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REMODELING THE FOSSIL RECORD:
ANALYSIS OF EMERGENT EVOLUTIONARY AND ECOLOGICAL PATTERNS

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In memory of my grandfather, George Handelman.

TABLE OF CONTENTS

LIST OF FIGURES	vi
LIST OF TABLES	viii
ACKNOWLEDGMENTS	ix
ABSTRACT	xii
1 INTRODUCTION	1
1.1 Emergent patterns, macroevolution, and macroecology	1
1.2 Structured data and modelling emergent patterns	6
1.3 Study summaries	9
2 EXPECTED TIME-INVARIANT DIFFERENCES IN MAMMAL SPECIES DURATION	12
2.1 Introduction	12
2.2 Results	15
2.3 Discussion	19
2.4 Materials and Methods	21
2.4.1 Species occurrence and covariate information	21
2.4.2 Survival model	23
2.4.3 Estimation	28
2.4.4 Posterior evaluation	29
2.5 Supplemental information for “Death and taxa”	30
2.5.1 Supertree inference	31
2.5.2 Modeling censored observations	32
2.5.3 Deviance residuals	32
2.5.4 Variance partitioning	33
2.5.5 Widely applicable information criterion	35
2.5.6 Results from posterior predictive checks	36
2.5.7 Data quality concerns	37
2.5.8 Concerns surrounding estimates of α	38
3 HOW MACROECOLOGY AFFECTS MACROEVOLUTION: THE INTERPLAY BETWEEN EXTINCTION INTENSITY AND TRAIT-DEPENDENT EXTINCTION IN BRACHIOPODS	42
3.1 Introduction	42
3.1.1 Factors affecting brachiopod survival	44
3.2 Materials and Methods	47
3.2.1 Fossil occurrence information	47
3.2.2 Details of model	50
3.2.3 Imputation of sampling probability	52
3.2.4 Posterior inference and posterior predictive checks	54

3.3	Results	55
3.4	Discussion	64
4	SPECIES TRAITS AND ENVIRONMENTAL CONTEXT: THE CHANGING FUNCTIONAL COMPOSITION OF THE NORTH AMERICAN SPECIES POOL	71
4.1	Introduction	71
4.1.1	Background	73
4.1.2	Foreground	77
4.2	Materials and Methods	79
4.2.1	Taxon occurrences and species-level information	79
4.2.2	Environmental and temporal covariates	87
4.2.3	Modeling species occurrence	88
4.2.4	Posterior inference and model adequacy	95
4.3	Results	99
4.3.1	Comparison of estimates from the pure-presence and birth-death models	99
4.3.2	Analysis of diversity	121
4.4	Discussion	130
4.4.1	Conclusions	138
5	CONCLUSION	140
5.1	Summary	141
5.2	Synthesis	143
5.3	Future	146
5.4	Final thoughts	147
	REFERENCES	149
A	TABLE OF SPECIES MASS ESTIMATES	177

LIST OF FIGURES

2.1	Posterior predictive checks	15
2.2	Effect of mammal ecotypes on survival	17
2.3	Partitioned variance for mammal survival	18
2.4	Effect of cohort on mammal survival	19
2.5	Estimate of hazard function for mammal survival	30
2.6	Deviance residuals of fitted model	36
2.7	Additional posterior predictive checks	37
2.8	Simulation of sample size and estimates of α	39
3.1	Posterior predictive check of survival	55
3.2	Posterior predictive check of congruence	56
3.3	Estimated relationship between environment and survival	57
3.4	Cohort specific relationship between environment and survival	58
3.5	Effect of environmental covariates on survival	61
3.6	Correlation matrix of effects of environmental covariates	62
3.7	Example imputed gap statistic distributions	63
4.1	Conceptual diagram of the paleontological fourth-corner problem	78
4.2	Conceptual figure of all possible occurrence histories for an observed species	96
4.3	Posterior predictive check of average occurrence	99
4.4	Ecotype occurrence probability estimated from the pure-presence model	101
4.5	Ecotype origination probability estimated from the birth-death model	102
4.6	Ecotype survival probability estimated from the birth-death model	103
4.7	Estimates of the effect of mass on observation probability	104
4.8	Effect of mass on probability of species occurrence as estimated from the pure-presence model	105
4.9	Effect of mass on probability of species origination as estimated from the birth-death model	106
4.10	Effect of mass on probability of species survival as estimated from the birth-death model	107
4.11	Effects of group-level covariates on log-odds of ecotype occurrence as estimated from the pure-presence model	108
4.12	Effects of group-level covariates on log-odds of ecotype origination as estimated from the birth-death model	109
4.13	Effects of group-level covariates on log-odds of ecotype survival as estimated from the birth-death model	110
4.14	Estimated correlations in origination probability between ecotypes	111
4.15	Ecotypes with strong correlations in origination probability	112
4.16	Estimated correlations in survival probability between ecotypes	113
4.17	Ecotypes with strong correlations in survival probability	120
4.18	Estimated mammal log-diversity and macroevolutionary rates for the Cenozoic .	124
4.19	Estimated mammal ecotype log-diversity for the Cenozoic	125

4.20 Relative mammal ectype log-diversity for the Cenozoic	129
4.21 Estimated per capita origination rates by mammal ectype	131
4.22 Estimated per capita extinction rates by mammal ectype	132

LIST OF TABLES

2.1	Posterior estimates of covariates on mammal survival	40
2.2	Cypher for ecotype assignments	40
2.3	Equations used to estimate mammal mass	41
3.1	Parameter estimates for brachiopod survival model	70
4.1	Species trait reassigments	81
4.2	Posture assignment based on taxonomy	82
4.3	Equations used to estimate mammal mass	87
4.4	Plant phase defintions	88
4.5	Transition matrices for the pure-presence and birth-death models	89
4.6	Parameters for the observation process part of the hidden Markov model.	90
4.7	Parameters for the model of occurrence in the pure-presence model	91
4.8	Parameters for the model of presence in the pure-presence model	94
4.9	Posterior probablity estimates of differences in occurrence by plant phase	114
4.10	Posterior probablity estimates of differences in origination by plant phase	115
4.11	Posterior probablity estimates of differences in survival by plant phase	116
4.12	Posterior probablity of effects of temperature on occurrence	117
4.13	Posterior probablity of effects of temperature on origination	118
4.14	Posterior probablity of effects of temperature on survival	119
4.15	Posterior probability estimates of a peak in diversity, diversification	126
A.1	Mammal body mass estimates	177

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ABSTRACT

Macroevolution and macroecology are concerned with the patterns in evolutionary and ecological data, respectively, which arise when observing multiple species over time and/or space. Species extinction and species pool functional composition are the macroevolutionary and macroecological patterns at the heart of this dissertation. The hypotheses and analyses in the three studies forming this dissertation were all framed in terms of how species functional traits can shape these emergent patterns.

In my first study, I analyzed the Cenozoic fossil record of North American mammals to test two long standing hypotheses: the survival of the unspecialized hypothesis, and the Law of Constant Extinction. My analysis centers around a model of species duration as a function of multiple species traits, species' phylogenetic relatedness, and species' origination cohort. My results support the conclusion that generalist species will, on average, have a greater duration than more specialized species. I also find that species extinction risk increases with species duration, a result that is counter the Law of Constant Extinction. Additionally, I find that only some of the factors associated with extinction risk for Modern mammals could be considered risk factors for mammals from the rest of the Cenozoic, indicating a difference between the modern biodiversity crisis and “normal” extinction dynamics.

My second study also deals with the survival of the unspecialized and the Law of Constant Extinction, but focuses on a different system: post-Cambrian Paleozoic brachiopods. An additional aspect of this study is an analysis of the relationship between extinction intensity and the strength of trait selection. I find support for greater survival among environmental generalists than specialists. I also find evidence that for geographic range and environmental preference, as extinction intensity increases, the selective importance of these traits increases. This result is evidence for a qualitative difference between background and mass extinction.

The final study is an analysis of the changing functional composition of the North American mammal regional species pool over the last 65 million years. The goals of this analysis are to understand when functional groups are enriched or depleted, and how changes to environmental context may shape these changes. I find that mammal diversity is more strongly shaped by changes to origination probability rather than changes to extinction probability. I also find that all arboreal ecotypes declined throughout the Paleogene and disappeared from the species pool by the Neogene. Additionally, I found that most herbivore ecotypes expand their relative contribution to functional diversity over time.

My desire with this dissertation is to present the types of analyses and results that are possible through a synthesis of macroevolution and macroecology. The first step to building any dialogue is to agree on a common language and I've emphasised an expressive statistical framework with which to phrase our questions in a common tongue. My hope is that this studies serve as an example of how to use paleontological data to unite questions about the processes underlying macroevolutionary and macroecological patterns.

CHAPTER 1

INTRODUCTION

Species traits are the bridge between evolution and ecology [196, 358]. A trait is an identifiable property of an organism, such as individual body size, while a species trait is an identifiable property of the entire species, such as the average body size or geographic range of a species [196]. A class of species traits called functional traits are those traits that describe a species means of interacting with its environment such as leaf surface area or trophic role [196]. In macroevolutionary studies, analyses are typically focused on a patterns associated with a single trait or are instead traitless analyses of diversity and the diversification process [291, 290, 230, 247, 304, 129, 130, 171, 222]. Macroecological studies are frequently concerned with describing the distribution of species and individuals over space or time, such as shifts in community composition along some gradient or axis [306, 305, 33, 32, 36, 65, 90, 151, 152]. My desire with this dissertation is to present the types of analyses and results that are possible through a synthesis of both macroevolution and macroecology; my approach is to develop inference devices (i.e. statistical models) to better understand the interactions between the effects of multiple species traits, as well as those of a species' temporal and environmental context, on diversity and differential diversification both in space and time.

1.1 Emergent patterns, macroevolution, and macroecology

An emergent pattern is one that is not observable from individual constituent parts, instead the observation of many individuals is necessary. Emergence is ubiquitous in biological systems: cells form tissues which in form organs with a complex functions, species extinction requires all individual members of that species to die for possibly unrelated reasons, and a species global geographic range is the product of many individual home ranges and their

collective distribution over space. The history of a species, or set of species, over time is inherently an emergent pattern as the temporal history of a species is not knowable from an instantaneous sample. Macroevolution and macroecology are the studies of emergent patterns in evolutionary and ecological data, respectively [32, 33, 321, 320]. Traditionally, macroevolution is the study of patterns over time while macroecology is concerned with patterns over space, but I find this division overly reductive.

Both macroevolution and macroecology are disciplines concerned with emergent patterns; they both implicitly and explicitly accept a hierarchical perspective on biology as emergence is not possible without different levels of organization, however they are defined. Even if an analysis is concerned with only a single species, the species' identity is itself an emergent property or label of a collection of populations and individuals which all share a common evolutionary history. While it may be argued that the species label or identity is a non-biological construct or is simply heuristic for understanding the complexity of populations and reproductive isolation, we are still concerned with patterns associated with that construct as well as its intrinsic properties (e.g. extinction, conservation, ecosystem services) [52, 137]. My opinion is that any level which has discernible, measurable, and assignable properties is potentially worth studying, though special attention to the species level most likely affords the greatest translation between paleontological and neontological studies. In the analyses presented in this dissertation the levels of organizations thare are studied are mammal species and brachiopod genera.

Macroevolution is much more than evolution above the species level [87]; this is overly reductive and assigns too much meaning to a single level of organization rather than embracing the multitudes of possible levels of organisation. A broad definition of macroevolution, *qua* field of study, is the study of emergent evolutionary patterns; these are patterns of speciation/extinction (diversity) and trait evolution (disparity) that are observable when

considering a collection of individuals. The key here is that it is the pattern which is emergent at the level of organization is a property of the distribution of the constituent members of that organization; the traits and factors which affect or cause this pattern need not be themselves emergent. In complement, macroecology is then the study of emergent ecological patterns, which means patterns in spatial distribution or community composition which are observable only when considering more than one taxon or when considering the temporal history of one or more taxa [32, 33, 306]. Because taxa inherently respond differently and individually to environmental changes, both biotic and abiotic, macroecological patterns are those due to the similarity in response across individuals [28].

Species selection is enshrined as one of the most important patterns in macroevolution [320, 321, 345, 137, 246, 295, 224]. Rabosky and McCune [246] portray species selection as the resultant phenomena due to the heritability of speciation and extinction rates. This definition is an expansion of which phenomenon fall under this category by divorcing the idea of levels of organization from species selection and removing any requirement surrounding the mechanisms behind differences in speciation and extinction rates. The result of these distinction is two fold. First, this definition avoids unnecessary semantic conflicts surrounding “effect macroevolution” versus “strict” species selection [136, 137, 344]. Second, this definition also avoids the unproductive debate surrounding the differences between species selection versus species sorting which has been the cause for a considerable amount of confusion and rhetorical fights [343, 345, 176, 224], not the least of which is the adoption of “sorting” as an important term for understanding community assembly [331, 178, 126, 51, 310, 334, 288]. One shortcoming of the definition presented by Rabosky and McCune [246] is that it provides no operational structure to understanding the causes for differences in speciation and extinction rate. Species inherit more than just speciation and extinction rates; they also inherit traits which themselves may be linked to differences in speciation or extinction rates due to their effects on species fitness. Species fitness is a concept that is vitally important to understanding

and estimating the effect of species selection in biological systems [49, 220].

Cooper [49] defined fundamental fitness for “populations” as the expected time to extinction, otherwise called mean survival time or average persistence time; this is a fundamentally statistical definition: one entity is fitter than another if and only if it is statistically expected to survive longer. This definition applies equally to the fitness of alleles, genotype populations, and species; the only requirement is a definition of extinction. The key with this definition of fitness is that it is a *fundamental* measure in that most other definitions of fitness are derivable from probability of survival [49]. Again, this definition of fitness is for “populations” and not individuals. Instead, individuals are considered stochastic draws from the population distribution; this distinction serves to separate fitness from luck. Ultimately, the logic behind this is that a unit that is fitter is more likely to occur [49].

Species fitness is an emergent property of a species because it is a property of the distribution of its constituents [136]. Given Cooper’s definition of fitness [49] species fitness has two derived forms depending on the definition of extinction. If species extinction is defined strictly as when the last individual of a species dies, then species fitness is the expected duration of a species. If species extinction is instead defined as the loss of the last descendent along a lineage, then species fitness is the expected duration of a species and all of its descendent species; this is a lines-of-descent definition [49]. While the former deals only with species extinction, the latter of these aspects encompasses speciation. By increasing the number of descendent species, the lineage has an increased probability of occurring for longer and thus a greater expected time till extinction [49]; in effect, a higher speciation rate can lead to a greater time till extinction. And while the mechanisms behind why speciation rates can vary both across species and time is not well understood [243, 245, 52], this derived definition is agnostic to whatever mechanism controls the generation of descendants.

Extinction is a property of, or a phenomenon affecting, species as it requires the death of

all organisms within a species which do not have to all occur for the same reason [295]. Extinction is a fundamentally emergent phenomenon that is the ultimate manifestation of selection; it is also central to macroevolutionary studies and the definition of (species) fitness used here [49]. Extinction is featured centrally in one of the few “laws” in macroevolution and paleobiology: the Law of Constant Extinction [335, 175]. This law states that a species risk of going extinct is independent of that species age [335, 175], a conclusion reached via analysis of patterns of (higher) taxon survival patterns. The Red Queen hypothesis was proposed as a process that would result in the observation underlying the Law of Constant Extinction [335], though it has obviously grown to have a life of its own [175].

Extinction also features prominently in the long standing macroevolutionary hypothesis of the survival of the unspecialized [298]. This hypothesis forms the expectation that generalist species are expected to have a greater duration than specialists. This hypothesis has received a great deal of attention over the years and appears to hold in most, but not all, situations [298, 169, 217, 218, 21, 250]. Because of the strong evidence of survival of the unspecialized as a good generalization and initial condition for all analyses, it should feature heavily in all comparative analyses of species durations, survival, and extinction. In particular, this hypothesis features prominently in the first two studies of this dissertation where generality is in the context of dietary and locomotor categories (Chapter 2) and environmental preference (Chapter 3).

The functional composition of a community or species pool is a property of that unit; observing a single species at a locality does not reveal the functional composition of the community in which it interacts. The composition of a community or species pool in terms of functional groups is a community ecology exercise. Comparing the distribution of functional groups across communities or species pools is where community ecology and macroecology intersect [196, 33, 306]. In paleobiology, a successful means of classifying marine invertebrate functional

groups has been a three dimensional classification scheme called an “ecocube” which uses consistently identifiable functional traits to label both possible and observed functional groups [36, 15]. This approach also emphasizes the presence or absence of different functional groups and how functional diversity can change over time. It is this strategy that inspires the third study presented in this dissertation.

1.2 Structured data and modelling emergent patterns

An inference device is a theoretical tool for improving our knowledge by processing new information and observations [195, 149]; this device has initial conditions describing what we know (e.g. nothing), mechanisms for updating this knowledge to reflect new information, and can then produce an updated “picture” that better represents our current knowledge as well as the uncertainty surrounding this knowledge. Each inference device has a specific and narrow purpose and functionality [195]; unless the mechanisms are similar, a device for processing the rate of imperfections in the manufacturing of widgets cannot process the queuing times of callers to a help line.

We can think of the well known battery of statistical tests [311] as re-usable inference devices with very narrow utility; these are unmodifiable tools for handling very specific questions and data. All Bayesian statistical models act as inference devices because they fulfill the requirements described above: initial conditions, updating mechanism, and output as updated knowledge along with the uncertainty surrounding that knowledge [195, 149]. By developing a new model for each new question there is a precision of translation; the model actually reflects the questions at hand, something that is preferable to forcing questions and data to fit into pre-made inference devices (e.g. models, tests) that do not update knowledge in a means relevant to the actual question(s) of interest.

Structure occurs naturally in the collection of data. For example, in my analyses of mammal species and brachiopod genus durations I grouped species that originated in the same temporal unit as a “cohort” (Chapter 2, and 3), the assumption being that species originating at the same time may be more similar than species that originate at different times. By grouping these species together the differences in extinction risk associated with each cohort can be estimated. If these groupings are ignored, species are all assumed to have identical properties. Additionally, by including these cohort labels in the model of species durations, the estimates for each cohort are tempered and drawn towards the overall average cohort effect through regularization [101, 195]. By sharing information across cohorts, the predictive properties of the model to out-of-sample data are improved [101, 195]. Finally, including these groupings in the actual model better reflect the actual data at hand.

Two of the most important analytical approaches at the core of macroevolutionary study are the birth-death process for diversification in both discrete and continuous time [257, 253, 212, 214, 211, 213, 316], and the random walk heuristic for continuous trait evolution [256, 70, 29, 104, 261, 262, 263, 287, 130, 129]. All three of the studies covered by this dissertation make use of some variant to the birth-death process. The first two studies are analyses of extinction, which is a pure-death processes. The third study utilizes a discrete-time birth-death process to model species presence in a species pool.

