

# Tempo of trophic evolution and its impact on mammalian diversification

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**Mammals are characterized by the complex adaptations of their dentition, which are an indication that diet has played a critical role in their evolutionary history. Although much attention has focused on diet and the adaptations of specific taxa, the role of diet in large-scale diversification patterns remains unresolved. Contradictory hypotheses have been proposed, making prediction of the expected relationship difficult. We show that net diversification rate (the cumulative effect of speciation and extinction), differs significantly among living mammals, depending upon trophic strategy. Herbivores diversify fastest, carnivores are intermediate, and omnivores are slowest. The tempo of transitions between the trophic strategies is also highly biased: the fastest rates occur into omnivory from herbivory and carnivory and the lowest transition rates are between herbivory and carnivory. Extant herbivore and carnivore diversity arose primarily through diversification within lineages, whereas omnivore diversity evolved by transitions into the strategy. The ability to specialize and subdivide the trophic niche allowed herbivores and carnivores to evolve greater diversity than omnivores.**

macroevolution | ecological specialization | character evolution

Living mammals are remarkably diverse: they span eight orders of magnitude in mass, occupy a variety of habitats across the globe, and exploit subterranean, aquatic, terrestrial, arboreal, and aerial niches. Living mammals also show striking differences in diversity between lineages of similar age, from the more than 2,200 species of rodent to the single species of armadillo (1, 2). Early mammals were small, homeothermic endotherms with tribosphenic molars. Homeothermic endothermy enabled mammals to survive in a wider range of ambient temperatures and achieve higher sustained activity levels, but it also increased energy demands (3). These increased energetic demands necessitated adaptations or behaviors that either allowed more efficient extraction of energy from the food consumed, entailed consumption of more energy rich foods, or required an increase in the time spent foraging and eating. The tribosphenic molar, which combines shearing and crushing functions in the precisely occluding teeth, is considered to be a key innovation that promoted more effective carnivory and omnivory in early mammalian lineages (4). This type of tooth is also frequently cited as facilitating the diversification of therian mammals (4–6). The tribosphenic molar is an evolutionarily and functionally highly versatile structure (4, 7) that, in combination with heterodonty (different tooth types within the jaw), enabled mammals to evolve a disparate array of specialized dentitions and thus adapt to a broad variety of niches. Indeed, the extraordinary dental diversity of mammals—to the extent that many species can be identified by the morphology of their molars alone (8)—is a testament to the importance of diet to mammalian evolution.

Although the adaptations of individual mammalian lineages to diet have been well studied, few studies have examined the impact of diet on large-scale macroevolutionary patterns. Here we present a quantitative macroevolutionary analysis of the tempos of lineage diversification and trophic transition across living mammals. Although the evolutionary history of mammals is intimately

linked to diet, no consensus exists on how trophic strategy (i.e., herbivory, carnivory, or omnivory) may bias transition rates into these different strategies and impact speciation and extinction. Many hypotheses have focused on specific trophic strategies. For example, theoretical, paleontological, and comparative analyses all suggest that carnivores are more prone to extinction (9, 10) resulting from their dependence on less abundant and stable food resources. It has also been common to make macroevolutionary predictions by considering omnivores, which use both plant and animal protein, as less specialized than species that use a narrower subset of the available food resources (i.e., herbivores or carnivores) (11, 12). Hypotheses linking greater diversification rates to ecological specialization have a long history dating back to Darwin (13) and they remain widespread (12, 14–16). Early macroevolutionary theorists suggested that diversification proceeds from generalist ancestors to specialist descendants (15, 17), implicitly predicting that diversification rates should be highest in specialist groups and transition rates should be highest out of omnivory into herbivory and carnivory.

Drawing from published data, we have compiled a dataset of diets for over one-third of mammalian species (Dryad repository, <http://dx.doi.org/10.5061/dryad.vr28vf67>). We applied these data to a virtually complete phylogeny of living mammals (5,020 species from ref. 1, based on ref. 18) and analyzed them using phylogenetic comparative methods that simultaneously estimate diversification and transition rates (19, 20).

## Results and Discussion

We compiled detailed dietary data on primary observations of mammalian diets from the scientific literature and then, using uniform criteria, categorized each species as either a herbivore (a specialist on primary producers), a carnivore (a specialist on consumers), or an omnivore (a generalist, eating both plant and animal material). Using these data and a virtually complete species-level time-calibrated phylogeny of living mammals (ref. 1, based on ref. 18) we can confirm that mammalian trophic strategies are not evenly distributed among mammalian taxa (Fig. 1). Some clades are almost exclusively carnivorous (e.g., aquatic Cetartiodactyla, such as whales and dolphins; aquatic Carnivora, such as seals and walrus) or herbivorous (e.g., terrestrial Cetartiodactyla, such as deer, cows, and antelope) and others appear to switch frequently between omnivory and herbivory (e.g., primates) or omnivory and carnivory (such terrestrial Carnivora as bears, dogs, and foxes). We used this variability

