Do Dietary Habits Predict Scale Counts in Snakes?

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Although studies on the biomechanical basis of feeding in snakes have revealed many links between dietary composition and feeding structures (e.g., Pough and Groves, 1983; Cundall, 1987), the strength of such links have rarely been examined quantitatively. In this paper, I explore a simple ecomorphological question: can the type of prey eaten by a snake species be used to predict two aspects of its feeding structures (the number of supralabial scales and dorsal scale rows)? The idea was first expressed more than 50 years ago by an innovative German scientist, Rudolf Mell (see Zhao and Adler, 1993). Based on his research in China, Mell noted that snake species eating larger prey (e.g., mammals rather than reptiles) had more supralabial scales and more midbody scale rows (Mell, 1929a,b). Mell suggested that an increased number of scale rows provides greater flexibility, enabling the skin between the scales to stretch over a large prey item. More recent authors have repeated this suggestion. For example, Gans (1974) interpreted the increased number of scale rows on the neck of egg-eating snakes in this way. Pough and Groves (1983) interpreted high supralabial and midbody scale-row counts in viperid snakes (compared to colubrids) in the same fashion. Mell's original formulation was simplistic, however, and recent analyses of snake feeding provide more sophisticated views on these topics. For example, Greene (1983, 1997) has pointed out that prey items can be "large" in a number of ways; Mell's hypothesis relies upon prey being large in diameter rather than (for example) mass. Equally, it is a gross oversimplification to rate all mammals that are consumed by terrestrial Australian snake species. Information on dietary composition and mean adult body sizes of snakes came from dissection and measurement of museum specimens (Shine, 1991, 1994). Diets were quantified in terms of the proportion of identifiable prey items falling into each major group (invertebrate, fish, amphibian, reptile, bird, mammal). Mean adult body size was calculated as the average of mean snout–vent lengths for adult males and adult females.

Three kinds of analysis were carried out on these data. First, I treated each species as a statistically independent unit and looked for the predicted correlations between the two types of scale characters (dorsal scale rows and supralabials) and between these traits and dietary composition (e.g., proportion of mammals in the diet). Second, because I found strong correlations between scale counts and body size (see below), I also compared dietary composition to size-corrected scale counts. To calculate size-corrected scores for each of the scale characters, I used residual scores from the general linear regressions of scale count versus mean body length. Thus, a species with a positive residual score for dorsal scale-rows would be one that had more scale-rows than would be expected for a species of that mean body size. Last, I combined the data with a putative phylogeny for these snakes (Fig. 1). This phylogenetic hypothesis was assembled by combining published suggestions about relationships among various subsets of the species involved (Kluge, 1991, 1993; Greer, 1997; Keogh et al., 2000; S. Keogh, pers. comm.). Hence, the resultant phylogeny (Fig. 1) is speculative and is not the result of any explicit phylogenetic analysis. Then I used this composite phylogeny to conduct a comparative analysis using independent contrasts (Purvis and Rambaut, 1995). This method overcomes the problem that many traits are highly conservative, so that correlations among traits may reflect phylogenetic inertia rather than functional relationship (Harvey and Pagel, 1991). Evolutionary changes in one variable (such as the proportional composition of the diet) within a clade can be compared to simultaneous changes in other traits (such as sculation) within the same clade. In the absence of reliable data on branch lengths within phylogenies, I assumed constant branch lengths (i.e., punctuated speciation events). Phylogenetically corrected relationships among variables were assessed by linear regression through the origin (Purvis and Rambaut, 1995). Statistical analyses were performed using CAIC (Purvis and Rambaut, 1995) and Statview 5 on an Apple Macintosh G4 computer. Prior to analysis, data were tested for conformity with assumptions of normality required for tests. A one-factor ANOVA on Pope's 1935 data (his table LX) with prey type as the factor and number of scale rows as the dependent variable, confirms Mell's (1929a,b) assertion that dietary habits correlate with scale counts within the Chinese snake fauna ($F_{1,51} = 3.88, P < 0.003$). Posthoc (Fishers' PLSD) tests show that species that eat mammals have significantly more
FIG. 1. Phylogenetic hypothesis for Australian snakes, as used in the present study. See text for list of sources for this phylogeny.

midbody scale rows than those that eat reptiles, frogs or invertebrates (P < 0.05 in each case).

Data were obtained for 108 Australian species, comprising 11 pythonids, nine colubrids, 76 elapids and 12 typhlopids. Mean adult snout–vent length ranged from 17–212 cm, midbody scale rows from 13–67, and supralabial scale counts from 4–14. The proportion of the diet composed of various prey types also varied widely (from 0–100% for each of fishes, reptiles, and mammals; from 0–97% for amphibians; 0–36% for birds).

