the method is that the true phylogeny linking the species in the data set be known in full, including the lengths of the branches in units of expected variance of change for the character(s) being studied (see Harvey and Pagel, 1991; Martins and Garland, 1991; Garland et al., 1992). In practice, we cannot be certain of the true phylogeny for any set of species. Instead, we must use an estimate of it. This working phylogeny (sensu Grafen, 1989) might differ from the real phylogeny in three ways: (1) taxa depicted as sister groups in the working phylogeny might not be so in reality, causing the wrong taxa to be compared when computing contrasts; (2) the working phylogeny might not depict all of the real sister-group relationships, i.e., a series of bifurcations in the true phylogeny might be represented as a polytomy in the working phylogeny; and (3) the lengths of branches between nodes (including tips) of the working phylogeny might be incorrect.

Differences of the first type clearly must be minimized because they violate the assumptions of Felsenstein's method to an unknown and possibly very serious degree. Differences of the third type also weaken the method, but apparently not fatally (see Grafen, 1989; Martins and Garland, 1991). However, we still want the branch lengths on the working phylogeny to reflect the truth as accurately as possible (Garland et al., 1992). Differences of the second type—incomplete resolution of the working phylogeny—raise some interesting conceptual and analytical issues. Unless we are prepared to assume that speciation is always a strictly dichotomous process, then we must recognize that some polytomies in working phylogenies will represent true cases of multiway speciation events; these are termed “hard” polytomies (Maddison, 1989). In the original description of the comparative method of phylogenetically independent contrasts, Felsenstein (1985:10) alluded to the correct way to analyze hard polytomies. However, other polytomies in working phylogenies simply reflect our ignorance of the true dichotomous branching pattern (“soft” polytomies, Maddison, 1989). Felsenstein's suggestion to set some internode branch lengths to zero when computing standardized independent contrasts leads to unbiased estimates of correlations and slopes for both hard and soft polytomies but does not indicate how many degrees of freedom are available for hypothesis testing with the latter. Grafen (1989), Harvey and Pagel (1991), and Pagel (1992) have offered other more complicated ways of dealing with soft polytomies that allow only one degree of freedom per node. Table 1 presents the differences between the two positions. We want to recognize any hard polytomies as such to claim the maximum degrees of freedom associated with them.

Hard and Soft Polytomies

Independent contrasts approaches use phylogenetic information to transform
TABLE 1. How the assumption of "hard" versus "soft" polytomies affects independent contrasts methods. When polytomous nodes are present, the number of contrasts computed and used to estimate the relationship between two traits should always be \( n - 1 \), as in a fully resolved \( n \)-taxon tree, whereas the associated degrees of freedom are reduced when soft polytomies are present. This table presumes bivariate regression (or correlation) through the origin; in multiple regression through the origin, one additional degree of freedom would be lost for each additional independent variable.

<table>
<thead>
<tr>
<th>All polytomies</th>
<th>All polytomies</th>
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</thead>
<tbody>
<tr>
<td>hard</td>
<td>soft</td>
</tr>
<tr>
<td>Contrasts at an ( n )-taxon polytomy</td>
<td>( n - 1 )</td>
</tr>
<tr>
<td>Contrasts in a fully resolved ( n )-taxon tree</td>
<td>( n - 1 )</td>
</tr>
<tr>
<td>Contrasts in an ( n )-taxon working phylogeny that contains some unresolved nodes</td>
<td>( n - 1 )</td>
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<tr>
<td>Degrees of freedom for an ( n )-taxon polytomy</td>
<td>( n - 1 )</td>
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<tr>
<td>Degrees of freedom for an ( n )-taxon working phylogeny that contains ( p ) nodes (including the root)</td>
<td>( n - 1 )</td>
</tr>
<tr>
<td>Test for significance</td>
<td>regression through origin</td>
</tr>
<tr>
<td>Degrees of freedom in test</td>
<td>( n - 1 - 1 )</td>
</tr>
</tbody>
</table>

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\(^b\) Except when the tree is completely unresolved (i.e., a star), in which case the minimum is 1, not 0, df. See also Figure 3.

...values for terminal taxa to eliminate statistical nonindependence. The working phylogeny specifies what pattern of non-independence we expect or, more specifically, the expected variances and covariances of the phenotypes of all terminal taxa. Hard and soft polytomies represent different specifications of phylogenetic pattern and so must be treated differently for hypothesis testing.