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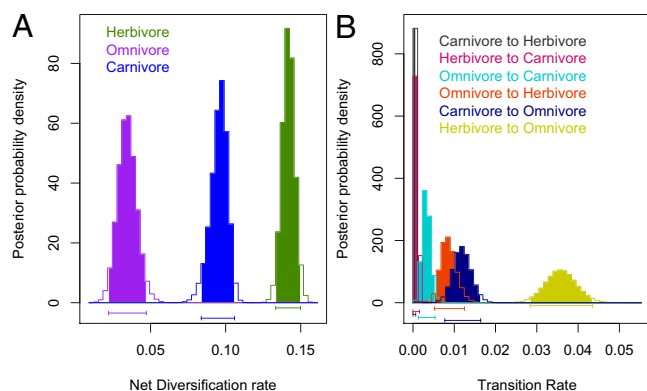
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**Fig. 2.** Plot of the posterior probability density of the parameter estimates of the all rates-free model (which was the best-fitting model from the maximum-likelihood analysis) analyzed using Bayesian MCMC methods on 10 of the 100 replicate phylogenies. (A) Depiction of the net diversification rate (speciation minus extinction) for all three trophic strategies and (B) the rate of transition between the trophic strategies.

Moreover, the tempo of transitions between the trophic strategies is highly biased. Surprisingly, the highest rates occur into omnivory from herbivory and carnivory and the lowest rates with almost zero transitions occur between herbivory and carnivory (Fig. 2B). These results are supported by the maximum-likelihood parameter estimates from the top four best-fitting models which range in  $\Delta AIC$  from 19.9 to 48.9 (Table 1); net diversification rate is the highest in herbivores, intermediate in carnivores, and lowest in omnivores; the highest transition rate is into omnivory. Thus, omnivore diversity evolves primarily through transitions into that strategy and rarely by diversification within omnivorous lineages, whereas herbivore and carnivore diversity is chiefly produced through diversification.

These results could have been produced by more than one scenario of evolutionary dynamics. Many discussions of diversification have made the distinction between explanations and models that assume differing intrinsic rates of extinction or speciation, on one hand, and differing ecological limits on the number of species the environment can accommodate (highest “carrying capacity”; according to ref. 24) on the other (24–28). If there is a global limit on the number of species that can be supported within a particular dietary strategy, net diversification may be diversity-dependent whereby rates slow through time as niche space fills up until diversity is at equilibrium (e.g., ref. 29). Indeed, fossil and molecular data frequently identify a slow-down in diversification rate (e.g., refs. 30 and 31), although there are other explanations for an apparent slow-down that do not involve niche-filling (e.g., ref. 32). Under this scenario the trophic group with a lower limit on its diversity will appear over time to have a lower average rate of net diversification than a group that continues on to reach a higher limit, even though they have grown with identical diversification rates (24). The constant-rate models we have used do not allow us to distinguish this situation from one in which the two groups have had consistently different rates. Nonetheless, whether the trophic strategies inferred to have higher rates have diversified at a consistently more rapid pace, or have simply spent more of their evolutionary history diversifying at a high rate, similar biological interpretations can be applied to these two alternative mechanisms.

There may be good biological reasons to expect herbivorous, carnivorous, and omnivorous niches to vary in the diversity they can support. In any ecosystem mammalian primary consumers have access to the largest resource base because primary producers account for the largest proportion of biomass. It is therefore possible

that, along with a larger number of individuals, a greater number of herbivore species can be supported globally. Additionally, the stability of plant-based food resources may facilitate diversification, whereas species dependent on food at higher trophic levels may be less able to subdivide niches in a stable fashion. There may also be strong limits on the number of omnivores that can be sustained because some food-web models suggest omnivory should be rare because it destabilizes food webs (e.g., refs. 33 and 34).

However, it is unlikely that the low net diversification rate in omnivores is entirely explained by the omnivorous niche supporting fewer species. In our dataset omnivores are not rare: species richness of carnivores and omnivores is similar (463 and 413 respectively), a pattern that is also evident in natural communities of mammals (35). Low diversification rates within omnivores may be caused by lower speciation rates if dietary specialists are more likely to undergo allopatric speciation than omnivores because specialists’ narrower food tolerances allow their geographic distributions to be more readily fragmented (36–38). Additionally, specialization can open opportunities for further specialization (39–41), thereby generating diversity through niche subdivision. Extreme illustrations of this process have been presented in several studies on phytophagous insects, which have documented higher rates of diversification in these plant-feeding groups (15). Niche subdivision, by definition, tends to increase specialization, so for omnivores (as generalists) the opportunities for producing descendant lineages that retain their dietary breadth might be limited. Accordingly, omnivores should diversify more slowly than carnivores or herbivores. The difference between herbivores and carnivores may be explained by higher rates of extinction in carnivores because carnivores are more vulnerable to disturbance of species lower down the food chain (9, 42). Indeed, higher trophic level has been shown to be correlated with higher extinction risk in extant mammals (10).

