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NOTES AND COMMENTS

HOMOLOGY AND COMPARATIVE METHODS IN THE STUDY OF AVIAN COOPERATIVE BREEDING

In an earlier article (Edwards and Naeem 1993), we presented a comparative analysis of cooperative breeding (CB) in perching birds (order Passeriformes). Using both discrete (categorical) and continuous data sets, we illustrated a number of ways in which comparative methods could help clarify the temporal origins of CB in various clades, ecological correlates of CB, and patterns of interspecific variation in attributes such as group size. The commentary on our article by McLennan and Brooks (1993) makes three main points, each of which raises important issues about the biology and ecology of CB in birds as well as the application of comparative methods to behavioral data. The first point is that comparative analyses of behavioral traits based on trees derived from taxonomies or noncladistic methods (phenetic clustering), “neither of which produces a robust phylogenetic hypothesis of genealogy” (McLennan and Brooks 1993, p. 790), are nonetheless useful for those researchers recognizing the importance of examining evolutionary hypotheses in a phylogenetic context. This point echoes our own plea for increased interest in building robust trees for ecologically and behaviorally diverse groups of birds (see, e.g., Richman and Price 1992), particularly for cooperative breeders (see, e.g., Edwards and Wilson 1990).

The analyses in our article were based on trees built with varied sources of data and methods of analysis: cladistic analysis of morphology (fairy wrens, *Malurus*), phenetic analysis of morphology (New World wrens, *Campylocercus*), taxonomy (e.g., Australian treecreepers, *Climacteris*), phenetic analysis of molecular distance data (e.g., New World jays), and cladistic analysis of DNA sequence data (Australian babblers, *Pomatostomus*). We agree that phylogenies are always preferable to taxonomies in comparative analyses, and, to the extent that it is difficult to model morphological evolution phenetically (Felsenstein 1988), our analysis in these cases should be treated as tentative. But to state that “phenetic” analyses categorically do not produce “robust” phylogenetic hypotheses—thereby implying that cladistic methods categorically produce better trees—is simply outdated. The philosophical underpinnings of cladistics are different from those of other methods, but this does not mean that “noncladistic” methods cannot yield truth in practice. Uncertainties in phylogenetic inference often lie more in limitations of data than in the particular method used (Miyamoto and
Cracraft 1991), and all tree-building methods have a specific region of parameter space in which they work best (Hendy and Penny 1989; Zharkikh and Li 1993). Under some circumstances, methods traditionally labeled as "cladistic" can converge on the wrong tree as more data are accumulated (Felsenstein 1978; Hendy and Penny 1989; Penny et al. 1992). Furthermore, several recently proposed comparative tests (see, e.g., Felsenstein 1985; Garland 1992) require estimates of branch lengths, which are arguably better obtained with noncladistic methods or, at the very least, with corrections on a purely cladistic approach (Fitch and Bruschi 1987). The above caveats particularly apply to the class of data of increasing use to comparative biologists, namely, DNA sequence data (Felsenstein 1988). As noted by McLennan and Brooks (1993), modern comparative biology is a relatively young field; in our opinion it would be unfortunate if the philosophical divisions characterizing the rise of quantitative systematic methods in the late 1960s and encapsulated in McLennan and Brooks’s statement were to recapitulate themselves in this new era of comparative methods.

ARE COMMUNAL BREEDING SYSTEMS HOMOLOGOUS?

McLennan and Brooks’s second point is that CB may in fact subsume a class or group of characters because of the inherent diversity among CB systems (e.g., in breeding systems, types and numbers of helpers and contexts of helping), in which case treating CB itself as homologous even among closely related species might be unwarranted. The concept of homology is still being refined (see, e.g., Patterson 1988; Wagner 1989), particularly as it is applied to behavior (Lorenz 1950; Mayr 1958; Atz 1970; McLennan et al. 1988; Wenzel 1992). For the purposes of this note, we define homology as "similarity due to inheritance from a common ancestor"; a homologous relationship of a character in different taxa can often be inferred only by determining whether that character displays congruence with (defines similar monophyletic groups as) other characters (Patterson 1982). Since we never denied that CB has developed multiple times in distantly related lineages of birds (Edwards and Naeem 1993, p. 771), we assume that McLennan and Brooks (1993) restrict the discussion of homology to closely related groups for which a hypothesis of homology of CB is reasonable; thus, for example, we agree that CB in New Zealand pukekos (Porphyrio porphyrio), a nonpasserine bird, probably shares few if any phylogenetic correlates with CB in passerines such as Florida scrub jays (Aphelocoma coerulescens coerulescens).