Instantaneous Radiations: Hard Polytomies

Felsenstein's (1985) original paper mentioned how hard polytomies should be analyzed: they "can always be represented as a series of bifurcations having some branch lengths zero" (Felsenstein, 1985:10). At an \( n \)-taxon node, then, Felsenstein advocated that the polytomy be resolved arbitrarily into a series of bifurcations separated by branches of zero length. For each bifurcation, a contrast is computed and standardized to unit variance by dividing it by the square root of the sum of its branch lengths. On first consideration, it might seem that the result of the analysis will depend on how the polytomy is resolved; a node with \( n \) taxa can be resolved in \( (2n - 3)!/2^{n-2}(n - 2)! \) different ways (Felsenstein, 1978) (e.g., for eight taxa, 135135 different resolutions are possible). However, all the different possible sets of contrasts give the same result when analyzed by regression through the origin; the slope and correlation coefficient, and hence the significance of the relationship between two variables as judged with \( n - 2 \) degrees of freedom, are the same in every case. Figure 1 shows the three possible resolutions of a three-way polytomy, the plot of the standardized independent contrasts from each of them, and the nonphylogenetic regression (in which species values are taken to be independent and degrees of freedom are \( n - 2 \) because an intercept is estimated). This congruence between phylogenetic and nonphylogenetic methods should not surprise us because the evolutionary models underlying the two approaches are identical: the subtaxa are all assumed to have radiated simultaneously from their common ancestor and hence to be statistically independent (Harvey and Pagel, 1991). The independent-contrasts method gives the correct estimates of correlation and \( P \) values even when applied to a star phylogeny; thus, it does not mislead, contrary to suggestions in Gittleman and Luh (1992:401).

Lack of Information: Soft Polytomies

In practice, working phylogenies often contain polytomies that definitely do not represent simultaneous radiations. For example, the bat genus *Rhinolophus* contains 64 extant species (Corbet and Hill, 1991), and little is known of its intrageneric phy-
A single lineage split instantaneously, giving rise to three modern descendants, A, B, and C. (a) Present-day values of two characters, $x$ and $y$, for each species and the regression of species values. (b) Three ways that the polytomy can be resolved into bifurcations separated by a branch of zero length (the other branches still have length $v$) and the standardized independent contrasts ($\Delta x$ and $\Delta y$) calculated according to Felsenstein (1985). Note that for each contrast the direction of subtraction is arbitrary and thus so are the signs. (c) All three resolutions give exactly the same slope, correlation, and $P$ value when the contrasts are analyzed by regression through the origin. Furthermore, these estimates are the same as those given by the regression of species values shown in (a).
logeny. Treating the phylogeny of this genus as a star leading to 64 tips most likely would not be a good representation of biological reality. Rather, some groups of species within the genus are closely related, but we do not know which ones. Such sister-group relationships that are not represented in the working phylogeny are known as "unrecognized phylogeny" (Grafen, 1989), which is essentially a synonym for soft polytomy. With respect to hypothesis testing, unrecognized phylogeny invalidates the procedure described above for independent contrasts approaches and for any other methods that assume all polytomies to be hard (e.g., Cheverud et al., 1985; Bell, 1989; Gittleman and Kot, 1990; Lynch, 1991) in exactly the same way that phylogeny in general invalidates non-phylogenetic regressions; some groups of taxa share character states by common descent rather than by convergent evolution, so their values are not statistically independent. Figure 2 highlights the difference between a hypothetical working phylogeny with soft polytomies and the associated "true" phylogeny. Garland et al. (1993) discussed how polytomies affect comparative methods that use computer-simulated null distributions for hypothesis testing (e.g., Martins and Garland, 1991).

Grafen (1989) suggested a way around the impasse of unrecognized phylogeny when using independent contrasts (Felsenstein, 1985) for multiple regression analyses. He assumed that hard polytomies are very much the exception in nature, so polytomies in the working phylogeny must be expressions of ignorance. Because of the statistical nonindependence among the \( n \) daughters of a node, it would be invalid to make \( n - 1 \) comparisons among them. Instead, Grafen's method collapses the information at a node into a single "comparison" (technically a linear contrast: see Grafen [1989, 1992] for a definition and details). By taking only one piece of information from each node and hence only one associated degree of freedom for hypothesis testing, Grafen's (1989) approach avoids inflated type I error rates due to unrecognized phylogeny. This principle that each node should contribute only one item of information to the statistical test was subsequently adopted in other linear contrasts approaches (i.e., modifications of Felsenstein's [1985] original presentation: Pagel and Harvey, 1989, 1992; Harvey and Pagel, 1991; Pagel, 1992). When applied to the working phylogeny shown in Figure 2a, these methods would compute only three contrasts: one within the taxon ABCDE, one within FGHI, and one between these two taxa.