Overall, my analyses for the Australian taxa revealed the same general pattern as noted by Mell for Chinese snakes but cast substantial doubt on the mechanism that he suggested to explain this pattern. Table 1 summarizes results from statistical tests of three predictions from Mell’s hypothesis.

Number of Midbody Dorsal Scale-Rows Is Positively Correlated with Number of Supralabial Scales.—As predicted, species with high numbers of midbody dorsal scale rows also had many supralabial scales (Fig. 2, Table 1). However, both of the scale counts also showed strong positive correlations with mean adult body size. These relationships differed among families. For example, at any given body size, pythons have more supralabial scales than elapids, and colubrids are intermediate in this respect (Fig. 3). The correlation between dorsal and labial counts remained high even after the effects of body size were removed (Table 1). That is, the trend for species with high numbers of scale-rows to also have many supralabial scales is not an indirect consequence of body size.

Comparative analysis further supported the notion of a functional relationship between scale numbers around the body versus along the upper lip. Increases in midbody scale rows during phylogeny were consistently associated with increases in supralabial counts (Table 1). After removing body-size effects from the comparative analysis, shifts in relative numbers of midbody dorsal scale-rows (i.e., residual scores) were associated with shifts in relative numbers of supralabial scales (i.e., residual scores; N = 70, r = 0.65, P < 0.0001).

Species that Eat Larger Prey Types Have More Dorsal Scale Rows.—As in the Chinese fauna, Australian snake species that feed on birds and mammals tend to have more dorsal scale rows than do other species (Table 1, Fig. 4). When the effects of body size on scale counts are removed, however, none of the probability values for these analyses attain the conventional level of significance (P < 0.05) after Bonferroni corrections are applied (Table 1).

Phylogenetically based analysis revealed a consistent association between dorsal scale rows and the proportion of mammalian prey in the diet but not for the other prey groups (Table 1). Using size-corrected scores (i.e., residual values from scale counts versus SVL) greatly weakened the apparent link between scatulation and diet: no dietary proportions were significantly correlated with size-corrected scale counts (P > 0.10 in all cases).

Species that Eat Larger Prey Types Have More Supralabial Scales.—Patterns for this variable are similar to those for dorsal scale rows (above), as might be expected from the high correlation between the two traits (Fig. 2). Species that feed on birds and mammals have more supralabial scales than do species that feed on ectotherms (Fig. 4), but this pattern disappears after body-size effects are removed from the analysis (Table 1). Phylogenetic shifts in supralabial counts were correlated (albeit weakly) with shifts in the proportion of mammalian prey (Table 1), but this effect disappeared when size-corrected values were used in the analysis (all P > 0.10).
The end result of these analyses is that three traits (body size, dietary habits, and scale counts) are significantly intercorrelated. Although it is difficult to distinguish direct from indirect effects within such constellations of traits, partial correlation offers a partial solution. I used this technique to evaluate the correlation between dietary habits and one variable (such as size) when another factor (such as body size) was held constant. Two sets of these analyses were conducted, one on the raw data and one on phylogenetic contrasts. In both cases, partial-correlation analyses suggested that dietary habits were more highly associated with body-size than with scale counts. For the raw data, the proportion of the diet composed of mammals was significantly correlated with mean adult SVL when either scale-row numbers or supralabial counts were held constant ($N = 108, r = 0.63, 0.71$, respectively, $P < 0.001$ in both cases) but neither of the scale characters was correlated with dietary habits if SVL was held constant ($N = 108, r = -0.03, 0.05$, respectively, $P > 0.50$). No other partial correlations between dietary habits and morphological traits were statistically significant ($P < 0.05$) after Bonferroni correction for multiple tests. The comparative analysis (independent contrasts) provided an almost identical result. The proportion of mammals in the diet was correlated with mean adult SVL when either of the scale characters was held constant ($N = 70, r = 0.56, 0.55$, respectively, $P < 0.001$ in both cases) but not with either of the scale counts when SVL was held constant ($N = 108, r = -0.13, -0.04$, respectively, $P > 0.50$). These results suggest that the correlations between scale counts and diet are secondary consequences of body-size effects on both scalation and diet.