Although the assumption (or inference) of interspecific homology among behaviors continues to prove useful for understanding the evolution of behavior (McLennan et al. 1988; Prum 1990), it has been applied primarily to individual actions, ritualized movements, and "motor" responses, rather than to social systems per se. Since the early comparative studies of CB, the persistent observation that alloparrental behavior could be generated by multiple ecological and demographic mechanisms has not only compromised most attempts to generalize about CB across taxa (Smith 1990) but opens the possibility that CB may not be homologous even between closely related species. For example, one could argue that the multiple mechanisms for the origin of CB in New World jays proposed by
Brown (1974) provide evidence that CB is not homologous among these species because it has arisen in diverse demographic contexts in different species (although other interpretations of this article are certainly possible). In another view, White et al. (1991) suggested that few aspects of CB are actually "inherited" or arise in any long-term historical sequence of events; rather, group living literally develops anew every generation because of a constellation of recurrent, uninherited (though perhaps biased) responses of individuals to a repeated set of social and physical environmental factors. That CB develops by such routes is evident in birds such as the Seychelles warbler (*Acrocephalus sechellensis*), which was reported to have "evolved" cooperative breeding in a five-year period of population growth (1968–1973; Komdeur 1992)—a time span probably too short for any appreciable population genetic change. In such cases, CB is more appropriately considered convergent (i.e., nonhomologous), not only between species, but within species across successive generations (Lambert and Hughes 1988).

McLennan and Brooks's claim that we and other evolutionary biologists confuse behavioral characters with character classes constitutes another reason not to consider CB systems homologues. Our article argues against this point of view. First and most important, although we acknowledged that CB has arisen multiple times within the Passeriformes, in most of the phylogenetic analyses we presented, CB arose very few times (often once) within specific groups; thus, CB (as defined in our article) fulfilled the primary criterion of homology, namely, congruence with other sets of characters (Patterson 1988; Prum 1990). Similarly, other authors have found evidence for congruence (hence, homology) in group behaviors that McLennan and Brooks would consider character classes because they occur in a variety of social and breeding contexts (e.g., territoriality and coloniality: de Queiroz and Wimberger 1993; queen number in social insects: Ross and Carpenter 1991). Second, we suspect that social systems, which are necessarily defined by interactions among multiple, easily delimitable parts (namely, individuals), merely bring into sharper focus the dichotomy between character and character class that we believe exists at nearly all levels in the biological hierarchy. Many biologists argue that the definition of a character is itself controversial (Pogue and Mickevich 1990). An organ considered a character by an anatomist might be considered a character class by a histologist or cell biologist (although arguments against attempts to consider characters or taxa at more than one biological level simultaneously have been proposed [Liden 1990]).

Finally, even if CB were not homologous between species, it was not the only character considered in our article. We speculated that phylogenetic analysis of traits considered prerequisites for CB and group living (e.g., the "stimulus-response" behaviors in Jamieson 1989, delayed breeding, mechanisms of kin recognition) would help clarify patterns of behavioral evolution in cooperative breeders; such prerequisite traits only seem less atomizable than communal breeding systems to organismic biologists but would presumably be considered good characters by McLennan and Brooks (1993).

A more stringent criterion of behavioral homology requires the identification of congruent physiological and genetic structures underlying that behavior (Atz 1970; Wenzel 1992; fig. 1); in our view, such a criterion would increase confidence
in the search for behavioral homologues in cooperative breeders. Identifying the molecular bases of morphological changes has proved extremely effective in clarifying homologies in morphological evolution (see, e.g., Patel et al. 1989). Such efforts could help pinpoint aspects of CB that are homologous to behaviors found in noncooperative species, or—a more exciting prospect—behavioral components in cooperative breeders that are truly derived with respect to noncooperative species (Jamieson 1989). Although some argue that there are no truly derived behavioral or physiological processes distinguishing cooperative breeders from
nonsocial species (White et al. 1991, p. 250), there are few data, and it is not known whether such derived processes exist. Patterns of hormonal expression are known to differ among, for example, bird species displaying varying levels of male parental care (Wingfield 1991). In many respects, whether derived homologues exist in cooperative breeders is the central issue in recent debates about CB (see references in Jamieson 1991); identifying similarities and differences in the tangible components of these behaviors could help resolve these debates.