Similar analytical foundations are harder to identify for macroecology as a whole, so instead I will focus on species distribution models (SDMs) as a powerful framework for understanding the distribution of species, due to environmental factors [64] and species traits [288]. SDMs are a class of models which attempt to operationalize the multitude of processes which result in the distribution of one or more species in space and/or time. In effect, SDMs are a means of operationalizing the concept of a species’ “realized niche” in order to understand the limits on a species distribution [64]. Typically, SDMs are used to analyze the relationship between

species presence at a locality and the environmental factors which characterize that locality. From this analysis, the possible distribution of a species in space can be then estimated and compared to the observed distribution of that species [64, 11, 229]. The maximum entropy theory of community assembly, and its related model, view community assembly as an ecological sorting process where traits mediate the effects of environmental filters [288, 356]; also called “community assembly via trait selection” (CATS regression). By analyzing the composition of species at localities based on their traits, the strength and relative importance of the traits that most directly structure community composition be elucidated.

Both of these approaches to analyzing species distributions can be united in a single fourth-corner model [356, 31]. The fourth-corner problem is an old problem in community ecology originating in the multivariate analysis literature: assuming species distribution is the result of functional traits interacting with environmental factors, how do we estimate which interactions are important and their relative strengths [164, 61]? By phrasing the fourth-corner problem as a model based framework, results are much more easily interpretable and actually provide estimates of the effects of species traits and environmental factors instead of the simple significance provided from the older Monte Carlo based methodology [31, 139, 232, 231].

The third study in this dissertation makes extensive use of this framework by casting the fourth-corner problem into an additional dimension: time. By combining the fourth-corner framework outlined above [356, 31] with the birth-death process used for modeling diversification into a single unified model of species occurrence through time as a function of both species traits and changing environmental context I’ve developed a powerful analytical bridge between macroecology and macroevolution.

I emphasize model-based approaches to analysis as well as question or study specific models because a common language is necessary for clear, coherent, and translatable results that actually relate to the question(s) at hand. Some of the greatest limits to paleobiological,

macroevolutionary, and macroecological study are a lack of strong, mechanistic predictive theories that can be expressed mathematically. Some of the greatest strides in advancing discussions of macroevolutionary and macroecological theory disputes have come from translating verbal theory into mathematical and statistical models [257, 212, 70, 130, 129, 288]. The complex realities of the biological processes which shape diversity are rarely integrated into paleobiological analyses of macroevolutionary and macroecological patterns. A move to a model-forward approach to paleobiology, heavily steeped in evolutionary and ecological theory, would be beneficial for advancement of theories in macroevolution and macroecology.

Paleobiologists historically believe that neontologists ignore their approaches and insights into macroevolution and macroecology study and theory [283, 282], but without a concerted effort to engage within the same theoretical framework and language when developing scientific questions and the related analytical tools (i.e. statistical models) this worry and resentment is all but preordained. Because the systems paleobiologists study are unknown to, have no direct impact on, or are contextualized with respect to the systems studied by neontologists, a push towards unification and synthesis most likely has to begin with the paleobiological community; luckily, it appears that the neontological community is receptive to paleobiological insight [97]. The simplest and fastest way to begin not just a dialogue but a unification is by translating verbal macroevolutionary and macroecological theories from paleobiology into statistical models that are readable by all researchers, both paleontological and neontological.

1.3 Study summaries

Each of the three studies that make up this dissertation involves developing a hierarchical model to describe structured data with the goal of making macroevolutionary and/or macroecological inference. The first two studies are decidedly macroevolutionary in bent as they

are analyses of trait-based extinction patterns in mammals and brachiopods, respectively. The third study is an analysis of mammal species pool temporal dynamics and is of a strong macroecological bent, though makes use of a macroevolutionary model of diversification in order to describe species turnover.

The first study presented is an analysis of North American mammal species durations and trait-based extinction risk (Chapter 2). This analysis is principally concerned with the long standing hypothesis of the survival of the unspecialized which states that average or generalist species are expected to have a greater duration than specialists or other extreme forms [298, 169, 170]. Species duration is a proxy for species extinction risk as species with a shorter duration experience a greater extinction risk than species with a long duration. In this study, differences in species extinction risk based on multiple functional traits are estimated while also taking into account time of species origination as well as its relative phylogenetic position. Finally, the possibility of species age affecting extinction risk is also considered because while the Law of Constant Extinction is extremely hard to “test” it has never definitively been proven [335, 175].

The second study presented is also an analysis of taxon durations, but this time focuses on all post-Cambrian Paleozoic brachiopod genera (Chapter 3). The question at the center of this study is “what happens to the effects of functional traits on survival when average survival increases or decreases?” Unlike the previously described study, which focused on the average effects of functional traits on survival, this study requires estimates of how the effects of functional traits vary through time. The key parameters are those of the correlation matrix of the effects of these traits on duration and the average duration of species originating at the same time. This study also has results relevant to the survival of the unspecialized hypothesis with respect to the effect of environmental preference on survival, and the Law of Constant Extinction by allowing survival to be a function of taxon duration.

As mentioned above, the third study presented is decidedly more macroecological in focus as it is an analysis of how a regional species pool changes over time due to species turnover and a changing environmental context (Chapter 4). The fundamental question is “when are certain ecotypes enriched or depleted with respect to their diversity history?” To that end, I analyze the set of North American mammals for the Cenozoic and the changing functional composition of that species pool from nearly the beginning of the Cenozoic to almost the very recent (64.2 million years ago). In this analysis, functional composition of the species pool is described as the relative diversity of 18 different mammal ecotypes which are defined for every species as its dietary and locomotor combination. The occurrence of an ecotype, in terms of origination and survival, is modeled as a function of that species environmental context as described by the dominant plant groups in North America as well as global temperature estimates.

All three of these analyses feature a hierarchical Bayesian model developed explicitly for each study in order to clearly attempt to answer the questions at hand. Each of these studies exemplifies my earlier rhetoric of how to build and advance macroevolutionary and macroecological study and theory through the explicit phrasing of scientific questions, precision of translation from question to analysis, and the mobilization of domain specific knowledge to cast results both in terms of the system specific insights as well as the theoretical insights.

CHAPTER 2

EXPECTED TIME-INVARIANT DIFFERENCES IN MAMMAL SPECIES DURATION

Determining which biological traits influence differences in extinction risk is vital for understanding the differential diversification of life and for making predictions about species' vulnerability to anthropogenic impacts. Here I present a hierarchical Bayesian survival model of North American Cenozoic mammal species durations in relation to species-level ecological factors, time of origination, and phylogenetic relationships. I find support for the "survival of the unspecialized" as a time-invariant generalization of trait-based extinction risk. Furthermore, I find that phylogenetic and temporal effects are both substantial factors associated with differences in species durations. Finally, I find that the estimated effects of these factors are partially incongruous with how these factors are correlated with extinction risk of the extant species. This parallels previous observations that background extinction is a poor predictor of mass extinction events and suggests that attention should be focused on mass extinctions to gain insight into modern species loss.¹

2.1 Introduction

Why extinction risk varies among species remains one of the most fundamental questions in paleobiology and conservation biology [298, 335, 255, 240, 347]. To address this issue, I test for similarities in associations between extinction risk and multiple species-level traits during times of background extinction and in the modern world; which traits have time-invariant effects on species duration; and whether extinction is age-independent. I approach

1. This chapter was previously published as an article in the *Proceedings of the National Academy of Sciences* [309] and is reproduced here as allowed by publisher.

these questions together by using a model of species duration whose parameter estimates act as direct tests of these questions. Cenozoic mammals are an ideal focus for this study because their fossil record is well sampled and well resolved both temporally and spatially, and because individual species ecology and taxonomic position are generally understood [298, 240, 6, 171, 305, 329, 191].

Time-invariant factors are those that have a constant directional effect even if their magnitude varies. Because change in the magnitude of extinction risk is not necessarily the best indicator of a shift from background to mass extinction [349], it is better to look for changes in either the direction of selection, the loss of a selective pressure, or the appearance of novel selective pressures [134].

The species-level traits studied here are bioprovince occupancy, body mass, and dietary and locomotor categories. These traits are related to aspects of a species' adaptive zone such as population density, expected range size, potential prey, and dispersal ability [305, 152] and are a combination of aggregate and emergent traits [137]. It is expected that species with larger geographic ranges have lower extinction rates than species with smaller geographic ranges [134, 270]; however, how traits more directly related to species–environment interactions may affect species extinction risk is more nebulous.

Body size is a complex trait related to many life history characteristics. There are three general hypotheses of how body size may affect extinction risk: 1) positive effect where an increase in body size causes an increase in extinction risk, potentially due to associated decrease in reproductive rate or other similar life history traits [171, 172]; 2) negative effect where an increase in body size causes a decrease in extinction risk because of an expected positive relationship between body size and geographic range; and 3) no effect of body size on extinction risk [329].

The strongest expectation for the effects of dietary category on extinction risk is that omnivores will have the lowest extinction risk of all species. This expectation is based on the long standing “survival of the unspecialized” hypothesis where more generalist species (e.g. omnivores) have greater survival than specialist species (e.g. carnivores/herbivores) [298, 169]. It has also been observed that both carnivores and herbivores have greater diversification rates than omnivores, with herbivores diversifying faster than carnivores [237]. How this result translates into differences in extinction risk is currently unknown [241]. In modern taxa, higher trophic levels (e.g. carnivores versus herbivores) have been associated with greater extinction risk, most likely because of human extermination of top predators [172, 238].

Similarly, there are few expectations of how locomotor category may effect extinction risk. During the Cenozoic, there was a shift at the Paleogene/Neogene boundary from predominately closed to predominately open environments [28, 142]. Based on this observation, a prediction is that arboreal taxa will have the greatest extinction risk of all, with both scansorial and ground dwelling taxa having lower extinction risks.

I use a hierarchical Bayesian survival model of species duration as predicted by the covariates of interest along with species’ temporal and phylogenetic context. Species duration, in 2 My bins, was modeled as realizations from either an exponential or Weibull distribution-based hierarchical model [101]. The exponential distribution corresponds to the Law of Constant Extinction, which states that extinction is age-independent [335]. Note that the exponential is a special case of the Weibull when its shape parameter, α , is 1. The Weibull distribution allows for extinction to be taxon-age dependent, where values of α greater than 1 corresponds to increasing risk with age and values less than 1 corresponds to decreasing risk with age. Origination cohort and phylogenetic position were modeled as independent effects. Phylogenetic effect was modeled assuming species duration may have evolved via a Brownian motion-like process [184, 128]. The results from the Weibull model are detailed

here because this model has a better fit to the data than the exponential (Weibull WAIC 6140.37, exponential WAIC 16697.35; Fig. 2.1, S1, S2).

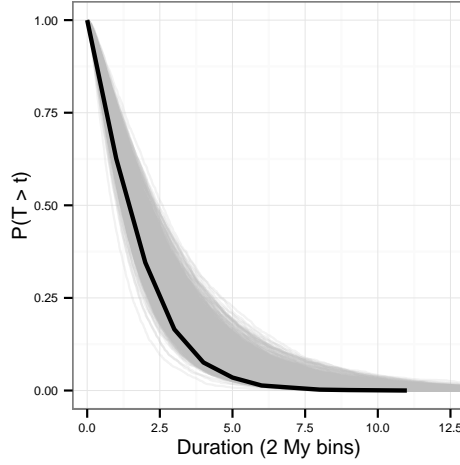


Figure 2.1: Weibull-based model estimates (grey) from 1000 posterior predictive data sets of the empirical survival function (black). The survival function is the probability that a species with duration t will not have gone extinct. Simulated data sets were generated by drawing parameter values randomly from their estimated posteriors and using the observed covariate information to estimate durations for all the observed species.

2.2 Results

A summary of the posterior distributions for the most relevant parameter estimates is presented in Table 2.1. All posterior inference is based on these estimates. For the results from the posterior predictive checks and discussion of the estimation of α , please see the accompanying Supplemental information (Section 2.5). Additionally, see the Supplemental information for discussion surrounding use of Paleobiology Database and accompanying data quality concerns.

Species with greater bioprovince occupancy are found to be associated with lower extinction risk than taxa with smaller bioprovince occupancy ($\beta_{occupancy}$ mean = -0.53 , std = 0.08). This is consistent with previous findings. Body size has nearly zero association with expected

duration (β_{size} mean = -0.05 , std = 0.05), a similar result to some previous studies [329]. However, previous studies were performed at the generic level and were unable to determine how body size may effect species-level extinction, as the effect of either extinction or speciation cannot be distinguished [171, 329].

Some clear patterns emerge from the pairwise differences in effect of each dietary category on expected duration (Fig. 2.2). Consistent with expectations from the “survival of the unspecialized” hypothesis [169, 298], omnivory appears to be associated with the lowest expected extinction risk. Carnivory is associated with a greater expected duration than either herbivory or insectivory, but a greater expected extinction risk than omnivory. Finally, herbivory and insectivory have approximately equal effects on expected duration. Given previous results, these results imply that carnivores have a greater origination rate than omnivores [237]. These results also imply that herbivores, which have the greatest extinction risk, must also have a very high origination rate in order to have the greatest diversification rate among these three categories [237].

For locomotor category, both scansoriality and ground dwelling life habitat are associated with a greater expected duration than arboreality (Fig. 2.2). Scansorial and ground dwelling life habits also have approximately equal expected effects on extinction risk. This is consistent with the expectation that arboreality will confer greater extinction risk due to the loss of associated environment with the shift from open to closed habitat at the Paleogene/Neogene boundary [28]. However, there are two possible processes which could lead to the observed pattern: arboreality confers an intrinsic difference in extinction risk or it might not be that arboreal taxa have an intrinsically higher risk but were instead “hit harder” by the environmental shift than other taxa. This analysis cannot distinguish between these two processes. Note that, while this is a study of North American Cenozoic mammals, for European Cenozoic mammals this transitionary period corresponds to the Vallesian which

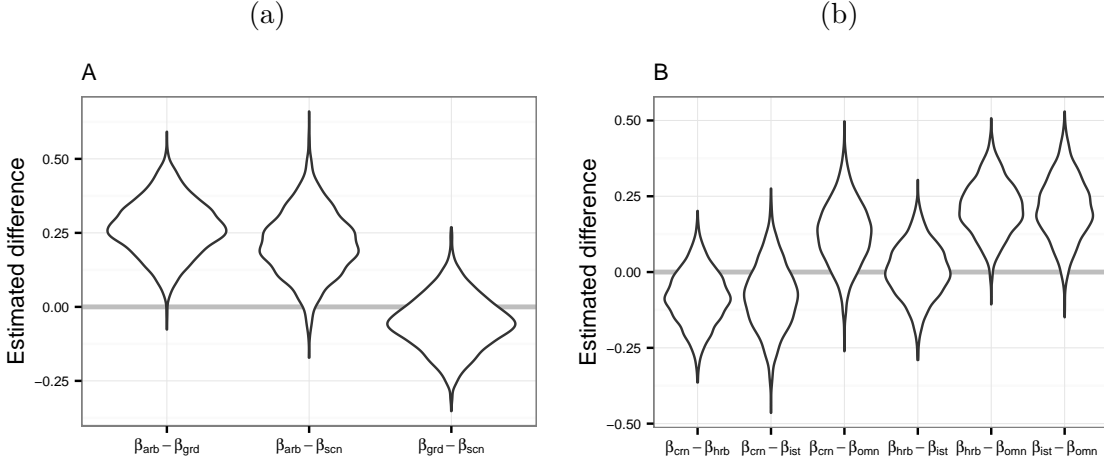


Figure 2.2: Pairwise differences in effect of the locomotor (**A**) and dietary categories (**B**) on expected duration from 1000 samples from the posterior distribution. Comparisons of locomotor categories, from top to bottom (**A**), are: arboreal ($\beta_{arb} = \beta_0$) versus ground dwelling ($\beta_{grd} = \beta_0 + \beta_g$), arboreal versus scansorial ($\beta_{scn} = \beta_0 + \beta_s$), and ground dwelling versus scansorial. For dietary category, from top to bottom (**B**): carnivore ($\beta_{crn} = \beta_0$) versus herbivore ($\beta_{hrb} = \beta_0 + \beta_h$), carnivore versus insectivore ($\beta_{ist} = \beta_0 + \beta_i$), carnivore versus omnivore ($\beta_{omn} = \beta_0 + \beta_o$), herbivore versus insectivore, herbivore versus omnivore, and insectivore versus omnivore. Negative values indicate that the first category is expected to have a greater duration than the second, while positive values indicate that the first category is expected to have a shorter duration.

was a sudden shift in species demography away from arboreality [1, 209].

Of the three sources of variance present in the model, individual species variance accounts for approximately 80% of the observed, unmodeled variance (Fig. 2.3). Note that the individual variance was approximated using an simulation approach [107] because the Weibull distribution does not have a variance term. Both cohort and phylogenetic effects account for the other 20% of the observed variance. This result means that extinction risk has both temporal and phylogenetic aspects, as both contribute greater than 0% of the observed variability in the data [128].

The estimates for the individual cohort effects show a weak pattern of greater extinction risk in older Cenozoic cohorts compared to younger cohorts (Fig. 2.4). This potential slowdown

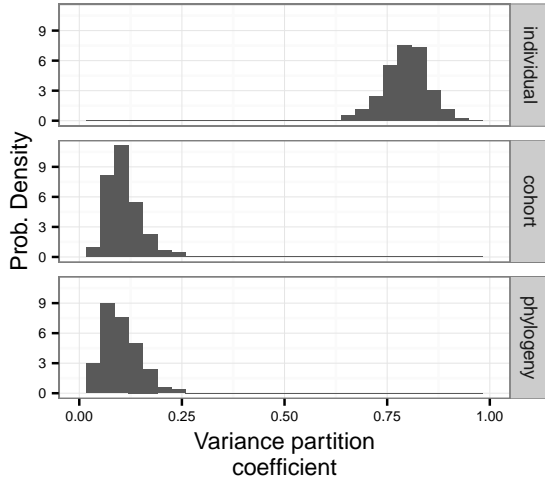


Figure 2.3: Estimates of the variance partitioning coefficients for the three different sources of variance: species, cohort, and phylogeny. Higher values correspond to greater contribution to total observed variance. Each of the estimates is a distribution of 1000 approximating simulations due to the model’s non-normally distributed errors.

in extinction risk is consistent with previous analyses of marine invertebrates [258, 82] and mammals [7, 10]. There are two prevailing hypotheses as to the cause of this slowdown: 1) extinction risk is constant within, but varies between, clades so over time clades with low extinction rates increases in proportion of total diversity thus bringing down expected extinction risk; or 2) over time taxa increase in mean fitness and thus decrease in expected extinction risk [258]. The observed decrease in extinction risk with age, along with the variance partitioning results (Fig. 2.3) are consistent with both of these hypotheses with neither being more “important” than the other.