Ecologically, omnivory can be a strategy for surviving variability in resource availability (34) and, if lineages shift to omnivory during times of environmental perturbation, this may explain why diversification is low (43) and transition rates are highest into, not out of, omnivory. Taken together, these results suggest that omnivory acts as an evolutionary “sink,” which at the broadest scale contradicts the expectation that ecological generalists are sources of future diversity and ecological specialists are evolutionary dead-ends (16, 44). Nonetheless, omnivory does appear to play an important macroevolutionary role because virtually all trophic transitions within mammals involve omnivory: the estimated transition rate between herbivory and carnivory is approximately zero. This finding is perhaps not surprising: In mammals, herbivory and carnivory often entail different and antagonistic physiological and morphological adaptations. Such changes would likely require large amounts of evolutionary time, increasing the probability of a branching event and the evolution of new lineages with intermediate ecology during the transition. Many omnivorous mammals exhibit intermediate phenotypes: the teeth of omnivores, like carnivores, generally have distinct cusps and low crowns; yet like herbivores, their enamel is thickened or crenulate (15, 17). Omnivores’ intestines are similarly intermediate, with gut lengths intermediate between herbivores and carnivores (45).

Our results imply that diet is a key factor in determining the rates at which different mammalian groups have diversified; however, the link between trophic strategy and diversification could be indirect and involve other important biological or ecological traits. Among the many factors that can drive patterns of diversification, body size and population density are often proposed as strong candidates (46). For body size, the typical expectation is for speciation rates to be higher in smaller species (e.g., refs. 47 and 48) and extinction higher in larger species (49). However, recent analysis of fossils found higher rates of both origination and extinction in larger mammals (10, 50); net diversification rates were either the same in small and large

mammals or higher in small species, depending on the data subset. Regardless, size is unlikely to be driving our patterns, as herbivorous mammals tend to be larger than carnivorous and omnivorous species (51) and on this basis should exhibit the lowest rates of diversification. Predictions based on population density are similarly problematic: species relying on resources at higher trophic levels usually live at lower densities (47) and should therefore experience much higher extinction rates (52); omnivores should have intermediate diversification rates. In our results this pattern is borne out in the difference between herbivores and carnivores but omnivores have the lowest diversification rates, contrary to the population density predictions.

## Conclusion

Although the determinants of mammalian diversity are undoubtedly complex (53, 54), diet has had profound consequences for mammalian macroevolution. Through trophic strategy the major features that characterize mammals, such as homeothermy and specialized dentition, are connected to the controls on diversification: herbivores diversify fastest, carnivores are intermediate, and omnivores are slowest. The estimated transition-rate bias between the trophic strategies contradicts some common expectations: omnivory appears to be an evolutionary sink rather than a source of diversity, and herbivory and carnivory are the basis of future diversity, not evolutionary dead-ends. The uneven distribution of lineages within mammals is thereby linked to differences in trophic strategy, yielding a conclusion that has broad implications for understanding mammalian evolution. The model we provide can be integrated with data from the fossil record and tested further using the increasingly sophisticated methods currently being developed for estimating speciation and extinction rates from molecular phylogenies.

## Methods

**Data Collection and Dietary Categorization.** We constructed a database of diets of mammalian species from published accounts of primary research reporting data obtained through analysis of stomach or cheek-pouch contents or the contents of food stores, direct behavioral observation, or fecal analysis. We recorded complete descriptions of diet from the sources; these descriptions were then converted to discrete character codings for the presence or absence of four food types in the diet: invertebrate protein, vertebrate protein, fibrous plant parts (mature leaves, stems, wood, and bark), and nonfibrous plant parts (any other parts of plants, along with fungi and lichens). Applying uniform and explicit criteria, we converted quantitative and qualitative descriptions of diet for each species to a repeatable coding of trophic strategies in three categories: carnivore, omnivore, and herbivore. This process yielded high-quality diet codings for 1,534 species of mammals. Of these 658 were herbivores, 463 carnivores, and 413 omnivores (see *SI Methods* and <http://dx.doi.org/10.5061/dryad.vr28vf67> for complete dataset and references used to generate it).

**Models of Trophic Transitions and Diversification Rate.** We used a model-testing approach to investigate the tempo of trophic evolution and its impact on diversification rate. We set up eight models that tested whether speciation, extinction, or transition rates were independent of trophic strategy, constrained by trophic strategy, or constrained by being a generalist (omnivore) or specialist (carnivore or herbivore). The most complex model had separate rates of speciation and extinction for each of the three trophic categories (six parameters) along with unconstrained rates of transition between each pair of trophic strategies (six parameters): herbivore to carnivore, carnivore to herbivore, herbivore to omnivore, omnivore to herbivore, omnivore to carnivore, and carnivore to herbivore.