We attempted to reconcile the plasticity observed so often in CB with our inference of phylogenetic conservatism by positing variation in the expression of more conservative behavioral or structural prerequisites of CB, as in a threshold character whose state depends on interactions with the environment (see also Mumme 1992). More thoroughly studied communal systems not only illustrate the relationship we envision between such prerequisites and their expression as social behavior but provide a mechanism for their phylogenetic conservatism. We refer to the recent identification of some of the genes responsible for mediating mating preferences, kin recognition, and social interactions in mice (Potts et al. 1991; Manning et al. 1992). Although the type of social behavior arising from such a genetic system in different species would depend on the rules by which individuals process the information provided by that system, the genetic component of the social behavior would still be homologous among taxa because a specific set of inherited factors common to mammals would have been identified (Klein 1986).

Referring to cases in which “different” (i.e., nonhomologous) forms of CB are mistakenly lumped together under one category, McLennan and Brooks (1993, p. 791) argue that “there is no a priori reason to expect to find ecological correlates for its [CB’s] evolution.” We disagree. On the contrary, the ecological correlates and the adaptive basis of behaviors are probably best detected in cases of behavioral convergence (nonhomology; Ridley 1983). The presence of auxiliaries to the breeding pair is the cornerstone of CB, and much of the literature on CB seeks to determine the ecological and demographic correlates of this trait despite the diversity observed in mating systems and sex and number of helpers. Different forms of CB are all nonetheless characterized by “helpers”; these variations can usefully be considered character states, comparison of which can yield an improved picture of the ecological correlates of CB (Peterson and Burt 1992). Thus the critical question regarding McLennan and Brooks’s (1993) concern is not whether CB is a character or a character class but whether it forms a group of traits with functions similar enough that ecological correlates may be expected.

**ECOLOGICAL CORRELATES OF CB DETECTED USING PHYLOGENIES: THE IMPORTANCE OF MODELS**

Finally, McLennan and Brooks (1993) point out that in most cases we did not have enough information to draw firm conclusions about ecological correlates of CB because CB was plesiomorphic (i.e., ancestral) to several groups. We agree; as stated above, finding potential correlates for the origin of CB requires multiple
derivations of CB within the clade of interest (Ridley 1983; Maddison 1990). We were surprised to find that CB arose so deeply in groups in which CB was well studied but for which close out-groups have not been identified (e.g., Australian wrens [Malurus + Stipiturus + Amytornis] and babblers [Pomatostomus]). In the one case in which the latter criterion was possibly fulfilled (Eopsaltria robins), the phylogenetic relationships of the species in question were ambiguous. McLennan and Brooks (1993) presented two possible resolutions of the ambiguous relationships of Eopsaltria georgiana (their fig. 2), but a third resolution exists: that in which E. georgiana composes the out-group to Eopsaltria australis, Eopsaltria flaviventris, and Eopsaltria griseogularis. This third alternative in fact corresponds to the cladistic analysis of Cracraft (1982) (whom we neglected to cite in our original article) and implies a scenario similar to that in McLennan and Brooks’s figure 2b. Clearly, an understanding of the origins of CB in Eopsaltria must await a comprehensive phylogenetic analysis of all its members and relatives.

Models for the maintenance of CB solely via specific ecological regimes predict (1) a phylogenetic coincidence between the invasion of specific ecological regimes and the origin of CB and (2) a loss of CB coincident with lineages leaving such regimes. McLennan and Brooks (1993) acknowledge that detecting the latter pattern does not necessarily require that CB be derived within the clade of interest. Thus we found that several hypothesized ecological correlates of CB were in fact ancestral to the entire clade composed of both cooperative and noncooperative species (fig. 1B of Edwards and Naem 1993), thereby weakening ecological explanations for macroevolutionary trends in CB. In Pomatostomus and Malurus, CB persists in descendant lineages despite radical shifts in ecological regime, which again weakens causal links with CB (fig. 1C of Edwards and Naem 1993). Our article was written in part to challenge claims based on nonphylogenetic analyses that there were well-supported ecological correlates of CB (e.g., in Australian passerines); with the caution that a more rigorous comparative analysis should be done, our preliminary findings are in agreement with those nonphylogenetic studies that failed to detect ecological correlates of CB (see, e.g., Dow 1980; Brown 1987).