Interestingly, the shift from older cohorts with a higher extinction risk to younger cohorts with lower extinction risk is approximately at the Paleogene–Neogene boundary. Given the association with arboreality and increased extinction risk (Fig. 2.2), the decrease in expected extinction risk over time might relate to the preferential loss of arboreal taxa over the Cenozoic. However, because the model used here does not allow for time-varying effects, I cannot identify whether this boundary is associated with a shift in the direction or magnitude

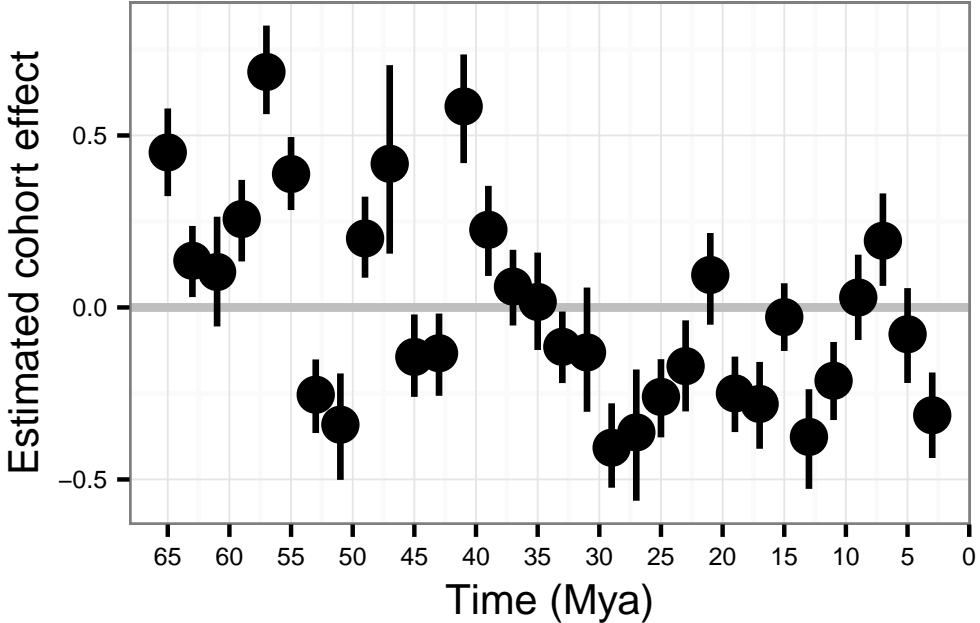


Figure 2.4: Summaries of posterior estimates of individual cohort effect depicted as medians and 80% credible intervals. High values correspond to shorter species durations while lower values correspond to greater species durations compared to the mean duration. Lines are placed at the middle of the 2 My origination cohorts.

of the expected effect of arboreality on extinction risk.

2.3 Discussion

My results indicate that Cenozoic North American mammal “generalists” are expected to have a lower extinction risk than “specialists.” This implies that the diversification of specialized taxa would have required either a driven trend away from generality [199] or an increase in speciation rate relative to extinction rate [320]. This requires that specialist traits should somehow increase or be associated with increases in speciation rate, perhaps via niche partitioning or changes in habitat heterogeneity. For example, descendant species of omnivores many divide available prey items more finely or arboreal taxa may increase in both extinction and speciation rates via increases in habitat heterogeneity. Possible evidence

to support this hypothesis would be to demonstrate differences in speciation rate associated with those traits analyzed here or other similar traits.

When these results are compared to factors contributing to increased extinction risk in extant mammals, there are some incongruencies. As expected, large range size is consistently associated with lower extinction risk in the modern world [172, 238, 95, 96]. While my analysis found body size to have almost no time-invariant effect on extinction risk, in extant mammals this is not necessarily the case as increased body size is associated with increased extinction risk [172, 238]. However, this pattern is partially clade dependent [95]. As stated earlier, higher trophic levels have been found to be associated with greater extinction risk in extant mammals [172, 238]. In contrast, I found that omnivores and carnivores have a lower expected extinction risk than either insectivores or herbivores (Fig. 2.2). Finally, phylogeny has been found to be a good predictor of differences in extinction risk in extant mammals as certain clades are at much higher risks than others [96]. This effect seems much greater in the Recent than for the whole Cenozoic, implying that current extinction risk is more phylogenetically concentrated than during times of background extinction levels during the Cenozoic.

Whether these incongruities are within the standard range of time-variant effects is unknown, though my comparisons do imply that current processes are different from those studied here. However, this is not a model of what makes taxa vulnerable during mass extinctions and that may account for these incongruities, assuming mass extinctions are qualitatively different than background extinction [134]. These results would also be inapplicable if the current biodiversity crisis is qualitatively different from either background or mass extinction as preserved in the fossil record.

By modeling how different ecologies and historical factors effect a species' expected extinction risk, it is possible to better understand what processes may have driven the resulting pattern of

selection (i.e. diversity) while also providing a baseline for evaluating the current biodiversity crisis. This analysis finds support for the “survival of the unspecialized” hypothesis [298, 169] as a time-invariant generalization about extinction risk. I also find that there are substantial effects of both cohort and phylogeny on extinction risk, which supports the idea that the decrease in extinction risk [258] over time has both temporal and phylogenetic components. Additionally, I found evidence of increasing extinction risk with species age, the cause of which is unknown. These results show that, like prior mass extinction events in the fossil record, the current biodiversity crisis is qualitatively different from the previous period of background extinction in the fossil record [134].

2.4 Materials and Methods

2.4.1 *Species occurrence and covariate information*

Fossil occurrence information was downloaded from the Paleobiology Database (PBDB; <http://paleodb.org/>). Occurrence, taxonomic, stratigraphic, and biological information was downloaded for all North American mammals. This data set was filtered so that only occurrences identified to the species-level, excluding all “sp.”-s. All aquatic and volant taxa were also excluded. Additionally, all occurrences without latitude and longitude information were excluded from the sample.

Species dietary and locomotor category assignments were done using the assignments in the PBDB, which were reassigned into coarser categories (Table 2.2). This was done to improve interpretability, increase sample size per category, and make results comparable to previous studies [152, 237].

All individual fossil occurrences were assigned to 2 My bins ranging through the entire

Cenozoic. Taxon duration was measured as the number of 2 My bins from the first occurrence to the last occurrence, inclusive. This bin size was chosen because it approximately reflects the resolution of the North American Cenozoic mammal fossil record [6, 10, 191]. Species originating in the youngest cohort, 0-2 My, were excluded from analysis because every species duration would be both left and right censored, which is illogical.

Species body size estimates in grams were sourced from a large selection of primary literature and database compilations. Databases used include the PBDB, PanTHERIA [156], and the Neogene Old World Mammal database (NOW; <http://www.helsinki.fi/science/now/>). Major sources of additional compiled body size estimates include [329, 30, 94, 198, 248, 307]. These were then supplemented with an additional literature search to try and fill in the remaining gaps. In many cases, species body mass was estimated using various published regression equations based on tooth or skull measurements (Table 2.3). If multiple specimens were measured, I used the mean of specimen measures as the species mean. See Appendix A for a complete list of mass estimates and sources.

Biogeographic network

Species geographic extent was measured as the mean of the relative number of bioprovinces occupied by a species for each 2 My bin the species was present. Bioprovinces were identified using a network-theoretic approach that has previously been applied to paleontological data [289, 341]. This approach relies on defining a biogeographic bipartite network of taxa and localities. In this study, taxa were defined as species and localities were grid cells from a regular lattice on a global equal-area cylinder map projection. The regular lattice was defined as a 70 x 34 global grid where each cell corresponds to approximately 250000 km². An advantage of this approach is that this approach reduces to occupancy when all localities are independent and to a single bioprovince when all localities are identical.

A biogeographic network was constructed for each of the 2 My bins used in this study. Emergent bioprovinces were then identified using the map equation [269, 268] as has been done before [289, 340, 341]. These bioprovinces correspond to taxa and localities that are more interconnected with each other than with other nodes.

The map projection and regular lattice were made using shape files from <http://www.naturalearthdata.com/> and the **raster** package for R [122]. Bioprovience identification was done using the map equation as implemented in the **igraph** package for R [56].

Supertree

As there is no single, combined formal phylogenetic hypothesis of all Cenozoic fossils mammals from North America, it was necessary to construct a semi-formal supertree. This was done by combining taxonomic information for all the observed species and a few published phylogenies using matrix representation parsimony [25]. For further explanation of the methodology used to construct this supertree, please see the Supplementary information in Section 2.5.

2.4.2 Survival model

Presented here is the model development process used to formulate the two survival models used in this study. First, define y as a vector of length n where the i th element is the duration of species i , where $i = 1, \dots, n$.

The simplest survival model where durations are assumed to follow an exponential distribution

with a single “rate” or inverse-scale parameter λ [160]. This is written

$$\begin{aligned} p(y|\lambda) &= \lambda \exp(-\lambda y) \\ y &\sim \text{Exp}(\lambda). \end{aligned} \tag{2.1}$$

The exponential distribution corresponds to situations where extinction risk is independent of age. To understand this, we need to define two functions: the survival function $S(t)$ and the hazard function $h(t)$. $S(t)$ is the probability that a species having existed for t 2 My bins will not have gone extinct while $h(t)$ corresponds to the instantaneous extinction rate for some taxon age t [160]. For an exponential model, $S(t)$ is

$$S(t) = \exp(-\lambda t) \tag{2.2}$$

and $h(t)$ is defined

$$h(t) = \lambda \tag{2.3}$$

The choice of the exponential distribution corresponds directly to the Law of Constant Extinction [335] as the right side of Eq. 2.3 does not depend on species age t .

The current sampling statement (Eq. 2.1) assumes that all species share the same rate parameter with no variation. To allow for variation in λ associated with relevant covariate information like species body size, λ is reparameterized as $\lambda_i = \exp(\sum \beta^T \mathbf{X}_i)$ with i indexing a given observation and its covariates, β is a vector of regression coefficients, and \mathbf{X} is a matrix of covariates. This is a standard regression approach, where one column of \mathbf{X} is all 1-s and its corresponding β coefficient is the intercept.

\mathbf{X} is an $n \times K$ matrix of species-level covariates. Three of the covariates of interest are the logit of mean relative occupancy, and the logarithm of body size (g). The discrete covariate

index variables of dietary and locomotor category were transformed into $n \times (k - 1)$ matrices where each column is an indicator variable (0/1) for that species's category, k being the number of categories of the index variable (3 and 4, respectively). Only $k - 1$ indicator variables are necessary as the intercept takes on the remaining value. For example, the difference in effect of arboreality versus scansoriality on extinction risk, given that arboreality is the reference category, is the coefficient for the scansorial indicator variable as that is the difference between the effect of arboreality (the intercept β_0) and scansoriality (the intercept + scansorial effect β_s); Fig. 2.2). Finally, a vector of 1-s was included in the matrix \mathbf{X} whose corresponding β coefficient is the intercept, making K equal eight.

β is the vector of regression coefficients. The intercept term was given a weak normal prior, $\beta_0 \sim \mathcal{N}(0, 10)$ while all of these other coefficients were slightly more informative priors, e.g. $\beta_{mass} \sim \mathcal{N}(0, 5)$. These priors were chosen because it is expected that the effect size of each variable on duration will be small, as is generally the case with binary covariates [102].

Regression coefficients are not directly comparable without first standardizing the input variables to have equal standard deviations. This is accomplished by subtracting the mean of the covariate from all values and then dividing by the standard deviation, resulting in a variable with mean of zero and a standard deviation of one. This linear transform greatly improves the interpretability of the coefficients as expected change in mean duration given a difference of one standard deviation in the covariate [275]. Additionally, this makes the intercept directly interpretable as the estimate of mean (transformed) σ (Eq. 2.7). However, because the expected standard deviation for a random binary variable is 0.5, in order to make comparisons between the binary and continuous variables, the continuous inputs were divided by twice their standard deviation [100].

Origination cohort is defined as the group of species which all originated during the same 2 My temporal bin. Because the most recent temporal bin, 0-2 My, was excluded, there are

32 total cohorts. The effect of origination cohort j was modeled with each group being a sample from a common cohort effect, η , which was considered normally distributed with mean 0, and standard deviation σ_c . The value of σ_c was then estimated from the data itself, corresponding to the amount of pooling in the individual estimates of η_j . This approach is a conceptual and statistical unification between dynamic and cohort survival analysis in paleontology [77, 252, 251, 336, 21], with σ_c acting as a measure of compromise between these two end members. The choice of the half-Cauchy prior for σ_c follows [99].

$$\begin{aligned}\eta_j &\sim \mathcal{N}(0, \sigma_c) \\ \sigma_c &\sim \text{C}^+(0, 2.5)\end{aligned}$$

The impact of shared evolutionary history, or phylogeny, was modeled as an individual effect where each observation, i , is modeled as a multivariate normal, h , where the covariance matrix Σ is known up to a constant, σ_p^2 [184, 128]. This is written

$$\begin{aligned}h &\sim \text{MVN}(0, \Sigma) \\ \Sigma &= \sigma_p^2 \mathbf{V}_{phy} \\ \sigma_p &\sim \text{C}^+(0, 2.5).\end{aligned}$$

\mathbf{V}_{phy} is the phylogenetic covariance matrix defined as an $n \times n$ matrix where the diagonal elements are the distance from root to tip, in branch length, for each observation and the off-diagonal elements are the amount of shared history, measured in branch length, between observations i and j . σ_p was given a weakly informative half-Cauchy hyperprior. Note that because the phylogeny used here is primarily based on taxonomy, estimates of σ_p represent minimum estimates [184, 128]. Improved phylogenetic estimates of all fossil Cenozoic mammals would greatly improve this estimate.

To relax the assumption of age-independent extinction of the Law of Constant Extinction, the Weibull distribution is substituted for the exponential [160]. The Weibull distribution has a shape parameter α and scale parameter σ . Conceptually, σ is the inverse of λ . α modifies the impact of taxon age on extinction risk. When $\alpha > 1$ then $h(t)$ is a monotonically increasing function, but when $\alpha < 1$ then $h(t)$ is a monotonically decreasing function. When $\alpha = 1$ then the Weibull distribution is equivalent to the exponential.

The Weibull distribution and sampling statement were defined

$$p(y|\alpha, \sigma) = \frac{\alpha}{\sigma} \left(\frac{y}{\sigma}\right)^{\alpha-1} \exp\left(-\left(\frac{y}{\sigma}\right)^\alpha\right)$$

$$y \sim \text{Weibull}(\alpha, \sigma). \quad (2.4)$$

The corresponding $S(t)$ and $h(t)$ functions are defined

$$S(t) = \exp\left(-\left(\frac{t}{\sigma}\right)^\alpha\right) \quad (2.5)$$

$$h(t) = \frac{\alpha}{\sigma} \left(\frac{t}{\sigma}\right)^{\alpha-1}. \quad (2.6)$$

To allow for σ to vary with a given observation's covariate information it is reparameterized in a similar fashion to λ with a few key differences. Because $\sigma = 1/\lambda$ in order to preserve the interpretation of β , while taking α into account, σ is reparameterized as

$$\sigma_i = \exp\left(\frac{-\beta}{\alpha}\right). \quad (2.7)$$

Given the above, the survival model was then fit in a Bayesian context using both exponential and Weibull distributions. The Weibull's α parameter was assumed constant across species, which is standard practice in survival analysis [160]. α was given a weakly informative half-

Cauchy (C^+) prior. σ was reparameterized as an exponentiated regression model (Eq. 2.7). This was further expanded (Eq. 2.8) to allow for two hierarchical factors as discussed above. This is written

$$\sigma_i = \exp\left(\frac{-(h_i + \eta_j[i] + \sum \beta^T \mathbf{X}_i)}{\alpha}\right) \quad (2.8)$$

where equivalent statement for the exponential distribution is defined

$$\lambda_i = \exp\left(h_i + \eta_j[i] + \sum \beta^T \mathbf{X}_i\right). \quad (2.9)$$

An important part of survival analysis is the inclusion of censored observations where the failure time has not been observed [133, 160]. The most common censored observation is right censored, where the point of extinction had not yet been observed in the period of study, such as taxa that are still present in the most recent time bin (0-2 My). Left censored observations, on the other hand, correspond to observations that went extinct any time between 0 and some known point. To account for this uncertainty, the probability of a left censored observation is found by integrating over all possible durations between 0 and 1 time bin. For an explanation of how censored observations are included in the sampling statement, please see the Supplementary information in Section 2.5.

2.4.3 Estimation

Parameter posteriors were approximated using a Markov-chain Monte Carlo (MCMC) routine implemented in the Stan programming language [318]. Stan implements a version of Hamiltonian Monte Carlo called the No-U-Turn sampler [124]. Posterior approximation was done using four parallel MCMC chains run for 30000 steps, thinned to every thirtieth sample, split evenly between warm-up and sampling. Convergence was evaluated using the scale reduction factor, \hat{R} . Values of \hat{R} close to 1, or less than or equal to 1.1, indicate approximate

convergence. Convergence means that the chains are approximately stationary and the samples are well mixed [101].

2.4.4 Posterior evaluation

The most basic assessment of model fit is that simulated data generated given the model should be similar to the observed. This is the idea behind posterior predictive checks. Using the covariates from each of the observed durations, and randomly drawn parameter estimates from their marginal posteriors, a simulated data set y^{rep} was generated. This process was repeated 1000 times and the distribution of y^{rep} was compared with the observed [101]. For results from the posterior predictive tests used in this study, please see the Supplementary information in Section 2.5.

The exponential and Weibull models were compared for out-of-sample predictive accuracy using the widely-applicable information criterion (WAIC) [357]. Because the Weibull model reduces to the exponential model when $\alpha = 0$, our interest is not in choosing between these models. Instead comparison of WAIC values is useful for better understanding the effect of model complexity on out-of-sample predictive accuracy. An explanation of how WAIC is calculated is presented in the Supplementary information (Section 2.5) following the recommended “WAIC 2” formulation [101].

There are three different variance components in this model: sample component, cohort σ_c^2 , and phylogenetic σ_p^2 . Partitioning the variance between these sources allows the relative amount of unexplained variance of the sample to be compared. The sample component is similar to the residual variance from a normal linear regression. However, the Weibull based model used here (Eq. 2.4) does not include an estimate of the variance similar to the squared scale term of the a Normal distribution. Instead, the sample component was approximated

via a simulation approach modified from [107]. For explanation of this method, please see Supplementary information (Section 2.5).

I used variance partitioning coefficients (VPC) to estimate the relative importance of the different variance components [102]. Phylogenetic heritability, h_p^2 [184, 128], is identical to the VPC of the phylogenetic effect. Phylogenetic heritability is a measure of how shared evolutionary history impacts differences in individual species trait values (e.g. duration). This is a broad sense “heritability” as it combines both genetic inheritance and other, non-genetic shared history factors. Importantly, because phylogenetic effect was estimated using a principally taxonomy based tree the estimates derived here can be considered minimum estimates of the phylogenetic effect.