**Phylogenetic Comparative Methods.** We combined the diet data with a time-calibrated phylogeny of virtually all living mammalian species (49) to estimate which of the eight evolutionary models was the best fit to our data. We used a version of the Fritz et al. tree (ref. 1, based on ref. 18), which resolved all polytomies by assigning branch lengths by the birth-death algorithm (54). Maximum-likelihood results were summarized across 100 such trees and the Bayesian MCMC results across 10 trees (because of constraints in time and CPU availability), including the two most disparate topologies according to the Robinson-Foulds symmetric distance metric.

We conducted a simultaneous analysis of the tempo of trophic evolution and its impact on diversification rate because these two processes are not independent (54). Analyses were conducted using the "Multiple State Speciation Extinction" (MuSSE) model in the diversitree package (55) within the statistical software R (56). This method is an extension of the "Binary State Speciation Extinction" (BiSSE) methods described in ref. 20. The fit of the eight models across the 100 trees was calculated using maximum likelihood; a full Bayesian analysis required more computational power or time than we had access to (~800 CPU's for ~25 d). We assessed model-fit using the AIC, which takes into account the number of parameters used in the model as well as the goodness of fit to the data. Akaike weights were then calculated, which indicate the probability that the model is the best among our set of eight candidate models. To examine the uncertainty in the parameter estimates, the best-fitting model from the maximum-likelihood analysis was analyzed using Bayesian MCMC methods. We assessed model adequacy of the best-fitting model through posterior predictive simulation using the parameter estimates from the MCMC analysis. We then compared the posterior predictive distribution of the number of herbivorous, carnivorous, and omnivorous mammals to our dataset (Fig. S1). Further methodological details are provided in *SI Methods*.

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- Fritz SA, Bininda-Emonds ORP, Purvis A (2009) Geographical variation in predictors of mammalian extinction risk: Big is bad, but only in the tropics. *Ecol Lett* 12:538–549.
- Wilson DE, Reeder DM (2005) *Mammal Species of the World. A Taxonomic and Geographic Reference* (Johns Hopkins Univ Press, Baltimore, MD), 3rd Ed, p 2142.
- Crompton AW, Taylor CR, Jagger JA (1978) Evolution of homeothermy in mammals. *Nature* 272:333–336.
- Luo Z-X (2007) Transformation and diversification in early mammal evolution. *Nature* 450:1011–1019.
- Jacobs LL, Winkler DA, Murry PA (1989) Modern mammal origins: Evolutionary grades in the Early Cretaceous of North America. *Proc Natl Acad Sci USA* 86:4992–4995.
- Woodburne MO, Rich TH, Springer MS (2003) The evolution of tribospheny and the antiquity of mammalian clades. *Mol Phylogenet Evol* 28:360–385.
- Simpson GG (1936) Studies of the earliest mammalian dentitions. *The Dental Cosmos* 78:791–800.
- Carroll RL (1988) *Vertebrate Paleontology and Evolution* (W. H. Freeman, New York).
- Diamond JM (1984) 'Normal' extinctions of isolated populations. *Extinctions*, ed Nitecki MH (Chicago Univ Press, Chicago), pp 191–246.
- Purvis A, Gittleman JL, Cowlishaw G, Mace GM (2000) Predicting extinction risk in declining species. *Proc Biol Sci* 267:1947–1952.
- Colles A, Liow LH, Prinzing A (2009) Are specialists at risk under environmental change? Neocological, paleoecological and phylogenetic approaches. *Ecol Lett* 12: 849–863.
- Futuyma DJ, Moreno G (1988) The evolution of ecological specialization. *Annu Rev Ecol Syst* 19:207–233.
- Darwin C (1859) *On the Origin of Species by Means of Natural Selection, or the Preservation of Favoured Races in the Struggle for Life* (J. Murray, London).
- Vrba ES (1995) On the connections between paleoclimate and evolution. *Paleoclimate and Evolution, with Emphasis on Human Origins*, eds Vrba ES, Denton GH, Partridge TC, Burckle LH (Yale Univ Press, New Haven, CT), pp 24–45.
- Simpson GG (1953) *The Major Features of Evolution* (Columbia Univ Press, New York).
- Eldredge N (1982) Phenomenological levels and evolutionary rates. *Syst Zool* 31: 338–347.
- Cope ED (1896) *The Primary Factors of Organic Evolution* (Open Court Publishing, Chicago, IL).
- Bininda-Emonds ORP, et al. (2007) The delayed rise of present-day mammals. *Nature* 446:507–512.
- FitzJohn RG (2010) Diversitree: Comparative phylogenetic tests of diversification, version 0.7-2. Available at <http://cran.r-project.org/web/packages/diversitree>.
- Maddison WP, Midford PE, Otto SP (2007) Estimating a binary character's effect on speciation and extinction. *Syst Biol* 56:701–710.
- FitzJohn RG, Maddison WP, Otto SP (2009) Estimating trait-dependent speciation and extinction rates from incompletely resolved phylogenies. *Syst Biol* 58:595–611.
- Rabosky DL (2010) Extinction rates should not be estimated from molecular phylogenies. *Evolution* 64:1816–1824.