If indeed we could find strong links between ecological traits (such as dietary composition) and morphological features (such as the numbers of dorsal scale-rows), such associations would be of considerable value. For example, we have quantitative information on dietary composition for only a small proportion of snake species. Unfortunately, my study is not encouraging with respect to inferring diet from scalation (or vice versa). There are at least three processes that might generate a correlation between dietary composition and scale counts in snakes. First, interspecific divergences in dietary habits (especially, relative prey size) might have imposed selection on scale-rows as envisaged by Mell. Second, the

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**Table 1. Relationships among scale characters and dietary composition in Australian snakes.** The table provides results from linear regression analysis of three types of data. First, values for each species were used as independent datapoints (“raw data”). Then, because both types of scale counts were highly correlated with overall body size, the effects of size were removed by calculating residual scores from the general linear regressions of the scale trait against mean adult snout–vent length (“size-corrected data”). Third, comparative analysis using independent contrasts was used to remove the effects of phylogenetic conservatism. Sample size was 108 for the first two types of tests and 70 for the comparative analysis. The table shows the correlation coefficient ($r$) and its associated probability ($P$). **Boldface** shows results that are statistically significant ($P < 0.05$) after Bonferroni correction.

<table>
<thead>
<tr>
<th>Traits being compared</th>
<th>Raw data</th>
<th>Size-corrected data</th>
<th>Comparative analysis</th>
</tr>
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<tr>
<td></td>
<td>$r$</td>
<td>$P$</td>
<td>$r$</td>
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<td>Dorsal scale rows vs supralabial scales</td>
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</tr>
</tbody>
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**Fig. 2.** The relationship between the numbers of dorsal scale rows and supralabial scale count in an interspecific comparison among Australian snakes. Each point represents one species. See Table 1 for results of a test of the statistical significance of this relationship.
Fig. 3. The relationship between body size (mean adult snout–vent length) and scale counts in Australian snakes. Lower graph shows data for the number of dorsal scale rows, and upper graph shows the data for the number of supralabial scales. Each point represents one species. Linear regression for these datasets shows: dorsal scale rows, $N = 108$, $r = 0.65$, $P < 0.0001$; supralabial scales, $N = 108$, $r = 0.62$, $P < 0.0001$.

correlation between scale counts and dietary composition might be an artifact of phylogenetic inertia, because some lineages of snakes have low values for all three traits (supralabials, dorsal scale rows, and proportion of the diet composed of endotherms) whereas other lineages have high values for all three traits. This circumstance might have arisen early in snake phylogeny, and require no adaptive explanation in terms of selective forces during the radiation of species within lineages. In practice, phylogenetic shifts in the degree of reliance on mammalian prey were consistently accompanied by shifts in scale counts (Table 1), suggesting a functional association between the traits. Nonetheless, phylogenetic conservatism undoubtedly amplifies the strength of these correlations (Table 1, Fig. 4). Last, correlation does not imply causation. A third factor (mean adult body size) is known to correlate both with dietary habits (e.g., Fitch, 1960; Arnold, 1993; Shine, 1994) and with scale counts (e.g., Klauber, 1956; Lindell, 1994). Thus, a shift in food habits toward larger prey could favor the evolution of larger body size, which in turn could favor an increase in scale counts (because larger snakes have more scales: see Fig. 3). In keeping with this hypothesis, correlations between scale counts and dietary habits generally disappeared when body-size effects were removed from the analysis (Table 1).

In summary, Mell was probably correct in recognizing a general association between scale counts and diets in snakes, but wrong about its cause. To fully understand the pattern identified by Mell 50 years ago, we will need to gather data on traits that relate much more closely to the actual selective pressures and functional challenges involved in prey ingestion. For example, many researchers in snake ecology still report only the species and mass of prey items, despite heartfelt appeals to take more extensive data (e.g., Greene 1997). Maximum prey diameter is simple to measure (even with partially digested prey) and is likely to be a better measure of the physical difficulty of ingesting the item than is prey mass. Until we have extensive datasets on such variables, many of the ideas espoused by visionaries like Rudolf Mell will remain untested.

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Testing for Equal Catchability of Triturus Nests by Dip Netting

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Testing the assumption of equal catchability is fundamental to understanding amphibian populations, modeling population dynamics, and testing hypotheses concerning life-history traits. Unequal catchability forms an important source of potential error in quantitative surveying, affecting the estimation of population parameters such as population size, in particular when the capture-recapture method is used (Bohlin and Sundström, 1977; Carothers, 1979; Pollock et al., 1990). Nonrepresentative sampling has been reported for many animal species in a wide range of taxa, and it has been questioned whether populations consisting of animals with equal catchability actually exist (Carothers, 1973). In amphibian studies, sampling bias has been mostly ignored, with some notable exceptions (Gelder and Rijsdijk, 1987; Shirose and Brooks, 1995; Wood et al., 1998; Wilson and Pearman, 2000).

The techniques available for the study of amphibians include some sophisticated approaches (Heyer et al., 1990; Mølle and Kupfer, 1998). However, the sampling of many different habitats will be facilitated by the use of simple, small, and inexpensive tools, such as the traditional dip net. Amphibians that breed over long periods in lentic water and those with larval stag-

es of long duration are especially amenable to netting. Paleartic nests of the genus Triturus fall into this cat-

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