2.5 Supplemental information for “Death and taxa”

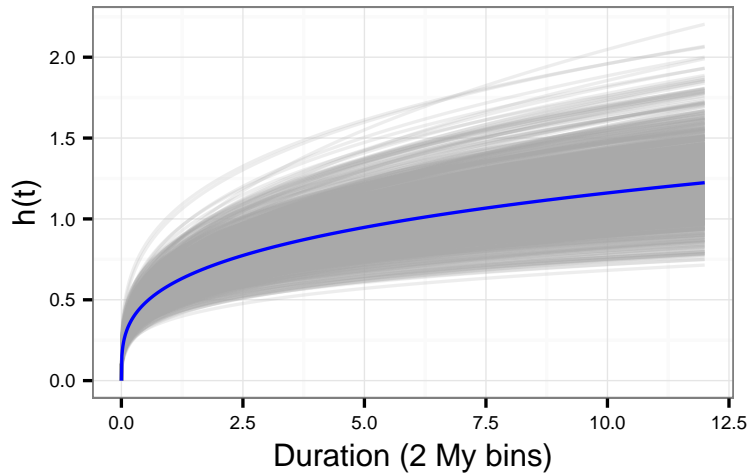


Figure 2.5: 1000 estimates of the hazard function ($h(t)$) for the observed species mean (grey), along with the median estimated hazard function (blue). $h(t)$ is an estimate of the rate at which a species of age t is expected to go extinct. Hazard functions were estimated from random draws from the estimated posterior distributions and evaluated with all covariate information set to 0, which corresponds to the expected duration of the mean species.

2.5.1 Supertree inference

As there is no single, combined formal phylogenetic hypothesis of all Cenozoic fossils mammals from North America, it was necessary to construct a semi-formal supertree. This was done by combining taxonomic information for all the observed species and a few published phylogenies.

The initial taxonomic classification of the observed species was based on the associated taxonomic information from the PBDB. This information was then updated using the Encyclopedia of Life (<http://eol.org/>) which collects and collates taxonomic information in a single database. This was done programatically using the `taxize` package for R [41]. Finally, this taxonomic information was further updated using a published taxonomy of fossil mammals [146, 145].

This taxonomy serves as an initial phylogenetic hypothesis which was then combined with a selection of species-level phylogenies [25, 248] in order to better constrain a minimum estimate of the actual phylogenetic relationships of the species. The supertree was inferred via matrix representation parsimony implemented in the `phytools` package for R [259]. Of the two most parsimonious trees found, I used only one for analysis.

Polytomies were resolved in order of species first appearance in order to minimize stratigraphic gaps. The resulting tree was then time scaled using the `paleotree` package via the “minimum branch length” approach with a minimum length of 0.1 My [17]. The minimum length is necessary to avoid zero-length branches which cause the phylogenetic covariance matrix not to be positive definite, which is important for computation (see below). While other time scaling approaches are possible [120, 18] this method was chosen for its simplicity and not requiring additional information about diversification rates which are the interest of this study.

2.5.2 Modeling censored observations

Censored data are modeled using the survival function of the distribution, $S(t)$, defined earlier for the Weibull distribution (Eq. 5, 6) with σ defined as above (Eq. 8, 9). $S(t)$ is the probability that an observation will survive longer than a given time t .

The likelihood of uncensored observations is evaluated as normal using equation 4 while right censored observations are evaluated at $S(t)$ and left censored observations are evaluated at $1 - S(t)$. Note, $1 - S(t)$ is equivalent to the cumulative distribution function and $S(t)$ is equivalent to the complementary cumulative distribution function [101].

The final sampling statement/likelihood for both uncensored and both right and left censored observations is then written

$$L \propto \prod_{i \in C} \text{Weibull}(y_i | \alpha, \sigma) \prod_{j \in R} S(y_j | \alpha, \sigma) \prod_{k \in L} (1 - S(y_k | \alpha, \sigma)),$$

where C is the set of uncensored observations, R is the set of right censored observations, and L is the set of left censored observations.

2.5.3 Deviance residuals

In standard linear regression, residuals are defined as $r_i = y_i - y_i^{est}$. For the model used here, this definition is inadequate. The equivalent values for survival analysis are deviance residuals. To define how deviance residuals are calculated, we first define the cumulative hazard function [160]. Given $S(t)$, we define the cumulative hazard function as

$$\Lambda(t) = -\log(S(t)).$$

Next, we define martingale residuals m as

$$m_i = I_i - \Lambda(t_i).$$

I is the inclusion vector of length n , where $I_i = 1$ means the observation is completely observed and $I_i = 0$ means the observation is censored. Martingale residuals have a mean of 0, range between 1 and $-\infty$, and can be viewed as the difference between the observed number of deaths between 0 and t_i and the expected number of deaths based on the model. However, martingale residuals are asymmetrically distributed, and can not be interpreted in the same manner as standard residuals.

The solution to this is to use the deviance residuals, D . This is defined as a function of martingale residuals and takes the form

$$D_i = \text{sign}(m_i) \sqrt{-2[m_i + I_i \log(I_i - m_i)]}.$$

Deviance residuals have a mean of 0 and a standard deviation of 1 by definition.

2.5.4 Variance partitioning

I calculated VPC using a resampling approach based on [107]. The procedure is as follows:

1. Simulate w (50,000) values of η ; $\eta \sim \mathcal{N}(0, \sigma_c)$.
2. For a given value of $\beta^T \mathbf{X}$, calculate σ^{c*} (Eq. 7) for all w simulations, holding h constant at 0.
3. Calculate v_c , the Weibull variance (Eq. 2.10) of each element of σ^{c*} with α drawn from the posterior estimate.

4. Simulate w values of h ; $h \sim \mathcal{N}(0, \sigma_p)$.
5. For a given value of $\beta^T \mathbf{X}$, calculate σ^{p*} (Eq. 7) for all w simulations, holding η constant at 0.
6. Calculate v_p , the Weibull variance (Eq. 2.10) of each element of σ^{p*} with α drawn from the posterior estimate.
7. $\sigma_{y*}^2 = \frac{1}{2} \left(\left(\frac{1}{w} \sum_i^w v_{pi} \right) + \left(\frac{1}{w} \sum_j^w v_{cj} \right) \right)$.
8. $\sigma_{c*}^2 = var(v_c)$ and $\sigma_{p*}^2 = var(v_p)$.

The simulated values of h were drawn from a univariate normal distribution because each simulated value is in isolation, so there is no concern of phylogenetic autocorrelation. The chosen value for $\beta^T \mathbf{X}$ was a draw from the posterior estimate of the intercept. Because input variables were standardized prior to model fitting, the intercept corresponds to the estimated effect on survival of the sample mean.

Weibull variance is calculated as

$$var(x) = \sigma^2 \left(\Gamma \left(1 + \frac{2}{\alpha} \right) - \left(\Gamma \left(1 + \frac{1}{\alpha} \right) \right)^2 \right), \quad (2.10)$$

where Γ is the gamma function.

The variance partitioning coefficients are then calculated, for example, as $VPC_{phylo} = \frac{\sigma_{p*}^2}{\sigma_{y*}^2 + \sigma_{c*}^2 + \sigma_{p*}^2}$ and similarly for the other components.

2.5.5 Widely applicable information criterion

WAIC can be considered fully Bayesian alternative to the Akaike information criterion, where WAIC acts as an approximation of leave-one-out cross-validation which acts as a measure of out-of-sample predictive accuracy [101]. The following explanation uses the “WAIC 2” formulation recommended by [101].

WAIC is calculated starting with the log pointwise posterior predictive density calculated as

$$\text{lppd} = \sum_{i=1}^n \log \left(\frac{1}{S} \sum_{s=1}^S p(y_i | \Theta^S) \right), \quad (2.11)$$

where n is sample size, S is the number posterior simulation draws, and Θ represents all of the estimated parameters of the model. This is similar to calculating the likelihood of each observation given the entire posterior.

A correction for the effective number of parameters is then added to lppd to adjust for overfitting. The effective number of parameters is calculated, following derivation and recommendations of [101], as

$$p_{\text{WAIC}} = \sum_{i=1}^n V_{s=1}^S (\log p(y_i | \Theta^S)). \quad (2.12)$$

where V is the sample posterior variance of the log predictive density for each data point.

Given both equations 2.11 and 2.12, WAIC is then calculated

$$\text{WAIC} = \text{lppd} - p_{\text{WAIC}}. \quad (2.13)$$

When comparing two or more models, lower WAIC values indicate better out-of-sample predictive accuracy. Importantly, WAIC is just one way of comparing models. When combined

with posterior predictive checks it is possible to get a more complete understanding of model fit.

2.5.6 Results from posterior predictive checks

With all marginal posterior estimates having converged ($\hat{R} < 1.1$) it is possible to examine the quality of model fit (Table 1). If the model is an adequate descriptor of the observed data, then relatively confident inference can be made [101].

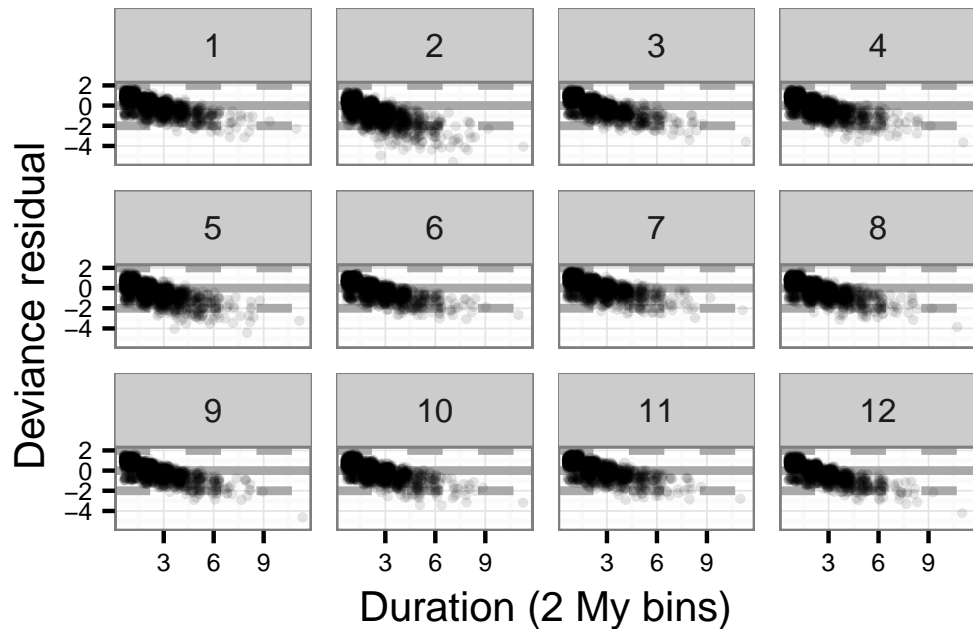


Figure 2.6: Deviance residuals from the fitted survival model compared to observed durations. Each graph depicts the residuals from single draws from the posterior distributions of all estimated parameters. Positive values indicate an underestimate of the observed duration, while negative values indicate an overestimate of the observed duration. A small amount of noise is added to each point to increase clarity. Twelve different examples are provided here to indicate consistency across multiple realizations.

Visual examination of the deviance residuals from twelve different sets of posterior predictive simulations indicates a systematic weakness estimating durations greater than 3 2-My bins (Fig. 2.6). However, the comparison of posterior predictive estimates of the 25th, 50th, and

75th quantiles to the observed indicate adequate fit. (Fig. 2.7). Importantly, this indicates that the model has approximate fit for 50% of the data. Because, the inferred model can be inferred to be approximately adequate at capturing the observed variation.

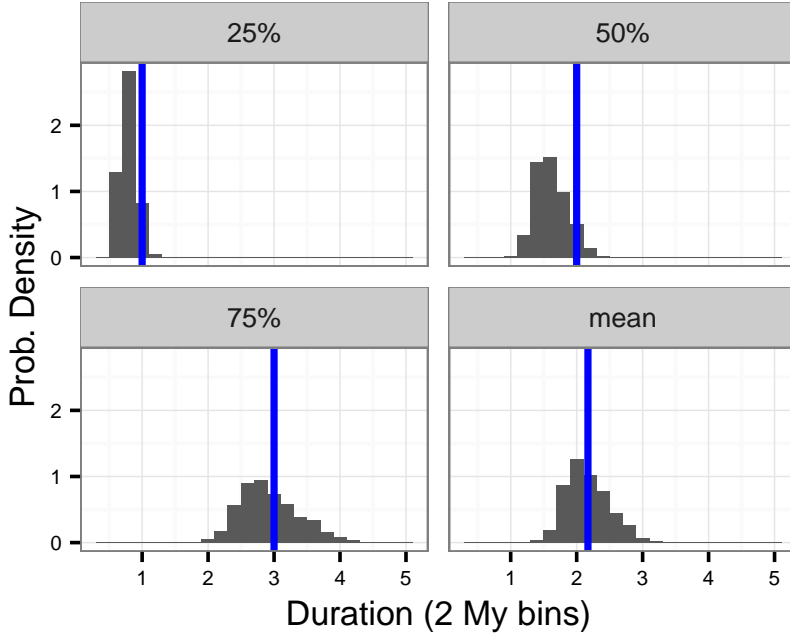


Figure 2.7: The results of additional posterior predictive checks for four summaries of the observed durations, as labeled. Blue vertical lines indicate the observed value. None of the observed values are significantly different from the posterior predictive distributions.

The Weibull model (6140.37) also had a much lower WAIC score than the Exponential model (16697.35). This large a difference indicates that the Weibull model probably has the lower out-of-sample predictive accuracy of the two.

2.5.7 Data quality concerns

A concern with using the PBDB as a primary data source, though this concerns are general to most paleontological data, is that the results are an artifact of taxonomy or the database itself [346]. However, to obtain the results obtained in this analysis there would need to be a

systematic error in assignments of all of diet, locomotor, and taxonomic categories for a large portion of the close to 2000 sampled species. It is important to note that species included have to have body size information, much of which is found from other sources (see Dataset S1). This means that, for many taxa, that species name has to appear in occur in more than one place. This is a strong filter for misspellings and potentially invalid taxa. Additionally, given that most mammal fossils are teeth which allows for relatively accurate dietary category assignment.

A possible concern, however, is that omnivorous taxa have feature poor morphology and thus longer durations may reflect a single anagenetic lineage as opposed to a single “species.” However it is possible to consider that, from a population genetic perspective, it can be argued that a single unbranching lineage is still a single biological “unit.”

2.5.8 Concerns surrounding estimates of α

The estimate of the Weibull shape parameter, α , is greater than 1 meaning that extinction risk is expected to increase with taxon age (Table 1). As the value of α is between 1 and 1.5, extinction risk for a given species only gradually increases with age (Fig. 2.5). There are three possible explanations for this result: 1) older taxa being outcompeted by younger taxa [347]; or 2) this is an artifact of the minimum resolution of the fossil record [284].

An additional concern is that there may be an upward bias in estimates of α at this sample size, similar to that for scale parameters [101]. The plausibility of third possibility in this example can be explored in simulation. I simulated from 10, 100, 1000, and 10000 samples from a Weibull($alpha = 1.3, sigma = 1$) 100 times each. For each of these simulated datasets, I then estimated the values of α and σ in a simple maximum likelihood context in order to just get the model estimate. The modal estimates of both parameters for the simulated datasets

were then compared to the known values (Fig. 2.8). The results from these simulations demonstrate that the estimates of α in the above analyses (Table 1) should not be particularly biased based on my sample size of approximately 2000 species.

The model used in this analysis, however, is unable to distinguish between the remaining two hypotheses [284, 347]. Further work on how to better constrain estimates α is necessary. A possibly is somehow incorporating these hypotheses as prior information.

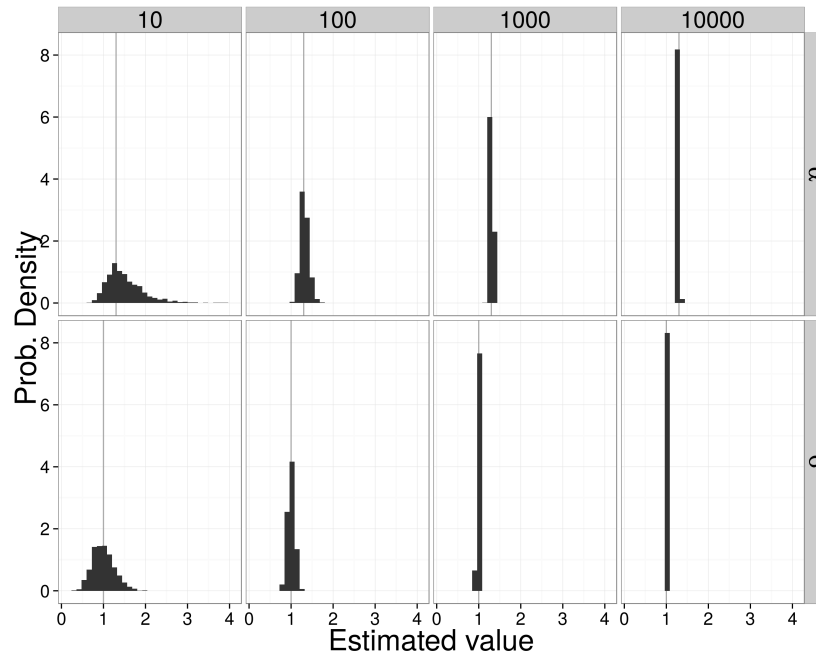


Figure 2.8: Comparison of maximum likelihood estimates of shape (α) and scale (σ) parameters from 1000 simulated data sets from 4 different sample sizes. Vertical lines are the actual parameter value used to generate the data. When sample size is approximately 100 or greater, estimates are not overly biased.

Table 2.1: Marginal posterior estimates for the parameters of interested based on 1000 posterior samples. The intercept β_0 can also be interpreted as the estimate for the mean observed species. The remaining β values can be interpreted as the effect of a trait on the expected species duration as expressed as deviation from the mean. The categorical variables are binary index variables where an observation is of that category or not. See Equation 2.6 for an explanation of the effect of α on extinction risk. \hat{R} values of less than 1.1 indicate approximate chain convergence for the posterior samples.

parameter	effect	mean	sd	2.5%	25%	50%	75%	97.5%	\hat{R}
α	“age”	1.29	0.03	1.23	1.27	1.29	1.31	1.36	1.00
β_0	arboreal/carnivore	-0.78	0.14	-1.05	-0.87	-0.78	-0.68	-0.51	1.00
β_o	occupancy	-0.53	0.08	-0.69	-0.59	-0.53	-0.48	-0.38	1.00
β_{size}	body size	-0.05	0.05	-0.14	-0.08	-0.05	-0.01	0.05	1.00
β_g	ground dwelling	-0.28	0.10	-0.47	-0.34	-0.28	-0.21	-0.09	1.00
β_s	scansorial	-0.22	0.11	-0.43	-0.29	-0.22	-0.14	-0.00	1.00
β_h	herbivore	0.09	0.09	-0.09	0.03	0.09	0.14	0.27	1.00
β_i	insectivore	0.10	0.11	-0.11	0.03	0.10	0.17	0.31	1.00
β_o	omnivore	-0.12	0.11	-0.33	-0.19	-0.12	-0.05	0.09	1.00
σ_c	sd cohort	0.33	0.06	0.23	0.29	0.33	0.37	0.48	1.00
σ_p	sd phylogeny	0.11	0.05	0.03	0.07	0.10	0.14	0.23	1.03

Table 2.2: Species trait assignments in this study are a coarser version of the information available in the PBDB. Information was coarsened to improve per category sample size and uniformity and followed this table.