23. Burnham KP, Anderson DR (2002) *Model Selection and Multimodel Inference: A Practical Information Theoretic Approach* (Springer, New York), 2nd Ed.
24. Rabosky DL (2009) Ecological limits and diversification rate: Alternative paradigms to explain the variation in species richness among clades and regions. *Ecol Lett* 12: 735–743.
25. Rabosky DL, Glor RE (2010) Equilibrium speciation dynamics in a model adaptive radiation of island lizards. *Proc Natl Acad Sci USA* 107:22178–22183.
26. Ricklefs RE (2007) Estimating diversification rates from phylogenetic information. *Trends Ecol Evol* 22:601–610.
27. Raup DM (1976) Species diversity in the Phanerozoic: An interpretation. *Paleobiology* 2:289–297.
28. Raup DM, Gould SJ, Schopf TJM, Simberloff DS (1973) Stochastic models of phylogeny and the evolution of diversity. *J Geol* 81:525–542.
29. Rabosky DL, Lovette IJ (2008) Explosive evolutionary radiations: Decreasing speciation or increasing extinction through time? *Evolution* 62:1866–1875.
30. Alroy J (1996) Constant extinction, constrained diversification, and uncoordinated stasis in North American mammals. *Palaeogeogr Palaeoclimatol Palaeoecol* 127: 285–311.
31. Phillimore AB, Price TD (2008) Density-dependent cladogenesis in birds. *PLoS Biol* 6: e71.
32. Etienne RS, Rosindell J (2012) Prolonging the past counteracts the pull of the present: Protracted speciation can explain observed slowdowns in diversification. *Syst Biol* 61: 204–213.
33. Pimm SL, Lawton JH (1978) On feeding on more than one trophic level. *Nature* 275: 542–544.
34. Ingram T, Harmon LJ, Shurin JB (2009) Niche evolution, trophic structure, and species turnover in model food webs. *Am Nat* 174:56–67.
35. Kelt DA, et al. (1996) Community structure of desert small mammals: Comparisons across four continents. *Ecology* 77:746–761.
36. Leschen RAB, Buckley TR (2007) Multistate characters and diet shifts: Evolution of Erotylidae (Coleoptera). *Syst Biol* 56:97–112.
37. Hunt T, et al. (2007) A comprehensive phylogeny of beetles reveals the evolutionary origins of a superradiation. *Science* 318:1913–1916.
38. Farrell BD (1998) “Inordinate Fondness” explained: Why are there So many beetles? *Science* 281:555–559.
39. Coll M, Guershon M (2002) Omnivory in terrestrial arthropods: Mixing plant and prey diets. *Annu Rev Entomol* 47:267–297.
40. Hamilton WJ III, Buskirk RE, Buskirk WH (1978) Omnivory and utilization of food resources by *Chacma* baboons, *Papio ursinus*. *Am Nat* 112:911–924.
41. Van Valen L (1965) Morphological variation and width of ecological niche. *Am Nat* 99: 377–390.
42. Van Valkenburgh B, Wang X, Damuth J (2004) Cope’s rule, hypercarnivory, and extinction in North American canids. *Science* 306:101–104.
43. Vrba ES (1980) Evolution, species and fossils: How does life evolve? *S Afr J Sci* 76: 61–84.
44. Vrba ES (1993) Turnover-pulses, the red queen, and related topics. *Am J Sci* 293-A: 418–452.
45. Hillson S (2005) *Teeth* (Cambridge Univ Press, Cambridge) 2nd Ed.
46. Chivers DJ, Langer P (1994) Gut form and function: Variations and terminology. *The Digestive System in Mammals: Food, Form and Function*, eds Chivers DJ, Langer P (Cambridge Univ Press, New York), pp 3–8.
47. Case TJ (1979) Optimal body size and an animal’s diet. *Acta Biotheor* 28:54–69.
48. Eldredge N (1979) Alternative approaches to evolutionary theory. *Bulletin of Carnegie Museum of Natural History* 13:7–19.
49. Hutchinson GE, MacArthur RH (1959) A theoretical ecological model of size distributions among species of animals. *Am Nat* 93:117–125.
50. McKinney ML (1997) Extinction vulnerability and selectivity: Combining ecological and paleontological views. *Annu Rev Ecol Syst* 28:495–516.
51. Liow LH, et al. (2008) Higher origination and extinction rates in larger mammals. *Proc Natl Acad Sci USA* 105:6097–6102.
52. Fa JE, Purvis A (1997) Body size, diet and population density in Afrotropical forest mammals: A comparison with neotropical species. *J Anim Ecol* 66:98–112.
53. Schmidt-Nielsen K (1984) *Scaling: Why Is Animal Size So Important?* (Cambridge Univ Press, New York).
54. Kuhn TS, Mooers AO, Thomas GH (2011) A simple polytomy resolver for dated phylogenies. *Methods in Ecology and Evolution* 2:427–436.
55. Maddison WP (2006) Confounding asymmetries in evolutionary diversification and character change. *Evolution* 60:1743–1746.
56. R Development Core Team (2011) R: A language and environment for statistical computing. (R Foundation for Statistical Computing, Vienna, Austria) <http://www.R-project.org/>.