The case of Eopsaltria robins illustrates an important methodological issue. As elsewhere (e.g., Brooks and McLennan 1990), McLennan and Brooks are here more concerned with qualitative evidence of causal relationships than with formal hypothesis testing and estimation of correlations (e.g., by Ridley’s [1983] or Maddison’s [1990] tests). As with any comparative analysis, the significance of correlations between CB and ecological parameters will depend on the model of character evolution chosen (Harvey and Purvis 1991). This effect is illustrated with simulations in figure 2, in which both the time span in which CB is assumed to arise (t) and the probability of change in CB (α) influence the strength of correlations of CB with ecological variables. Early in our analyses we were well aware that, “because the reconstructions of CB on the trees generally involved one or a few origins per group, we could not hope to achieve statistically significant associations of CB and other variables within any taxonomic group using existing tests that assume parsimony. . . . Although the trait of interest [CB] may be
Fig. 2.—Assumptions about the evolution of cooperative breeding influence the statistical significance of correlations with ecological variables. A, Hypothetical tree of four species, with branch lengths indicated in arbitrary time units. Solid circles indicate transitions to cooperative breeding from an ancestral state (open circle) with probability \( \alpha \) (inset). Solid squares indicate a derived ecological regime. The two terminal branches bracketed with a \( t \) were varied in the simulations in B, as was \( \alpha \) in C. The tree and character distributions are identical to those in the example in box 4.6 of Harvey and Pagel (1991). B, Simulations showing the effect of varying the two terminal branch lengths \( t \) in A (X-axis) on the probability of observing the associations of cooperative breeding and the ecological variable in A (Y-axis) by chance. This probability is \( P(Y|X) \), where \( Y \) is cooperative breeding and \( X \) is the ecological variable (Maddison 1990; Harvey and Pagel 1991). Ecological correlations with behavioral traits will in general appear much stronger when the time span during which the transitions are assumed to have occurred decreases. The horizontal dashed line indicates the traditional \( P = .05 \) cutoff value. As \( t \) was varied from 0 to 40, the branch below \( t \) was varied from 40 to 0 such that the sum of these two length was always 40. For each \( t \) chosen, the corresponding maximum-likelihood value of \( \alpha \), the probability of a transition to cooperative breeding, was calculated by reiteration as described (Harvey and Pagel 1991, pp. 101–103). The method used to compute \( P(Y|X) \) is that of Maddison (1990), as formulated in a maximum-likelihood framework by Harvey and Pagel (1991). C, Effect of varying the probability of a transition to cooperative breeding, \( \alpha \), on the strength of correlations of cooperative breeding with an ecological variable. In this particular example, as the evolution of cooperative breeding becomes increasingly likely, the significance of correlations with ecological increases, although the specific relationship between \( P(Y|X) \) and \( \alpha \) is strongly dependent on the particular reconstruction of characters, as we have observed other types of relationships for other reconstructions (e.g., positive or “humped” curves). Branch lengths are as in A, dashed horizontal line as in B. The model used is again that of Harvey and Pagel (1991), except here arbitrary values of \( \alpha \) were chosen and applied to the reconstruction on the tree.
deemed an adaptation to a hypothesized factor because it displays a few instances of appropriate relationship to that factor, nonetheless, inferring a causal relationship from these associations would not be supported by the criteria of existing parsimony tests” (Edwards and Naeem 1993, p. 758; italics added). We do not imply that more complex models would enhance associations in our study; the conservatism of both CB and ecological traits on many of the trees ensures that the probabilities of change and time spans required to achieve significant associations would probably be unrealistic. In addition, in the case of robins, it would be difficult to interpret significance or nonsignificance by a method other than parsimony since the branch lengths in this case are unknown. But it is a striking result that several recent tests of correlated evolution of discrete behavioral and ecological characters have failed to detect statistically significant associations of traits that traditional analyses had suggested were highly correlated (see, e.g., Höglund 1989; Maddison 1990; Read and Nee 1991), either because the associations are indeed unsupported given the size of available data sets or because the models of character (e.g., parsimony) are unrealistic.

In practice, the results of parsimony and statistically consistent methods such as maximum likelihood converge when branch lengths as measured by expected amounts of character change are small but can conflict under other circumstances (Farris 1973; Felsenstein 1979, 1983; Goldman 1990). As a result, the statistical significance of a particular distribution of CB and ecological variables will depend in part on the model assumed (fig. 2; Harvey and Pagel 1991; Martins and Garland 1991). For example, we pointed out that both the frequency (about 3% among all birds) and complexity of CB suggest that it is much more difficult for lineages to gain CB than to lose it. This suspicion justifies the use of models able to detect and incorporate asymmetries in probability of change (Sanderson 1993). More realistic models for both discrete and continuous characters are being developed (see, e.g., Martins and Garland 1991; Janson 1992) and will be crucial to extracting more information from comparative data in conjunction with quantitative estimates of the phylogenies themselves.

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