This study		PBDB categories
Diet	Carnivore	Carnivore
	Herbivore	Browser, folivore, granivore, grazer, herbivore.
	Insectivore	Insectivore.
	Omnivore	Frugivore, omnivore.
Locomotor	Arboreal	Arboreal.
	Ground dwelling	Fossorial, ground dwelling, semifossorial, saltatorial.
	Scansorial	Scansorial.

Table 2.3: Regression equations used in this study for estimating body size. Equations are presented with reference to taxonomic grouping, part name, and reference.

Group	Equation	log(Measurement)	Source
General	$\log(m) = 1.827x + 1.81$	lower m1 area	[165]
General	$\log(m) = 2.9677x - 5.6712$	mandible length	[91]
General	$\log(m) = 3.68x - 3.83$	skull length	[183]
Carnivores	$\log(m) = 2.97x + 1.681$	lower m1 length	[337]
Insectivores	$\log(m) = 1.628x + 1.726$	lower m1 area	[26]
Insectivores	$\log(m) = 1.714x + 0.886$	upper M1 area	[26]
Lagomorph	$\log(m) = 2.671x - 2.671$	lower toothrow area	[329]
Lagomorph	$\log(m) = 4.468x - 3.002$	lower m1 length	[329]
Marsupials	$\log(m) = 3.284x + 1.83$	upper M1 length	[108]
Marsupials	$\log(m) = 1.733x + 1.571$	upper M1 area	[108]
Rodentia	$\log(m) = 1.767x + 2.172$	lower m1 area	[165]
Ungulates	$\log(m) = 1.516x + 3.757$	lower m1 area	[201]
Ungulates	$\log(m) = 3.076x + 2.366$	lower m2 length	[201]
Ungulates	$\log(m) = 1.518x + 2.792$	lower m2 area	[201]
Ungulates	$\log(m) = 3.113x - 1.374$	lower toothrow length	[201]

CHAPTER 3

HOW MACROECOLOGY AFFECTS MACROEVOLUTION: THE INTERPLAY BETWEEN EXTINCTION INTENSITY AND TRAIT-DEPENDENT EXTINCTION IN BRACHIOPODS

As extinction intensity increases, how do the effects of traits on taxonomic survival change? Does the extinction rate associated with certain traits increase while that of others decreases? Using a hierarchical Bayesian approach, I develop a model of how the effects of biological traits on extinction risk can vary with respect to extinction intensity, origination cohort (i.e. time of origination), and in relation to each other. The emergent traits I analyze for each brachiopod genus are geographic range, affinity for epicontinental seas versus open ocean environments, and body size. Additionally, I estimate the effects of environmental generalization versus specialization on taxonomic survival by allowing environmental preference to have a nonlinear effect on duration. My analytical framework eschews the traditional distinction between background and mass extinction, and instead considers extinction intensity as a continuum. I find that the cohort-specific effects of geographic range and environmental preference are negatively correlated with baseline extinction intensity. Additionally, I find support for greater survival of environmental generalists versus specialists in all origination cohorts. These results support the conclusion that for Paleozoic brachiopods, as extinction intensity increases overall extinction selectivity increases.

3.1 Introduction

Extinction is one half of the diversification process [255, 321, 320], second only to speciation or origination; it can also be the ultimate manifestation of selection as a taxon with a

beneficial trait should persist for longer on average than a taxon without that beneficial trait [246, 137, 255, 320].

While estimation of both trait-dependent speciation and extinction rates from phylogenies of extant taxa has grown dramatically [190, 75, 105, 106, 247, 316, 315, 317], there are two major ways to estimate trait-dependent extinction: analysis of phylogenies, and analysis of the fossil record. These two directions, phylogenetic comparative and paleobiological, are complementary and intertwined in the field of macroevolution [246, 137, 131]. In the case of extinction, analysis of the fossil record has the distinct advantage over phylogenies of only extant taxa because extinction is observable; this means that extinction rate is possible to estimate [241, 239, 174]. The approach used here is thus complementary to the analysis of trait-dependent extinction based on a phylogeny.

Jabonksi [134] observed that for bivalves at the end Cretaceous mass extinction event, the effects of some biological traits on taxonomic survival decreased. However, this pattern was not the case for the effect of geographic range on survival [134, 222]. There are multiple possible macroevolutionary mechanisms which may underlie this pattern: the effect of geographic range on survival remains constant and those of other biological traits decrease, the effect of geographic range on survival increases and those of other biological traits stay constant, or the effects of all traits decrease potentially by different degrees.

While Jablonski [134] phrased his conclusions in terms of background versus mass extinction, these states are not distinguishable in terms of extinction rate alone; my analysis treats the time period analyzed as part of the same continuum [349, 222, 296]. Additionally, in order to test the proposed macroevolutionary mechanism behind this scenario [134]; not only do the taxon trait effects need to be modeled, but the correlations between trait effects must be modeled as well.

Here I model brachiopod taxon durations because trait based differences in extinction risk should manifest as differences in taxon durations. Brachiopods are an ideal group for this study as they are well known for having an exceptionally complete fossil record [89, 80]. I focus on the brachiopod record from the post-Cambrian Paleozoic, from the start of the Ordovician (approximately 485 My) through the end Permian (approximately 252 My) as this represents the time of greatest global brachiopod diversity [8] meaning a large sample size for this analysis.

The analysis of taxon durations, or time from origination to extinction, falls under the purview of survival analysis, a field of applied statistics commonly used in health care and engineering [160] and also has a long history in paleontology [298, 299, 335, 336, 54]. I adopt a hierarchical modeling approach [102, 101], which represents both a conceptual and statistical unification of the paleontological dynamic and cohort survival analytic approaches [335, 336, 252, 251, 77, 21, 294, 54, 68].

3.1.1 Factors affecting brachiopod survival

Conceptually, taxon survival can be considered an aspect of “taxon fitness” [49, 220]. Traits associated with taxon survival are thus examples of species (or higher-level) selection, as differences in survival are analogous to differences in fitness. The traits analyzed here are all examples of emergent and aggregate traits [137, 246]; specifically I analyze genus-level traits. Emergent traits are those which are not measurable at a lower level (e.g. species versus individual organism) such as geographic range, or even fossil sampling rate. Aggregate traits, like body size or environmental preference, are the average of a shared trait across all members of a lower level.

Geographic range is widely considered the most important biological trait for estimating

differences in extinction risk at nearly all times, with large geographic range associated with low extinction risk [134, 135, 138, 222, 137, 114, 73]. This stands to reason even if extinction is completely at random; a taxon with an unrestricted range is less likely to go extinct at random than a taxon with a restricted range.

Epicontinental seas are a shallow-marine environment where the ocean has spread over the continental interior or craton with a depth typically less than 100m. In contrast, open-ocean coastline environments have much greater variance in depth, do not cover the continental craton, and can persist during periods of low sea level [203]. Because of this, a simple hypothesis that taxa which favor epicontinental seas would be at great risk during periods of low sea levels, such as during glacial periods, when epicontinental seas are drained. During the Paleozoic (approximately 541–252 My), epicontinental seas were widely spread globally but declined over the Mesozoic (approximately 252–66 My) and have nearly disappeared during the Cenozoic (approximately 66–0 My) as open-ocean coastlines became the dominant shallow-marine setting [286, 226, 203, 155]. Taxa in epicontinental environments could also have a greater extinction susceptibility than taxa in open-ocean environments due to anoxic events due to enhanced water stratification or poor water circulation [225].

Miller and Foote [203] demonstrated that during several mass extinctions taxa associated with open-ocean environments tend to have a greater extinction risk than those taxa associated with epicontinental seas. During periods of background extinction, however, they found no consistent difference between taxa favoring either environment. Miller and Foote [203] hypothesize that open-ocean taxa may have a greater extinction rate because these environments would be more strongly affected by waterborne hazards such as fallout from impacts or volcanic events which would propagate more quickly than in epicontinental environments with sluggish circulation. These two environment types represent the primary identifiable environmental dichotomy observed in ancient marine systems [203, 286]. Given these findings,

I would hypothesize that as extinction risk increases, the extinction risk associated with open-ocean environments should generally increase.

Because environmental preference is defined here as the continuum between occurring exclusively in open-ocean environments versus epicontinental environments, intermediate values are considered “generalists” in the sense that they favor neither end member. A long-standing hypothesis is that generalists or unspecialized taxa will have greater survival than specialists [298, 169, 170, 217, 218, 21, 309]. Because of this, the effect of environmental preference was modeled as a quadratic function where a concave down relationship between preference and expected duration indicates that generalists are favored over specialists end-members.

Body size, measured as shell length, is also considered as a trait that may potentially influence extinction risk [223, 115]. Body size is a proxy for metabolic activity and other correlated life history traits [223]. Harnik *et al.* [116] analyzed the effect of body size selectivity in Devonian brachiopods in both a phylogenetic and non-phylogenetic context; finding that body size was not found to be associated with differences in taxonomic duration. It has also been found that, at least in the case of some bivalve subclades, body size can be as important a factor as geographic range size in determining extinction risk [115]. Given these results, I expect that if body size has any effect on brachiopod taxonomic survival it is very small.

It is well known that, given the incompleteness of the fossil record, the observed duration of a taxon is an underestimate of that taxon’s true duration [313, 348, 351, 173, 9, 89]. Because of this, the concern is that a taxon’s observed duration may reflect its relative chance of being sampled and not any of the effects of the covariates of interest. In this case, for sampling to be a confounding factor there must be consistent relationship between the quality of sampling of a taxon and its apparent duration (e.g. greater sampling, longer duration). If there is no relationship between sampling and duration then interpretation can be made clearly; while observed durations are obviously truncated true durations, a lack of a relationship would

indicate that the amount and form of this truncation is not a major determinant of the taxon’s apparent duration. By including sampling as a covariate in the model, this effect can be quantified and can be taken into account when interpreting the estimates of the effects of the other covariates.

3.2 Materials and Methods

3.2.1 Fossil occurrence information

The brachiopod dataset analyzed here was sourced from the Paleobiology Database (<http://www.paleodb.org>) which was then filtered based on taxonomic (Rhynchonelliformea: Rhynchonellata, Chileata, Obolellida, Kutorginida, Strophomenida, Spiriferida), temporal (post-Cambrian Paleozoic), stratigraphic, and other occurrence information used in this analysis. Analyzed occurrences were restricted to those with paleolatitude and paleolongitude coordinates, assignment to either epicontinental or open-ocean environment, and belonging to a genus present in the body size dataset [223]. Epicontinental versus open-ocean assignments for each fossil occurrence are partially based on those analyzed by Miller and Foote [203], with additional occurrences assigned similarly (Miller and Foote, personal communication). These filtering criteria are very similar to those used by Miller and Foote [88] with an additional constraint of being present in the body size data set from Payne *et al* [223]. In total, 1130 genera included in the dataset.

Fossil occurrences were analyzed at the genus level which is common for paleobiological, macroevolutionary and macroecological studies of marine invertebrates [8, 88, 114, 158, 203, 217, 218, 222, 296, 341]. While species diversity dynamics are frequently of much greater interest than those of higher taxa, though see [85, 123], the nature of the fossil record makes

accurate, precise, and consistent taxonomic assignments at the species level difficult for all occurrences. As such, the choice to analyze genera as opposed to species was in order to assure a minimum level of confidence and accuracy in the data analyzed here.

Genus duration was calculated as the number of geologic stages from first appearance to last appearance, inclusive. Durations were based on geologic stages as opposed to millions of years because of the inherently discrete nature of the fossil record; dates are not assigned to individual fossils themselves but instead fossils are assigned to a geological interval which represents some temporal range. In this analysis, stages are effectively irreducible temporal intervals in which taxa may occur. Genera with a last occurrence in or after Changhsingian stage (e.g. the final stage of the study interval) were right censored at the Changhsingian; genera with a duration of only one stage were left censored [160]. How the likelihood of censored observations is calculated is detailed in section 3.2.2.

The covariates of duration included in this analysis are geographic range size (r), environmental preference (v, v^2), body size (m), and sampling (s).

Geographic range was calculated as relative occupancy corrected for incomplete sampling. First, the paleolatitude-paleolongitude coordinates for all occurrences were projected onto an equal-area cylindrical map projection. Each occurrence was then assigned to one of the cells from a 70×34 regular raster grid placed on the map. Each grid cell represents approximately $250,000 \text{ km}^2$. The map projection and regular lattice were made using shape files from <http://www.naturalearthdata.com/> and the `raster` package for R [122]. For each stage, the total number of occupied grid cells was calculated. Then, for each temporal bin, the relative occurrence probability of the observed taxa was calculated using the JADE method developed by Chao *et al.* [42]. This method accounts for the fact that taxa with an occupancy of 0 cannot be observed which means that occupancy follows a truncated Binomial distribution. This correction is critical when comparing occupancies from different times with different

geographic sampling. Finally, for each genus, the mean relative occurrence probability was calculated as the average of that genus' occurrence probabilities for all stages it was sampled to yield relative occupancy, my proxy for geographic range.

Environmental preference was defined as probability of observing the ratio of epicontinental occurrences to total occurrences ($\theta_i = e_i/E_i$) or greater given the background occurrence probability θ'_i as estimated from all other taxa occurring at the same time (e'_i/E'_i). This measure of environmental preference is expressed.

$$\begin{aligned} p(\theta'_i | e'_i, E'_i) &\propto \text{Beta}(e'_i, E'_i - e'_i)\text{Beta}(1, 1) \\ &= \text{Beta}(e'_i + 1, E'_i - e'_i + 1), \end{aligned} \tag{3.1}$$

where v is the percent of the distribution defined in equation 3.1 less than or equal to θ_i . The Beta distribution is used here because it is a continuous distribution bounded at 0 and 1, which is ideal for modeling percentages.

Body size, measured as shell length, was sourced directly from Payne *et al.* [223]. These measurements were made from brachiopod taxa figured in the *Treatise on Invertebrate Paleontology* [359].

The sampling probability for individual taxa was calculated using the standard gap statistic [89, 79]. The gap statistic is calculated as the number of stages in which the taxon was sampled minus two divided by the duration of the taxon minus two. Subtracting two from both the numerator and denominator is because the first and last appearance stages are by definition sampled. Because taxa that were right censored only include a first appearance, one was subtracted from the numerator and denominator instead of two.

The minimum duration for which a gap statistic can be calculated is three stages, so I chose to impute the gap statistic for all observations with a duration less than 3. Imputation

is the “filling in” of missing observations based on the observed values [273, 102]. This is fairly straightforward in a Bayesian framework because both covariates and parameters are considered random variables, meaning that the missing values of sampling can be modeled as coming from some probability distribution. The model for imputing sampling probability is described in section 3.2.3.

Prior to analysis, geographic range was logit transformed and body size was natural-log transformed; both of these transformations make these variables defined for the entire real line. Sampling probability was transformed as $(s(n - 1) + 0.5)/n$ where n is the sample size as recommended by Smithson and Verkuilen [308]; this serves to slightly shrink the range of the data so that there are no values of 0 or 1. All covariates except for sampling were standardized by subtracting the mean from all values and dividing by twice its standard deviation [102]. This standardization means that the associated regression coefficients are comparable as the expected change per 1-unit change in the rescaled covariates. Finally, D is defined as the total number of covariates, excluding sampling, plus one for the intercept term.

3.2.2 Details of model

Hierarchical modelling is a statistical approach which explicitly takes into account the structure of the observed data in order to model both the within and between group variance [101, 102]. The units of study (e.g. genera) each belong to a single group (e.g. origination cohort). Each group is considered a draw from a shared probability distribution (e.g. prior) of all cohorts, observed and unobserved. The group-level parameters, or the hyperparameters of this shared prior, are themselves given (hyper)prior distributions and are also estimated like the other parameters of interest (e.g. covariate effects) [101]. The subsequent estimates are partially pooled together, where parameters from groups with large samples or effects remain large while those of groups with small samples or effects are pulled towards the overall group

mean. All covariate effects (regression coefficients), as well as the intercept term (baseline extinction risk), were allowed to vary by group (origination cohort). The covariance between covariate effects was also modeled.

Genus durations were assumed to follow a Weibull distribution which allows for age-dependent extinction [160]: $y \sim \text{Weibull}(\alpha, \sigma)$. The Weibull distribution has two parameters: scale σ , and shape α . When $\alpha = 1$, σ is equal to the expected duration of any taxon. α is a measure of the effect of age on extinction risk where values greater than 1 indicate that extinction risk increases with age, and values less than 1 indicate that extinction risk decreases with age. Note that the Weibull distribution is equivalent to the exponential distribution when $\alpha = 1$.

In the case of the right- and left-censored observations mentioned above, the probability of those observations has a different calculation [160]. For right-censored observations, the likelihood is calculated $p(y|\theta) = 1 - F(y) = S(y)$ where $F(y)$ is the cumulative distribution function. In contrast, the likelihood of a left-censored observation is calculated $p(y|\theta) = F(y)$.

The scale parameter σ was modeled as a regression with both varying intercept and varying slopes and the effect of sampling [161]; this is expressed

$$\sigma_i = \exp\left(\frac{-\mathbf{X}_i B_{j[i]} + \delta s_i}{\alpha}\right) \quad (3.2)$$

where i indexes across all observations where $i = 1, \dots, n$ where n is the total number of observations, $j[i]$ is the cohort membership of the i th observation where $j = 1, \dots, J$ where J is the total number of cohorts, X is a $N \times D$ matrix of covariates along with a column of 1's for the intercept term, B is a $J \times D$ matrix of cohort-specific regression coefficients, and δ is the regression coefficient for the effect of sampling s . δ does not vary by cohort.

Each of the rows of matrix B are modeled as realizations from a multivariate normal distribution with length D location vector μ and $J \times J$ covariance matrix Σ : $B_j \sim \text{MVN}(\mu, \Sigma)$.

The covariance matrix was then decomposed into a length J vector of scales τ and a $J \times J$ correlation matrix Ω , defined $\Sigma = \text{diag}(\tau)\Omega\text{diag}(\tau)$ where “diag” indicates a diagonal matrix.