# Supporting Information

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## SI Methods

**Dietary Categorization.** A database of diets of mammalian species was constructed from published accounts drawn from primary research with data obtained through analysis of stomach or cheek-pouch contents or the contents of food stores, direct behavioral observation, or fecal analysis. Only observations of noncaptive animals that were presented at the species level were included. Dietary inferences based on morphology, phylogeny, or other indirect means were not used. During primary data collection, we recorded complete descriptions of diet from the sources; these descriptions were then converted to discrete character codings for the presence or absence of four food types in the diet: invertebrate protein, vertebrate protein, fibrous plant parts (mature leaves, stems, wood, and bark), and nonfibrous plant parts (any other parts of plants). A category was scored as present in the diet of a species if it constituted at least 5% of the food consumed by volume, weight, or feeding time. In many cases, the data available for diet were not quantitative, but relative amounts of various dietary elements could be deduced from verbal descriptions. In these instances, a food was scored as present in the diet if the descriptors “sometimes,” “some,” “supplemented by,” or “small amounts” were used. Foods that were consumed “occasionally,” “periodically,” “probably,” or “when no other food is available” were considered to constitute less than 5% of the diet and were therefore coded as absent. The four food-type categories were used to determine whether a species was herbivorous (only plant parts present in the diet), carnivorous (only invertebrate or vertebrate protein in the diet), or omnivorous (including both plant parts and animal protein in the diet).

Efforts in our literature searches emphasized broad taxonomic/phylogenetic representation; thus, for example, in the course of the searches more effort would be directed at finding information for species in an unrepresented family than on obtaining exhaustive sampling for any given genus. As a result, the dataset is as unbiased as possible given the biases to collection of dietary observations. To the degree we can assess it, our sampling should not bias our results. Well-studied clades for diet data are often also those that have good phylogenetic data, and those are distributed fairly evenly across the dietary categories. Compilation of the database was distributed by taxonomic group among the four authors. To maintain consistency and provide cross-validation of the scoring, all four authors independently scored subsets of each of our compilations and explicit guidelines were developed for handling accounts that were descriptive but not quantitative. A few species (23 species) were described in our references on the basis of direct observations as omnivorous, carnivorous, or herbivorous; these categorizations were used where no detailed dietary data were available if descriptions were based on direct observations.

Although the coding into only three categories is coarse, it has advantages both for robustness of our results and for analytical tractability. Because the types of available diet data are so diverse, coding beyond the three simple categories would reduce the number of species that could be included in the analysis. This process would create more problems with missing data and biased data toward well-studied groups. Additionally, adding more categories to the analysis would not only reduce the number of datapoints on which character state reconstructions depend, but would add exponentially more complexity to the computation of models of diversification and character change. Given that the model we used here is already demanding of time and processor capability, adding further complexity would likely yield data that

could not be analyzed using these methods. During preliminary analyses we ran one model that used four instead of three categories, splitting herbivores into folivores and frugivores, and the likelihood did not converge.

Our process yielded high-quality diet codings for 1,530 species of mammals. Of these 656 were herbivores, 461 carnivores, and 413 omnivores. The resulting database (<http://dx.doi.org/10.5061/dryad.vr28vf67>) is deposited in Dryad ([www.datadryad.org](http://www.datadryad.org)).

**Phylogeny.** Diet data were combined with the time-calibrated mammalian phylogeny of Fritz et al. (1, 2) to analyze patterns of diversification with diet in mammals. Diversification patterns were analyzed over the whole tree (5,020 species), even though dietary data are available only for a subset of them, using the method developed by Fitzjohn et al. for incomplete character state knowledge on a complete tree (3). This practice avoids the problems in distributions of branch lengths created by cropping out species for which we lack diet data.

The original tree (1) was terminally unresolved. To overcome this, we used a version of this tree in which the unresolved nodes were arbitrarily resolved and assigned branch length by the birth-death algorithm of Kuhn et al. (4). This method was chosen in preference to the methods given in ref. 3, as unresolved clades slow down the analysis considerably. To avoid effects of this arbitrary resolution on our result, we used 100 trees representing random replicates of this resolution process in the maximum-likelihood analyses and 10 trees (including the two most disparate according to the Robinson-Foulds symmetric distance metric) for the Bayesian Markov Chain Monte Carlo (MCMC) analyses. The results we report summarize the findings that were consistent across all tree replicates.

**Data Analysis.** Analyses were conducted using the “Multiple State Speciation Extinction” (MuSSE) model in the diversitree package (5) within the statistical software R. This method is an extension of the “Binary State Speciation Extinction” (BiSSE) methods described in ref. 6. The fit of the eight evolutionary models across the 100 trees was calculated using maximum likelihood, as a full Bayesian analysis required more computational power or time than we had access to (~800 CPU’s for 25 d or ~8 CPU’s for 6+ y). Diversitree was used to make a likelihood function using the “make.musse” command, and then, using the starting point it generated using the “starting.point.musse” function, we optimized the function by maximum likelihood using “find.mle.” This analysis was repeated for each of the 100 replicate trees and run in batches of 10 to allow parallel processing.