The distinction between statistical estimation and hypothesis testing is important here. The various works by Grafen, Harvey, and Pagel have emphasized hypothesis testing. The primary aim was to avoid overestimating the degrees of freedom associated with an incompletely resolved working phylogeny. Table 1 lists the degrees of freedom associated with resolved versus unresolved nodes and with working phylogenies containing an arbitrary number of nodes (\( p \); including the root node), regardless of whether they are dichotomous or polytomous, in a way fully consistent with the procedures of Grafen, Harvey, and Pagel. When unresolved nodes are present, it is only possible to bound the degrees of freedom; they will be no more than \( n - 2 \) (i.e., the number of contrasts minus one, when all polytomies are hard) and no fewer than the number of nodes minus 1 (or 1, as opposed to 0, in the special case of a completely unresolved star phylogeny). Significance tests will be liberal with \( n - 2 \) degrees of freedom and conservative with \( p - 1 \) degrees of freedom. Statistical power will be affected accordingly. Figure 3 illustrates an example for a phylogeny of eight species. When reporting such uncertainty about \( P \) values, it may help to indicate the fewest degrees of freedom that would be required for a given relationship to be considered statistically significant, e.g., at \( \alpha = 0.05 \) (cf. Walton, 1993: table 3).

We differ from Grafen, Harvey, and Pagel as to how many contrasts should be computed for an unresolved node and hence how one should estimate an evolutionary relationship. Computing only a
single contrast for each unresolved node will lead to cases in which only one (e.g., Fig. 3f) or a few (e.g., Figs. 2a, 3e) contrasts are computed for an entire tree, resulting in trimodal distributions of slopes or correlations (e.g., Gittleman and Luh, 1992) and poor estimates of evolutionary correlations (cf. Martins and Garland, 1991). We therefore recommend that the full $n - 1$ contrasts should always be computed and used for estimation by correlation or regression through the origin. This procedure is simple and less arbitrary than previous proposals, and it may lead to better estimates (e.g., lower mean squared error), although this has not yet been studied.

Near-simultaneous Radiations: "Firm" Polytomies

In some cases, an unresolved node may be fully dichotomous in reality but may actually resemble a hard polytomy. For example, the topology illustrated in Figure 4 is the same as that in Figure 2b, but the tree looks much more like that in Figure 2a because the shared branches within ABCDE and within FGHI are very short. Because the shared branches (unrecognized phylogeny; they are not present in Fig. 2a, which is being used as the working phylogeny) are so very much shorter than the unshared ones (independent evolution), the working phylogeny shown in Figure 2a is actually a rather good descrip-
FIGURE 4. The same topology as Figure 2b but with much less unrecognized phylogeny because the shared branches within taxa ABCDE and FGHI are very short.

1. The polytomy is based on one or more cladograms with many characters per taxon but no synapomorphies resolving the polytomy. The absence of characters suggesting an internal branch implies that this branch persisted for only a short time at most.

2. The polytomy is based on molecular distance analyses using full data matrices (all species contrasted to all others). If so, then branches not shown in the working phylogeny cannot be longer than the confidence limits associated with the estimates of node ages.

3. Fossil evidence shows the radiation to have been rapid.

In contrast, the following features are danger signs and indicate that a polytomy must be treated as soft to avoid inflated type I error rates.

1. The polytomy comes from a standard taxonomy featuring taxonomic levels.

2. The polytomy is from a molecular distance analysis where the data matrix is incomplete. If only some species are used as tracers, polytomies will inevitably result unless the tracers happen always to be the most distal taxa (cf. Bledsoe, 1987; Lanyon, 1992).

3. The polytomy is taken from one or more cladograms where the ratio of informative characters to taxa is low: resolution in such cladograms is likely to be poor whatever the true phylogeny (Henderson et al., 1989; Penny et al., 1991).

When in doubt as to the “firmness” of a given polytomy, assume the worst—that it hides lots of unrecognized phylogeny—and test hypotheses with the fewer degrees of freedom, as indicated in Table 1.

CONCLUSION

Comparative tests must be based on an estimate of phylogeny, which is often incompletely resolved. Most polytomies are
soft, hiding an unknown bifurcating structure (unrecognized phylogeny). For the method of phylogenetically independent contrasts, we consider how to treat soft polytomies for both estimation and hypothesis testing. Studies of interspecific allometry, for example, often aim to estimate the form of an evolutionary relationship between characters. Some previous workers have suggested complicated ways to extract a single independent contrast from a given soft polytomy. Instead, we argue (consistent with Felsenstein’s [1985] original suggestion) that soft polytomies should be resolved arbitrarily by setting branch lengths between internal nodes to zero. This procedure will yield a single slope or correlation coefficient, irrespective of how the polytomy is arbitrarily resolved, and unlike previous suggestions, it does not depend on the tip data being analyzed. When using independent contrasts to test hypotheses, however, unrecognized phylogeny must be taken into account or the test will invalidly claim too many degrees of freedom, thus yielding inflated type I errors. When soft polytomies are present, it is possible to bound the degrees of freedom available for hypothesis testing. Our pragmatic guidelines can be followed for assessing just how soft a polytomy might be so that the maximum reasonable degrees of freedom can be claimed.

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