The elements of μ were given independent normally distributed priors. The effects of geographic range size and the breadth of environmental preference were given informative priors reflecting the previous findings while the others were given weakly informative favoring no effect. The correlation matrix Ω was given an LKJ distributed prior [166] that slightly favors an identity matrix [319]. These priors are defined

$$\begin{aligned}\mu^0 &\sim \mathcal{N}(0, 5) \\ \mu^r &\sim \mathcal{N}(-1, 1) \\ \mu^v &\sim \mathcal{N}(0, 1) \\ \mu^{v^2} &\sim \mathcal{N}(1, 1) \\ \mu^m &\sim \mathcal{N}(0, 1) \\ \tau &\sim C^+(1) \\ \Omega &\sim \text{LKJ}(2).\end{aligned}\tag{3.3}$$

The log of the shape parameter α was given a weakly informative prior $\log(\alpha) \sim \mathcal{N}(0, 1)$ centered at $\alpha = 1$, which corresponds to the Law of Constant Extinction [335].

3.2.3 Imputation of sampling probability

The vector of sampling values s has two parts: the observed part s^o , and the unobserved part s^u . Of the 1130 total observations, 539 have a duration of 3 or more and have an observed gap statistic. The gap statistic for the remaining 591 observations was imputed. As stated above, the unobserved part is the imputed, or filled in, based on the observed part

s^o . Because sampling varies between 0 and 1, I chose to model it as a Beta regression with matrix W being a $N \times (D - 1)$ matrix of covariates (i.e. geographic range size, environmental preference, body size) as predictors of sampling; this assumes that the sampling value for all taxa come from the same distribution. Importantly, I make no assumptions of causal structure.

Predicting sampling probability using the other covariates that are then included in the model of duration is acceptable and appropriate in the case of imputation where the sample goal is accurate prediction [273, 102]. Not including these covariates can lead to biased estimates of the imputed variable; if the covariates themselves are related, not including them will bias this correlation towards zero which then leads to improper imputation and inference [273].

The Beta regression is defined

$$s^o \sim \text{Beta}(\phi = \text{logit}^{-1}(X^o\gamma), \lambda), \quad (3.4)$$

where γ is a length D vector of regression coefficients, and X defined as above. The Beta distribution used in the regression is reparameterized in terms of a mean parameter

$$\phi = \frac{\alpha}{\alpha + \beta} \quad (3.5)$$

and total count parameter

$$\lambda = \alpha + \beta \quad (3.6)$$

where α and β are the characteristic parameters of the Beta distribution [101].

The next step is to then estimate $s^u|s^o, X^o, X^u, \gamma$, the posterior distribution of which are folded back into s and used as a covariate of duration (Eq. 3.2). All the elements of γ , and

both δ (Eq. 3.2) and λ (Eq. 3.4) were given weakly informative priors where

$$\begin{aligned}\gamma &\sim \mathcal{N}(0, 1) \\ \delta &\sim \mathcal{N}(0, 1) \\ \lambda &\sim \text{Pareto}(0.1, 1.5).\end{aligned}\tag{3.7}$$

3.2.4 Posterior inference and posterior predictive checks

The joint posterior was approximated using a Markov-chain Monte Carlo routine that is a variant of Hamiltonian Monte Carlo called the No-U-Turn Sampler [125] as implemented in the probabilistic programming language Stan [319]. The posterior distribution was approximated from four parallel chains run for 10,000 steps each, split half warm-up and half sampling and thinned to every 10th sample for a total of 4000 posterior samples. Chain convergence was assessed via the scale reduction factor \hat{R} where values close to 1 ($\hat{R} < 1.1$) indicate approximate convergence. Convergence means that the chains are approximately stationary and the samples are well mixed [101].

Model adequacy was evaluated using a couple of posterior predictive checks. Posterior predictive checks are a means for understanding model fit or adequacy where the basic idea is that replicated data sets simulated from the fitted model should be similar to the original data and systematic differences between the simulations and observations indicate weaknesses of the model fit [101]. For both approaches used here, each posterior predictive datasets were generated from a unique draw from the posterior distribution of each parameter. The two posterior predictive checks used in this analysis are a comparison of a non-parametric estimate of the survival function $S(t)$ from the empirical dataset to the non-parametric estimates of $S(t)$ from the 100 posterior predictive datasets, and comparison of the observed genus durations to the average posterior predictive estimate of $\log(\sigma)$ (Eq. 3.2). The former is to

see if simulated data has a similar survival pattern to the observed, while the latter is to see if the model systematically over- or under- estimates taxon survival.

3.3 Results

Comparison of the posterior predictive estimates of $S(t)$ to the empirical estimate reveal few obvious biases except for the case of values from the far right tail of observed durations (Fig. 3.1). This result is reinforced by the additional posterior predictive comparison where most estimates are not systematically biased except for a consistent under-estimate of $\log(\sigma)$ for older taxa (Fig. 3.2). The results of both posterior predictive checks indicate that, for the majority of observations, model fit is generally not biased.

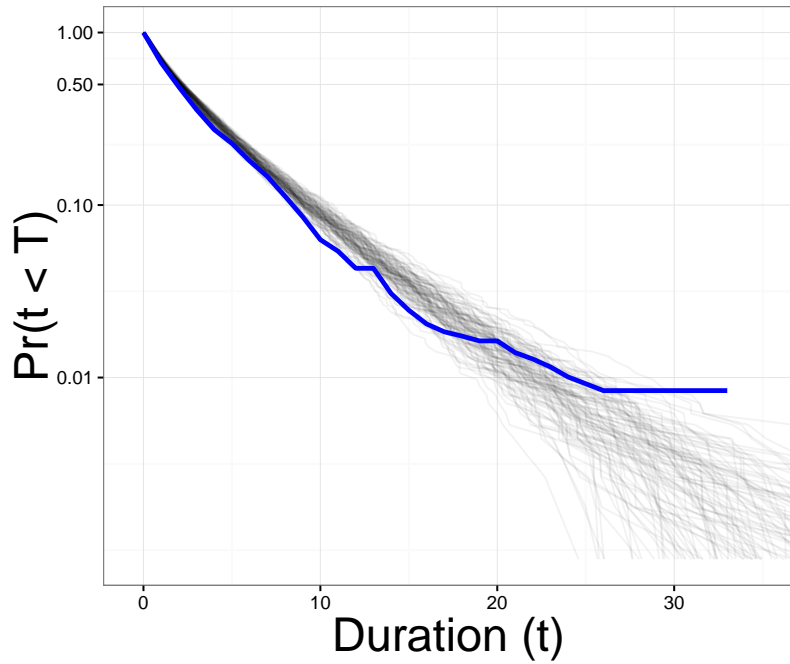


Figure 3.1: Comparison of the empirical estimate of $S(t)$ (highlighted) versus estimates from 100 posterior predictive data sets (black). $S(t)$ corresponds to the probability that the age of a genus t is less than the genus' ultimate duration T . The vertical axis is log10 transformed.

The cohort-level estimate of the effect of geographic range size indicates that as a taxon's

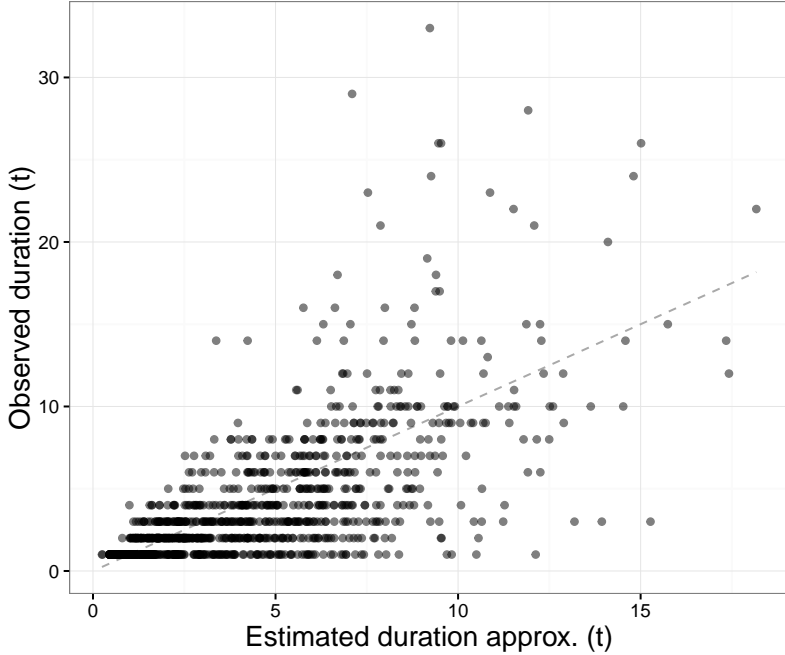


Figure 3.2: Comparison of all observed genus durations in number of geological stages to the average posterior predictive estimates of $\log(\sigma)$. The dashed, diagonal line corresponds to $x = y$.

geographic range increases, that taxon's duration is expected to increase (Table 3.1). Given the estimates of μ^r and τ^r , there is a less than 3.7% ($\pm 0.04\%$ SD) probability that this relationships would be reversed ($\Pr \mathcal{N}(\mu^r, \tau^r) > 0$). The between-cohort variance τ^r is the lowest of all the regression coefficients (Table 3.1).

Body size is estimated to have no effect on taxon duration, with the estimate being nearly 0 (Table 3.1). The variance between the cohort-specific estimates of the effect of body size τ^m is estimated to be greater than the variance of between-cohort estimates of the effect of geographic range size τ^r .

The group-level estimate of the effect of environmental preference is estimated from both μ^v and μ^{v^2} .

The estimate of μ^v indicates that epicontinental favoring taxa are expected to have a greater

duration than open-ocean favoring taxa (Table 3.1). Additionally, given the estimate of between-cohort variance τ^v , there is approximately 18% ($\pm 7\%$ SD) probability that, for any given cohort, taxa favoring open-ocean environments would have a greater expected duration than taxa favoring epicontinental environments ($\Pr(\mathcal{N}(\mu^v, \tau^v) > 0)$).

The estimate of μ^{v^2} indicates that the overall relationship between environmental preference and $\log(\sigma)$ is concave down (Fig. 3.3), with only a 2.7% ($\pm 3\%$ SD) probability that any given cohort is convex up ($\Pr(\mathcal{N}(\mu^{v^2}, \tau^{v^2}) < 0)$).

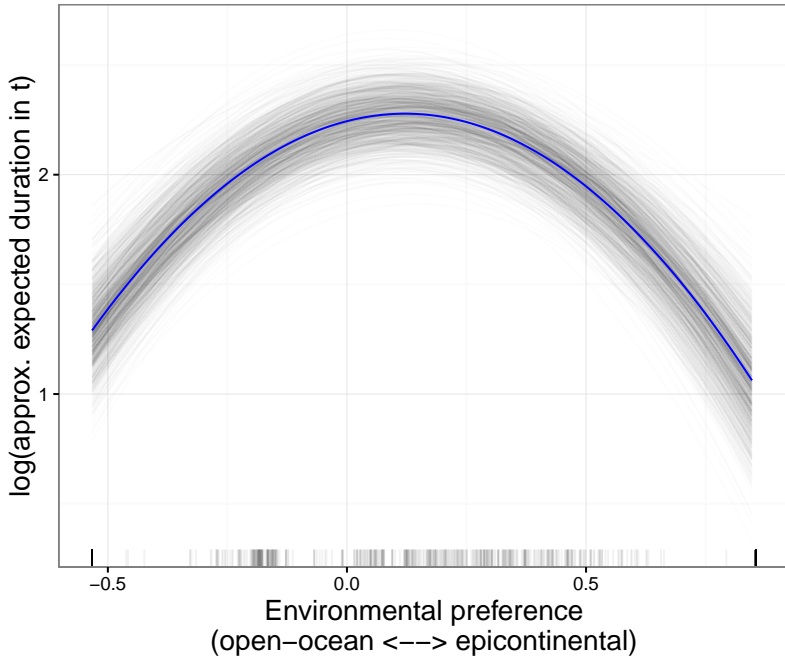


Figure 3.3: The overall expected relationship between environmental affinity v_i and a $\log(\sigma)$ when $r = 0$ and $m = 0$. The 1000 semi-transparent lines corresponds to a single draw from the posterior predictive distribution, while the highlighted line corresponds to the median of the posterior predictive distribution. The overall relationship is concave down with an optimum greater than 0, which means that taxa favoring epicontinental environments are expected to have longer durations than those favoring open-ocean environments. The tick marks along the bottom of the plot correspond to the (rescaled) observed values of environmental preference.

The cohort-specific estimates of all the regression coefficients demonstrate a lot of between cohort variance, with no obvious trends. As indicated in Table 3.1 and detectable visually (Fig. 3.4), the between-cohort estimates for β^0 , β^r , and β^m all have much lower variance

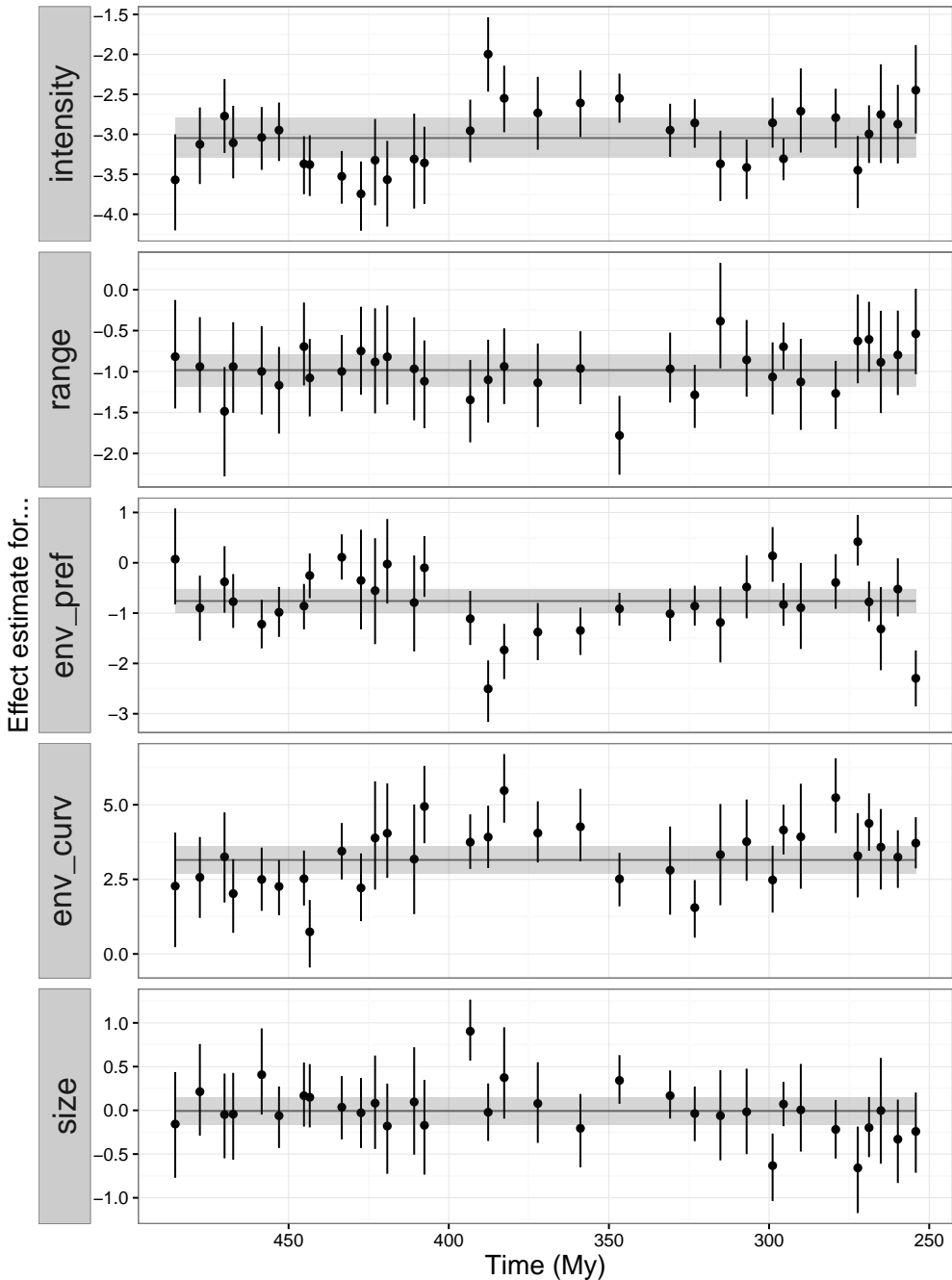


Figure 3.4: Comparison of cohort-specific estimates of β^0 , the effect of geographic range on extinction risk β^r , the effect of environmental preference β^v and β^{v^2} , and body size β^m . Points correspond to the median of the cohort-specific estimate, along with 80% credible intervals. Points are plotted at the midpoint of the cohorts stage of origination in millions of years before present (My). Black, horizontal lines are the overall estimates of covariate effects along with 80% credible intervals (shaded).

than the between-cohort estimates of both β^v and β^{v^2} .

While most cohort-specific estimates are very similar to the overall cohort-level estimate, there are a few notable excursions away from the overall mean (Fig. 3.4). There are simultaneous excursions in both β^0 and β^v for cohorts originating in the Givetian (387-382 My) and Frasnian (382-372 My) stages; both of which directly precede the late Devonian mass extinction event at the Frasnian/Famennian boundary. These cohorts are marked by both a high extinction intensity and an increase in expected duration for taxa favoring epicontinental environments over open-ocean ones.

Cohorts originating from the Silurian through the Early Devonian have a slightly lower extinction intensity than the overall mean; these cohorts are those originating in the Llandovery (443-443 My) through the Emsian (407-393 My). These time periods are when there is the lowest overall probability that epicontinental favoring taxa are expected to have greater duration than open-ocean favoring taxa. Both the Silurian and Devonian periods are notable for having been periods with a mostly “hothouse” climate, with no polar icecaps and a high sea-level [62, 153, 210].

The cohort-specific relationships between environmental preference and $\log(\sigma)$ were calculated from the estimates of β^0 , β^v , and β^{v^2} (Fig. 3.5) and reflect how these three parameters act in concert and not just individually (Fig. 3.4). Beyond results already discussed above in the context of the parameters individually, it is notable that the cohort originating in the Kungurian (279-272 My) is least like the overall expected relationship and has the most sharply curved appearance due to a high estimate β^{v^2} (Fig. 3.4). This cohort has the biggest difference in extinction risk between environmental generalists and specialists. The cohorts originating during the Emsian (407-393 My) and Frasnian (382 - 372 My) are tied for second in sharpness of curvature. The least sharply curved cohorts include those originating during

Tremadocian (484-477 My), Hirnantian (445-443 My), Llandovery (443-433 My), and Ludlow (427-423 My). Except for the Tremadocian cohort, most of these cohorts originate during the Silurian through the Early Devonian range identified earlier as having lower expected extinction intensity than what is expected from the group-level estimate.