For each tree, eight different models were run; these tested whether speciation, extinction, or transition rates were independent of trophic strategy, constrained by trophic strategy, or constrained by being a generalist (omnivore) or specialist (carnivore or herbivore). (i) The most complex model allowed speciation, extinction, and transition rates to vary independently in each of the three trophic categories (12 parameters). (ii) Constrained speciation allowed rates to be the same across trophic strategies and allowed extinction and transition rates to vary independently (10 parameters). (iii) Constrained extinction allowed rates to be the same across trophic strategies and allowed speciation and transition rates to vary independently (10 parameters). (iv) Constrained diversification allowed rates (speciation and extinction) to be the same across all trophic strategies with transition rates allowed to vary independently (8 parameters). (v) Constrained diversification allowed rates (speciation

and extinction) to be the same within specialists and generalists and allowed transition rates to vary independently (10 parameters). (vi) Constrained transition allowed rates to be the same between all trophic strategies and allowed speciation and extinction rates to vary independently in each of the three trophic categories (7 parameters). (vii) Constrained transition allowed rates to have three different rates: between specialist strategies, from specialist to generalist, and from generalist to specialist; speciation and extinction rates were free to vary independently in each of the three trophic categories (9 parameters). (viii) Constrained specialist and generalist allowed transition rates and diversification rates, combining the constraints of models 5 and 7 (7 parameters).

Because the search for a maximum-likelihood point can be affected by starting point, we also generated parameter estimates using starting points with substantially different (up to an order of magnitude greater or less than the) initial  $q$  (transition rate),  $\lambda$  (speciation rate), and  $\mu$  (mutation rate) values; none of these estimates generated a model with a likelihood value superior to the model produced by the initial analysis. MCMC analyses for a subset of the trees also did not produce any models with likelihoods superior to the best model generated by the initial analysis. This testing of assumptions in the diversitree model allowed us to be confident in our conclusion that the model fitting each of the three categories with a significantly different diversification rate is the best description of our data.

We have reported differences among models using the Akaike Information Criterion (AIC), which takes into account the number of parameters used in the model as well as the goodness of fit to the data. All of these comparisons were made for each of the 100 trees independently and reported as medians and interquartile ranges,

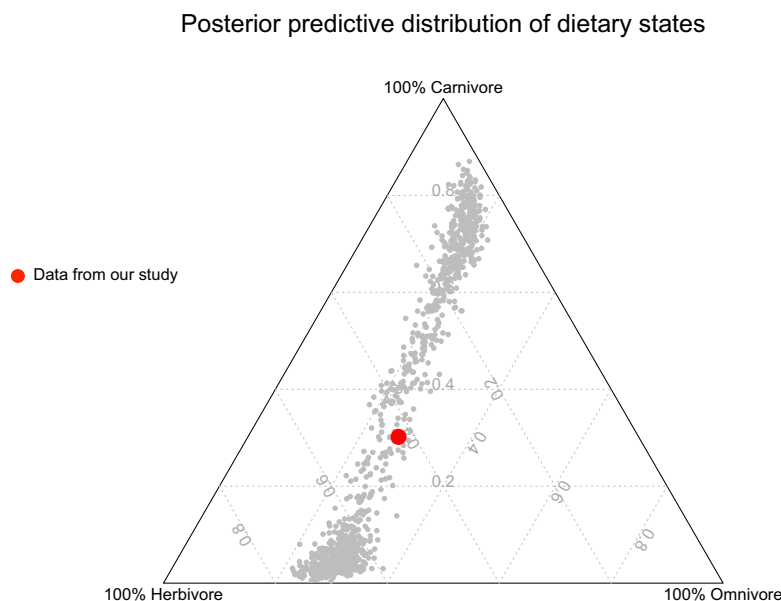
because mean values may not be meaningful for the analyses performed. All patterns we report hold for each of the 100 trees analyzed separately.

To examine the uncertainty in the parameter estimates, the best-fitting model from the maximum-likelihood analysis was analyzed using Bayesian MCMC methods. We used an exponential prior with rate  $1/(2 \times \text{character independent rate})$  and a step-size that was the range of the observed samples from a preliminary MCMC run of 100 steps. We ran a chain of 10,000 steps on 10 trees, which resolved the polytomies in the original phylogeny (1); each tree was run on a separate CPU and took ~25 d. The output for the 10 trees was combined after examination of the individual runs to estimate burn-in, ensure convergence, and check for autocorrelation among samples.

We assessed model adequacy of the best-fitting model through posterior predictive simulation to provide a posterior predictive distribution (7) of the number of herbivorous, carnivorous, and omnivorous mammals to compare with our dataset. To do this, we randomly sampled 1,000 sets of parameters estimated from the MCMC analysis of the best model and used them to simulate a new dataset, fixing the extant taxon number to be 5,020, using the trees function in diversitree (5). Because the simulations need to be given a root state (there is no option to set the model to sample from the equilibrium-state frequencies) we ran the simulations in proportion of the number of species in each category, as those are used as the stationary frequencies in the model. We then randomly sampled 1,530 species from each simulated tree, which is the number of species in our dataset, and compared the number of herbivores, carnivores, and omnivores using a ternary diagram (Fig. S1) implemented in the vcd R package (8).

1. Bininda-Emonds ORP, et al. (2007) The delayed rise of present-day mammals. *Nature* 446:507–512.
2. Fritz SA, Bininda-Emonds ORP, Purvis A (2009) Geographical variation in predictors of mammalian extinction risk: Big is bad, but only in the tropics. *Ecol Lett* 12: 538–549.
3. FitzJohn RG, Maddison WP, Otto SP (2009) Estimating trait-dependent speciation and extinction rates from incompletely resolved phylogenies. *Syst Biol* 58:595–611.
4. Kuhn TS, Mooers AO, Thomas GH (2011) A simple polytomy resolver for dated phylogenies. *Methods in Ecology and Evolution* 2:427–436.

5. FitzJohn RG (2010) Diversitree: Comparative phylogenetic tests of diversification, version 0.7-2. Available at <http://cran.r-project.org/web/packages/diversitree>.
6. Maddison WP, Midford PE, Otto SP (2007) Estimating a binary character's effect on speciation and extinction. *Syst Biol* 56:701–710.
7. Rubin DB (1984) Bayesianly justifiable and relevant frequency calculations for the applied statistician. *Ann Stat* 12:1151–1172.
8. Meyer D, Zeileis A, Hornik K, Friendly M (2012) vcd: Visualization techniques, data sets, summary and inference procedures aimed particularly at categorical data, version 1.2-13. Available at <http://cran.r-project.org/web/packages/vcd>.



**Fig. S1.** Ternary diagram depicting the number of species generated from the posterior predictive simulations. These results from the posterior predictive simulations indicate that our real data are encompassed by the proportions of herbivores, carnivores, and omnivores generated by our best-fitting model. It is also clear that the best-fitting model does a much better job of predicting the number of omnivores than it does for either herbivores or carnivores.

**Table S1. Median speciation and extinction rates from the eight evolutionary models**

Model specification			No. of parameters	Speciation			Extinction		
Speciation rate	Extinction rate	Transition rate		Herbivore	Omnivore	Carnivore	Herbivore	Omnivore	Carnivore
Free	All constrained	Free	10	0.150 ± 0.002	0.064 ± 0.004	0.106 ± 0.003	0.012 ± 0.004	0.012 ± 0.004	0.012 ± 0.004
All constrained	Free	Free	10	0.121 ± 0.003	0.121 ± 0.003	0.121 ± 0.003	0.000 ± 2.2e-6	0.062 ± 0.015	0.045 ± 0.003
All constrained	All constrained	Free	8	0.118 ± 0.002	0.118 ± 0.002	0.118 ± 0.002	0.028 ± 0.003	0.028 ± 0.003	0.028 ± 0.003
Specialist/generalist	Specialist/generalist	Free	10	0.124 ± 0.001	0.086 ± 0.005	0.124 ± 0.001	0.000 ± 0.3e-6	0.051 ± 0.007	0.000 ± 0.3e-6
H = C	H = C								
Free	Free	All constrained	7	0.125 ± 0.013	0.130 ± 0.031	0.098 ± 0.008	0.000 ± 0.002	0.047 ± 0.028	0.009 ± 0.011
Free	Free	Specialist/generalist	9	0.137 ± 0.002	0.088 ± 0.006	0.106 ± 0.001	0.000 ± 0.000	0.058 ± 0.005	0.000 ± 0.000
		H → O = C → O							
		O → H = O → C							
		H → C = C → H							
Specialist/generalist	Specialist/generalist	Specialist/generalist	7	0.123 ± 0.013	0.09 ± 0.046	0.123 ± 0.013	0.000 ± 0.027	0.056 ± 0.055	0.000 ± 0.027
		H → O = C → O							
H = C	H = C	O → H = O → C							
		H → C = C → H							
Free	Free	Free	12	0.143 ± 0.002	0.082 ± 0.005	0.102 ± 0.002	0.000 ± 1.5e-6	0.049 ± 0.005	0.000 ± 8.2e-5

Median speciation and extinction rates estimated for the eight models analyzed ± interquartile range over the 100 replicate phylogenies. The model is specified by indicating whether the rates are free to vary depending on trophic strategy (Free), constrained by specialist or generalist (Specialist/generalist), or constrained so that rates are the same regardless of trophic strategy (All constrained). Model fit is indicated by  $\Delta AIC$ , which is the AIC value relative to that of the best-fitting model ( $\Delta AIC$  0); the best-fitting model is the most complex model that allows every rate to vary according to trophic strategy.