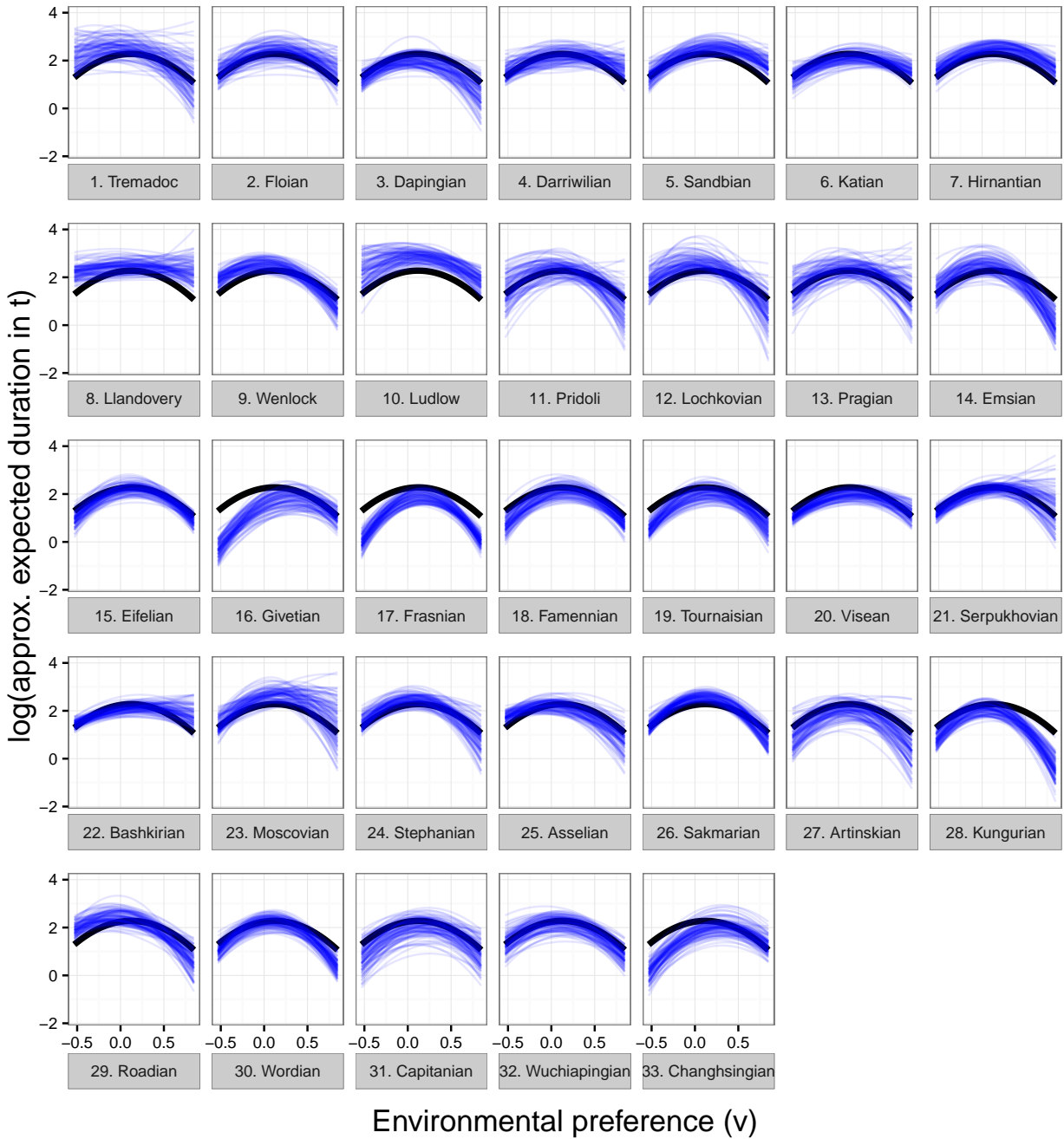
The correlations of the cohort-specific estimates of the regression coefficients are estimated as the off-diagonal elements of the correlation matrix Ω . Only two of the elements of Ω are distinguishable from 0: the correlation of β^0 (extinction intensity) with both β^r and β^v (Fig. 3.6).

There is an approximate 90% probability that the cohort-specific estimates of baseline extinction intensity β^0 and the effect of geographic range β^r are negatively correlated; this means that for cohorts experiencing a lower extinction intensity (β^0 decreases), the magnitude of the effect of geographic range is expected to decrease as well, and *vice versa*; this is in contrast to the observation made by Jablonski [134] with regards to late Cretaceous bivalves.

Similarly, there is an approximate 97.4% probability that the cohort-specific estimates of β^0 and β^v are negatively correlated; this means that as extinction intensity increases it is expected that epicontinental taxa become more favored over open-ocean environments (i.e. as β^0 increases, β^v decreases).

There is only an approximate 30% probability that β^r and β^v are positively correlated. This lack of cross-correlation may be due in part to the much higher between-cohort variance of the effect of environmental preference τ^v than the very small between-cohort variance in the effect of geographic range τ^r (Table 3.1); the effect of geographic range might simply not vary enough relative to the much noisier environmental preference.

Comparison of observed values of sampling, as measured by the gap statistic, to random draws from the posterior estimates of the imputed sampling values indicate that they are



Environmental preference (v)

Figure 3.5: Comparison of origination cohort-specific (posterior predictive) estimates of the effect of environmental preference on $\log(\sigma)$ to the mean overall estimate of the effect of environmental preference. Cohort-specific estimates are from 100 posterior predictive simulations across the range of (transformed and rescaled) observed values of environmental preference. The oldest cohort is in the top-right and younger cohorts proceed left to right, with the youngest cohort being the right-most facet of the last row. Panel names correspond to the name of the stage in which that cohort originated.

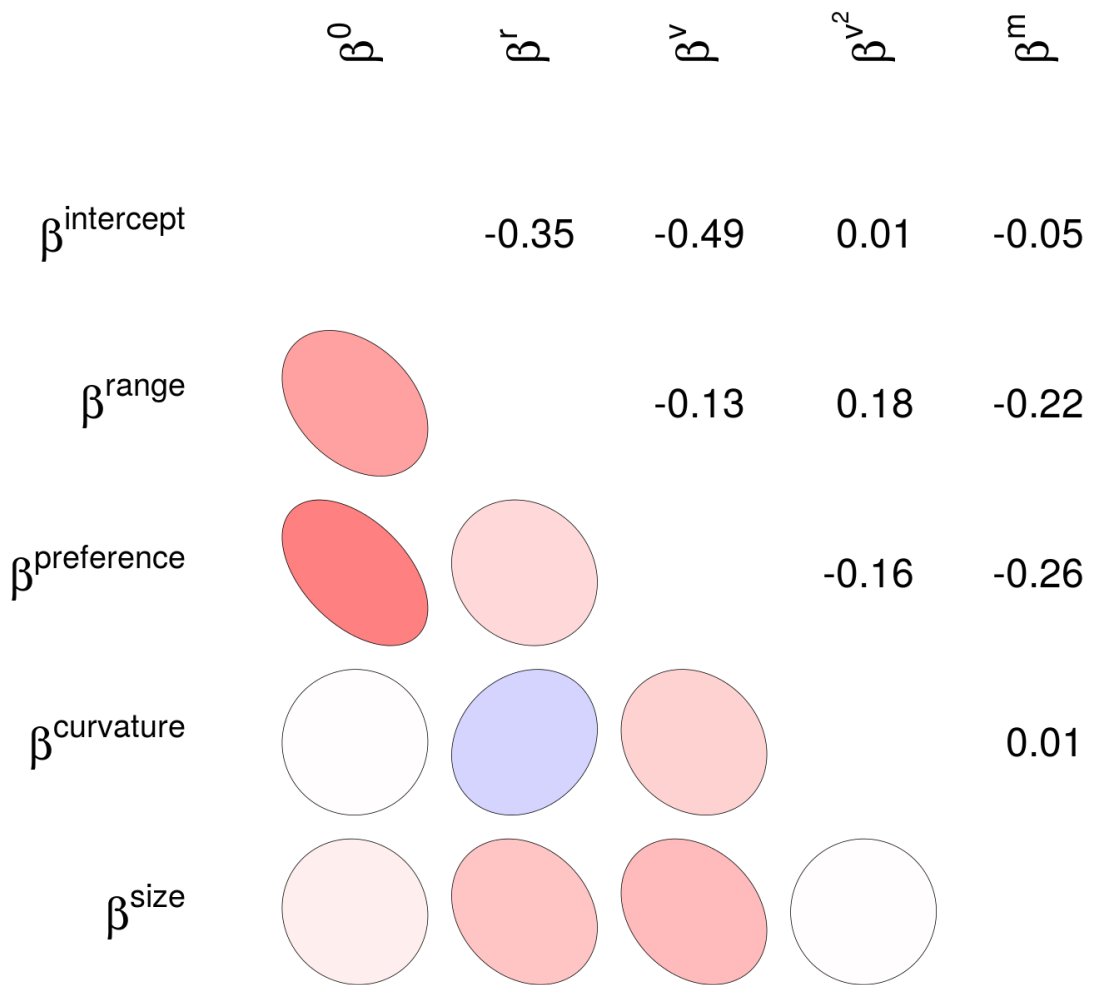


Figure 3.6: Mixed graphical and numerical representation of the correlation matrix Ω of variation in cohort-specific covariate estimates. These correlations are between the estimates of the cohort-level effects of covariates, along with intercept/baseline extinction risk. The median estimates of the correlations are presented numerically (upper-triangle) and as idealized ellipses representing that much correlation (lower-triangle). The darkness of the ellipse corresponds to the magnitude of the correlation.

very similar (Fig. 3.7). This result is very encouraging as this is the ultimate goal of multiple imputation: to fill in missing data with values similar to the observed while taking into account the randomness of that variable [273, 102]. The estimates of δ are based on the set of observed values and the entire posterior of imputed values.

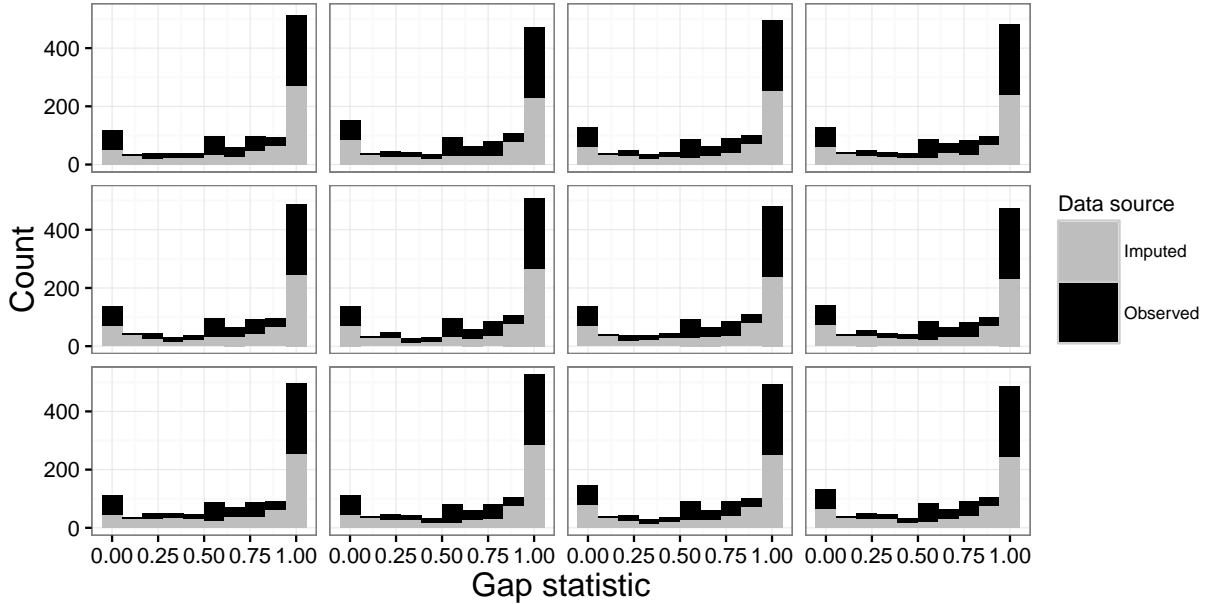


Figure 3.7: Histograms of the distribution of gap statistic values from both the observed values and the imputed values from 12 unique posterior realizations. For each panel the observed values are identical but the imputed values are from a single set of their posterior estimates.

Sampling was found to have a negative effect (positive δ) on duration: greater sampling, shorter duration (Table 3.1). While potentially counter intuitive, this result is most likely due to some long lived taxa only be sampled in the stages of the first and last appearance. Also, longer lived taxa have more opportunities to not be sampled than shorter lived taxa. These two factors will lead to this result.

While the effect of sampling appears large compared to the other regression coefficients, this is only because sampling was not standardized like the other covariates. To make the coefficients comparable, δ is multiplied by twice the posterior mean of the standard deviation

of sampling probability; the transformed value of δ is then 0.642 (± 0.1 SD). This effect is relatively small compared to the other covariate effects (Table 3.1). There is then a 98.6% probability that the effect of geographic range μ^r has a greater magnitude than δ . Similarly, μ^v has a 71.8% probability of having a greater magnitude of effect than δ . Finally, μ^{v^2} has a 100% probability of having a greater magnitude of effect than δ .

The Weibull shape parameter α was found to be approximately 1.36 (± 0.05 SD) with a 100% probability of being greater than 1. This result is not consistent with the Law of Constant Extinction [335] and is instead consistent with accelerating extinction risk with taxon age. This may indicate that older taxa are out-competed by younger taxa, a result consistent with some empirical results [347, 240, 309] and (ironically) with a recently proposed Red Queen-like model of evolution [267]. This results, however, is not consistent with other empirical results [74, 54] and could potentially be caused by the minimum resolution of the fossil record [284]. It is thus unclear if a strong biological inference can be made from this result, which means that further work is necessary on the effect of taxon age on extinction risk.

3.4 Discussion

The generating observation behind this study was that for bivalves at the end Cretaceous mass extinction event, the only biological trait that was found to effect extinction risk was geographic range while traits that had previously been beneficial had no effect [134]. This observation raises two linked questions: how does the effect of geographic range change with changing extinction intensity, and how does the effect of other biological traits change with changing extinction intensity?

I find that as intensity increases (β^0 decreases), the magnitude of the effect of geographic

range increases. I also find that as intensity increases, the effect of favoring epicontinental environments of open-ocean environments is expected to be increase; this is consistent with the results obtained by Miller and Foote [203]. There is no evidence for a correlation between the effect of geographic range and environmental preference. Additionally, the between-cohort variance in effect of geographic range is much less than the between-cohort variance of the effect of environmental preference which may underlie the lack of correlation between these two effects.

Additionally, the lower between-cohort variance in the effect of geographic range versus that higher between-cohort variance implies that for cohorts with a greater than average extinction intensity, the difference in the effect geographic range and the group-level effect of geographic range is expected to be smaller than the difference between the effect of environmental preference and the group-level effect of environmental preference.

I find consistent support for the “survival of the unspecialized,” with respect to epicontinental versus open-ocean environmental preference, as a time-invariant generalization of brachiopod survival; taxa with intermediate environmental preferences are expected to have lower extinction risk than taxa specializing in either epicontinental or open-ocean environments (Fig. 3.3), though the curvature of the relationship varies from rather shallow to very peaked (Fig. 3.5). However, this relationship is not symmetric about 0, as taxa favoring epicontinental environments are expected to have a greater duration than taxa favoring open-ocean environments. This description of environment only describes one major aspect of a taxon’s environmental context, with factors such as bathymetry and temperature being further descriptors of a taxon’s adaptive zone [217, 114, 115, 121]; inclusion of these factors in future analyses would potentially improve our understanding of the “survival of the unspecialized” hypothesis [298].

Hopkins *et al.* [127], in their analysis of niche conservatism and substrate lithological preference

in marine invertebrates, found that brachiopods were among the least “conservative” groups; taxa were found to easily change substrate preference on short time scales. While substrate preference is not the same as environmental preference (as defined here), a question does arise: are there three classes of environmental preference instead of two? These classes would be taxa with broad tolerance (“true” generalists), inflexible specialists (“true” specialists), and flexible but with a narrow tolerance. A flexible taxon is one with a narrow habitat preference at one time, but with preference that changes over time with changing environmental availability. My analysis assumes that traits are constant over the duration of the taxon meaning that this scenario is not detectable; taxa with broad tolerances and flexible taxa with narrow per-stage preference end up being treated the same way. Future work should explore how environmental preference changes over lineage duration in relation to environmental availability to estimate if the generalists–specialists continuum is actually ternary relationship.

An alternative approach for specifically modeling survival that can take into account imperfect observation than the method used here is the Cormack-Jolly-Seber (CJS) model [271, 171, 329, 173]. This model is a type of hidden Markov model with an absorbing state (i.e. extinction). In this model, survival is defined as the probability of surviving from time t to time $t + 1$. Additionally, the effect of preservation and sighting is estimated as probability of observing a taxon that is present; this can extend the duration of a taxon beyond its last occurrence. This approach is a fundamentally different from the method used in my analysis: I am estimating the biasing effect of sampling probability on taxon duration to then compare with effects of other covariates, while the CJS model estimates the pre-sampling fossil record and then estimates per-time unit survival probability.

The use of genera as the unit of the study and how to exactly interpret the effects of the biological traits is an important question. For example, if any of the traits analyzed here are associated with increases in speciation rates, this might increase the duration of genera

through self-renewal [254, 255], which would be an example of the difference in biological pattern between species and genera [135, 136, 137]. This could lead to a trait appearing to decrease generic level extinction risk by that trait increasing species level origination rate instead of decreasing species level extinction risk.

The model used here could be improved through either increasing the number of analyzed traits, expanding the hierarchical structure of the model to include other major taxonomic groups of interest, and the inclusion of explicit phylogenetic relationships between the taxa in the model as an additional hierarchical effect. An example trait that may be of particular interest is the affixing strategy or method of interaction with the substrate of the taxon, which has been found to be related to brachiopod survival where, for cosmopolitan taxa, taxa that are attached to the substrate are expected to have a greater duration than those that are not [3].

It is theoretically possible to expand this model to allow for comparisons both within and between major taxonomic groups which would better constrain the brachiopod estimates while also allowing for estimation of similarities and differences in cross-taxonomic patterns. The major issue surrounding this particular expansion involves finding a similarly well sampled taxonomic group that is present during the Paleozoic. Potential groups include Crinoidea, Ostracoda, and other members of the “Paleozoic fauna” [285].

With significant updates, it would also be possible to compare the brachiopod record with with Modern groups such as bivalves or gastropods [285], though remembering that the groups may not necessarily share all cohorts with the brachiopods. This particular model expansion would act as a test of any universal cross-taxonomic patterns in the effects of emergent traits on extinction such as has been proposed for geographic range [222]. Additionally, this expanded model would also act as a test of the distinctness of the Sepkoski’s three-fauna hypothesis [285] in terms of trait-dependent extinction.

Traits like environmental preference or geographic range [135, 132] are most likely heritable. Without phylogenetic context, this analysis assumes that differences in extinction risk between taxa are independent of the shared evolutionary history of those taxa [70]. In contrast, the origination cohorts only capture shared temporal context. For example, if taxon duration is phylogenetically heritable, then closely related taxa may have more similar durations as well as more similar biological traits. Without taking into account phylogenetic similarity the effects of these biological traits would be inflated solely due to inheritance. The inclusion of phylogenetic context as an additional individual-level hierarchical effect, independent of origination cohort, would allow for determining how much of the observed variability is due to shared evolutionary history versus shared temporal context versus actual differences associated with biological traits [309].

The combination and integration of the phylogenetic comparative and paleontological approaches requires both sources of data, something which is not possible for this analysis because there is no phylogenetic hypothesis for all Paleozoic taxa, something that is frequently the case for marine invertebrates with a good fossil record. When both data sources are available has been possible, however, the analysis can more fully address the questions of interest in macroevolution [309, 303, 304, 297, 329, 302, 249, 248, 116, 97].

In summary, patterns of Paleozoic brachiopod survival were analyzed using a fully Bayesian hierarchical survival modelling approach while also eschewing the traditional separation between background and mass extinction. I find that cohort extinction intensity is negatively correlated with both the cohort-specific effects of geographic range and environmental preference. These results imply that as extinction intensity increases (β^0) it is expected that both effects will increase in magnitude. However, the change in effect of environmental preference is expected to be greater than the change in the effect of geographic range. Additionally, I find support for greater survival in environmental generalists over specialists

in all origination cohorts analyzed; this is consistent with the long standing “survival of the unspecialized” hypothesis [169, 170, 298, 299, 309, 218, 217, 21]. The results of this analysis support the conclusion that for Paleozoic brachiopods, as extinction intensity increases overall extinction selectivity is expected to increase as well.

Table 3.1: Estimates of various parameters in the model used here. These include group-level estimates of the effects of biological traits on brachiopod generic survival, the standard deviation of the between-cohort effects, as well as the estimates of both the effect of sampling δ and the Weibull shape parameter α . The mean, standard deviation (SD), 10th, 50th, and 90th quantiles of the marginal posteriors are presented.

type	parameter	effect of	mean	SD	10%	50%	90%
Mean	μ^i	intercept	-3.05	0.20	-3.30	-3.05	-2.80
	μ^r	geographic range	-0.98	0.16	-1.18	-0.98	-0.79
	μ^v	environmental preference	-0.76	0.19	-0.99	-0.76	-0.52
	μ^{v^2}	environmental preference ²	3.15	0.36	2.69	3.15	3.62
Standard deviation	μ^m	body size	-0.01	0.13	-0.17	-0.01	0.15
	τ^i	intercept	0.51	0.11	0.38	0.50	0.65
	τ^r	geographic range	0.50	0.16	0.30	0.49	0.71
	τ^v	environmental preference	0.84	0.17	0.63	0.82	1.05
Other	τ^{v^2}	environmental preference ²	1.51	0.36	1.08	1.48	1.97
	τ^m	body size	0.47	0.13	0.32	0.46	0.64
	δ	sampling	0.90	0.15	0.71	0.90	1.09
	α	“time”	1.36	0.04	1.30	1.36	1.42

CHAPTER 4

SPECIES TRAITS AND ENVIRONMENTAL CONTEXT: THE CHANGING FUNCTIONAL COMPOSITION OF THE NORTH AMERICAN SPECIES POOL

4.1 Introduction

Changes to species diversity are the result of evolutionary and ecological processes acting both in concert and continually. Local communities are shaped by dispersal and local ecological processes such as resource competition and predator-prey relationships. The constituent species of these communities are drawn from a regional species pool, or the set of all species that are present in at least one community within a region [206, 331, 117]. Species dispersal from the regional species pool to the local communities is a sorting process shaped by biotic and abiotic environmental filters which are mediated by those species' traits [288, 64, 331, 178, 51, 117]. Regional species pools are shaped by speciation, extinction, migration, and extirpation. The gain or loss of regional diversity is the result of macroevolutionary dynamics which, in turn, shape the downstream macroecological dynamics of the regional species pool and its constituent local communities [331, 206, 117].

Fundamentally, all species respond differently to climate and environmental change [28]. Similarities in ecological roles of species within a regional species pool can be described as a collection of guilds or functional groups [333, 14, 32, 292, 363]. Species within the same functional group are expected to have more similar macroecological dynamics to each other than to species of a different functional group. By focusing on the relative diversity of functional groups, changes to diversity are interpretable as changes to the set of ways species within a species pool could interact with the biotic and abiotic environment.

A key question when comparing communities or regional species pools based on their functional composition is whether specific functional groups are enriched or depleted and why; what are the processes that led to a species pool having the functional composition it does [196, 358, 32, 306, 28]? Comparisons between contemporaneous regional species pools only reveal if a functional group is enriched or depleted in one species pool relative to other species pools. This type of comparison does not reveal if that functional group is enriched or depleted relative to its diversity in the regional species pool over time [28]. While a species pool may be depleted of a functional group relative to other contemporaneous species pools, that same functional group may be actually be enriched in that species pool relative to its historical diversity. Because the processes which shape regional species pool diversity (e.g. origination, extinction) operate on much longer time scales than is possible for studies of the Recent, paleontological data provides a unique opportunity to observe and estimate the changes to functional diversity and how species functional traits and environmental context can shape the enrichment or depletion of functional groups within a regional species pool [28, 306]. Being able to identify if the diversity of any functional groups are depleted relative to their long term average diversity in the species pool is particularly useful in conservation settings; species in depleted groups are most likely more at risk of extinction than species in enriched groups, even if those enriched groups are relatively rare when compared to the functional composition of other contemporaneous species pools.

The paleontological record of North American mammals for the Cenozoic (\sim 66 million years ago to present) provides one of the best opportunities for understanding how regional species pool functional diversity changes over time. The North American mammal record is a relatively complete temporal sequence for the entire Cenozoic which is primarily, but not exclusively, based on fossil localities from the Western Interior/Great Plains region of North America [5, 10, 6]. Additionally, mammal fossils preserve a lot of important functional information, such as teeth, so that important ecological traits like the dietary/trophic category

of species are easy to estimate [234, 233, 67].

The goals of this study are to understand when are unique functional groups, called ecotypes, enriched or depleted in the North American mammal regional species pool and to estimate the relationship between changes to regional ecotypic diversity and changes to their environmental context.

4.1.1 Background

The diversity history of North American mammals for the Cenozoic is relatively well understood as it has been the focus of considerable study [6, 5, 143, 10, 72, 230, 93, 309, 240, 304, 290, 12, 28, 147]. Previous approaches to understanding mammal diversity, both in North America and elsewhere, fall into a number of overlapping categories: total diversity [10, 5, 72, 171], with/between guild comparisons [140, 141, 152, 147, 230, 144], within/between clade comparisons [240, 304, 290, 93, 37], and estimating the impact of environmental process on diversity [28, 147, 143, 93, 66, 12, 13, 10]. Each of these individual perspectives provide a limited perspective on the macroevolutionary and macroecological processes shaping diversity and diversification. Integration across perspectives is necessary for producing a holistic and internally consistent picture of how the North American mammal species pool has changed through time. One of the goals of this study is to present a framework for approaching hypotheses about diversity and diversification through multiple lenses simultaneously so that our inferences are better constrained and the relative importance of various functional traits and environmental factors may be better elucidated.

The narrative of the diversification of North American mammals over the Cenozoic is one of gradual change. There is little convincing evidence that there have been any major or sudden cross-functional group or cross-taxonomic turnover events in mammal diversity at any point

in the Cenozoic record of North America [6, 5, 66, 143, 10]. Instead of being concentrated at specific time intervals, species turnover has been found to be distributed through time. It is then expected that, for this analysis, turnover events or periods of rapid diversification or depletion should not occur simultaneously for all functional groups under study. Additionally, changes to mammal diversification seem to be primarily driven by changes to origination rate and not to extinction [5, 10, 6]. An unresolved aspect of the general history of mammal diversification is whether that diversity is limited or self-regulating; namely, to what extent is mammal diversification diversity-dependent [6, 244, 113, 242]. Similarly, this question can also be asked of specific functional groups [152, 338, 290, 240].

Within the overall narrative of mammal diversity, the histories of a selection of taxonomic and functional groups are better understood. These groups have particularly good fossil records and/or have been the focus of previous analyses.

The diversity history of ungulate herbivores has been characterized by more recently originating taxa having longer legs, higher crowned teeth, and a shift from graze-dominated to browse-dominated diets than their earlier originating counterparts [140, 141, 143, 144, 37, 93]. The mechanisms which drive this pattern are theorized to be some combination of tectonic activity driving environmental change such as the drying of the western interior of North America due mountain building and global temperature and environmental change such as the formation of polar icecaps [144, 66, 28, 13].

In contrast, the origination of modern cursorial pursuit carnivore forms was not until later in the Cenozoic; this is not to say that carnivore diversity only grew in the late Cenozoic, but that those forms were late entrants [147]. Instead, the diversity history of carnivores reflects density-dependence or some other form of self-regulation [338, 290, 304]. Specifically, it has been proposed that different canid clades have replaced each other as the dominant members of their functional group within the species pool [290, 338]. It is then expected that, for this

analysis, the diversity of digitigrade and plantigrade carnivores combined (i.e. the “carnivore” guild [338]) should be relatively constant for the Cenozoic or at least have plateaued by the Neogene, though the relative diversities of digitigrade versus plantigrade forms are subject to change [235].

In a relevant study, Smits [309] found that functional traits such as a species dietary or locomotor category structure differences in mammal extinction risk. In particular, arboreal taxa were found to have a shorter duration on average than species from other locomotor categories [309]. Two possible scenarios that could yield this pattern were proposed: the extinction risk faced by arboreal species is constant and high for the entire Cenozoic or the Paleogene and Neogene represent different regimes and extinction risk increased in the Neogene, thus driving up the Cenozoic average extinction risk. These two possible explanations have clear and testable predictions with respect to the diversity history of arboreal taxa: 1) if arboreal taxa always have an elevated extinction risk when compared to other taxa, then the diversity history of arboreal taxa is expected to be constant with time, albeit possibly at low diversity; and 2) if the Paleogene and Neogene represent difference selective regimes with the former being associated with lower extinction risk than the latter, then the diversity history of arboreal taxa are expected to be present in the Paleogene but depleted or absent from the species pool during the Neogene.

The climate history of the Cenozoic can be broadly described as a gradual cooling trend, with polar ice-caps forming in the Neogene [368, 367, 53]. There are of course exceptions to this pattern such as the Eocene climatic optimum, the mid-Miocene climatic optimum, and the sudden drop in temperature at the Eocene/Oligocene boundary [368, 367]. In terms of the North American biotic environment, the Cenozoic is additionally characterized by major transition from having closed, partially forested biomes being common in the Paleogene to the landscape being dominated by savannah and grasslands biomes by the Neogene

[28, 143, 141, 326]. Additionally, the landscape structure and topology of North America changed substantially over the Cenozoic with mountain uplift and other tectonic actives in Western North America [28, 66, 144, 12]. This type of geological activity affects both local climates as well as continental weather patterns while also mobilizing increased grit into the environment, something which may be responsible for increasing trend of hyposodony (high crowned teeth) among herbivores [148, 151, 58].

The effect of climate on mammal diversity and its accompanying diversification process has been the focus of considerable research with a slight consensus favoring mammal diversification being more biologically-mediated than climate-mediated [10, 72, 46]. However, differences in temporal and geographic scale seem to underlie the contrast between these two perspectives. For example when the mammal fossil record analyzed at small temporal and geographic scales a correlation between diversity and climate is observable [46]. However, when the record is analyzed at the scale of the continent and most of the Cenozoic this correlation disappears [10]. This result, however, does not go against the idea that there may be short periods of correlation between diversity and climate and that this relationship can change or even reverse direction over time; this type result means that there is no single direction of correlation between diversity and climate [72].

In the case of a fluctuating correlation between diversity and climate it is hard to make the argument for an actual causal link between the two without modeling the underlying ecological differences between species; after all, species respond differently based on their individual ecologies [28]. When analysis is based on diversity or taxonomy alone no mechanisms are possible to infer. Taxonomy, like body size, stands in for many important species traits to the point that mechanistic or process based inference is impossible. While emergent patterns might correspond to taxonomic grouping, this itself is an emergent phenomenon. Instead, by framing hypotheses in terms of species traits and their environmental context, these emergent

phenomena can be observed and analyzed rather than assumed.

4.1.2 Foreground

Fourth-corner modeling is an approach to explaining the patterns of either species abundance or presence/absence in a community as a product of species traits, environmental factors, and the interaction between traits and environment [31, 356, 232, 139]; effectively uniting climate-based species distribution modeling (SDMs) with trait-based community assembly models (CATS, MaxEnt). In modern ecological studies, what is being modeled is species occurrences at localities distributed across a region [232, 139]. In this study, what is being modeled is the pattern of species occurrence over time for most of the Cenozoic record for North America (Fig. 4.1). By analyzing assemblages over time instead of space in fourth-corner framework we can gain better inference of how an instantaneous species pool (i.e. the Recent) is assembled over time. These two approaches, modern and paleontological, are different views of the same three-dimensional pattern: species at localities over time. The temporal limitations of modern ecological studies and difficulties with uneven spatial occurrences of fossils in paleontological studies means that these approaches are complementary and reveal different patterns of how species are distributed in time and space.

My approach to delimiting and assigning mammal functional groups is inspired on the ecocube heuristic used to classify marine invertebrate species by three functional traits [36, 15, 35, 34, 216, 342] and creodont mammals in a similar fashion [208]. Unique combinations of traits represent ecotypes, which are equivalent to functional groups defined by species functional traits instead of a holistic understanding how a taxon interacts with its environment. In this study, the two functional traits used to define a species' ecotype are dietary (e.g. herbivore, carnivore, etc.) and locomotor category (e.g. arboreal, unguligrade, etc.). Species body mass was also included as a species trait in this analysis, but not as a functional trait for defining

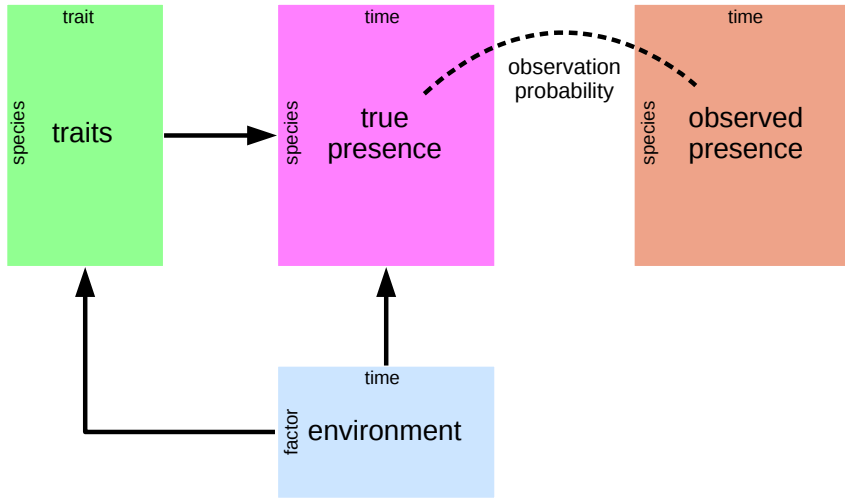


Figure 4.1: Conceptual diagram of the paleontological fourth corner problem. The observed presence matrix (orange) is the empirical presence/absence pattern for all species for all time points; this matrix is an incomplete observation of the “true” presence/absence pattern (purple). The estimated true presence matrix is modeled as a function of both environmental factors over time (blue) and multiple species traits (green). Additionally, the effects of environmental factors on species traits are also modeled, as traits are expected to mediate the effects of a species environmental context. This diagram is based partially on material presented in Brown *et al.* [31] and Warton *et al.* [356].

ecotypes; instead, its inclusion is principally to control for differences in species dynamics that driven by mass and not ecotype.

The environmental factors included in this study are estimates of global temperature and the changing floral groups present in North America across the Cenozoic [53, 109]. These covariates were chosen because they provide high level characterizations of the environmental context of the entire North American regional species pool for most of the Cenozoic. Importantly, the effects of a species ecotype on diversity are themselves modeled as functions of environmental factors (Fig. 4.1) allowing for inference as to how a species ecology can mediate selective pressures due to its environmental context.

All observations, paleontological or modern, are made with uncertainty. With presence/absence data this uncertainty comes from not knowing if an absence is a “true” absence or just a failure to observe [271, 272, 76, 81, 177, 352]. For paleontological data, the incomplete preservation and sampling of species means that the true times of origination or extinction may not be observed [76, 81, 350, 352]. The model(s) I propose below represent an attempt to translate the verbal/visual model described here (Fig. 4.1) into a statistical model for estimating the relative diversity of mammal ecotypes over time and how those ecotypes respond to changes to environmental context while taking into account the fundamental incompleteness of the fossil record.

Ultimately, the goals of this analysis are to understand when different ecotypes enriched or depleted in the North American mammal regional species pool and how these changes in ecotypic diversity are related to changes in species’ environmental context. In the analyses done here, many covariates which describe a species’ macroecology and its environmental context are considered. In order to analyze this complex and highly structured data set, I developed a hierachal Bayesian model combining the fourth-corner modeling approach with a model of an observation-occurrence or observation-origination-extinction process.

4.2 Materials and Methods

4.2.1 *Taxon occurrences and species-level information*

All fossil occurrence information used in this analysis was downloaded from the Paleobiology Database (PBDB). The initial download restricted occurrences to Mammalia observed in North America between the Maastrichtian (72-66 Mya) and Gelasian (2.58-1.8 Mya) stages [47]. Occurrences were then further limited to those occurring between 64 and 2 million years

ago (Mya); this age restriction was to insure that observation time series lines up with the temperature time series [53]. Taxonomic, stratigraphic, and ecological metadata for each occurrence and species was also downloaded. A new download for a raw, unfiltered PBDB datafile following the same criterion used here is available at <http://goo.gl/2s1geU>. The raw datafile used as a part of this study, along with all code for filtering and manipulating this download is available at <http://github.com/psmits/coping>.

After being downloaded, the raw occurrence data was then sorted, cleaned, and manipulated programmatically before analysis. Many species taxonomic assignments as present in the raw PBDB data were updated for accuracy and consistency. For example, species classified in the order Artiodactyla were reclassified as Cetartiodactyla. These re-assignments follow previous work [309] which were based on taxonomies present in the Encyclopedia of Life (<http://eol.org>) and other volumes [146, 145]. All taxa whose life habit was classified as either volant (i.e. Chiroptera) or aquatic (e.g. Cetacea) were excluded from this analysis because of their lack of direct applicability to the study of terrestrial species pools.

Species ecotype is defined based on a combination of locomotor and diet categories; the goal is to classify species based on the manner with which they interact with their environment. Most mammal species records in the PBDB have life habit (i.e. locomotor category) and dietary category assignments. In order to simplify interpretation, analysis, and per-ecotype sample size these classifications were coarsened in a similar manner to [309] following Table 4.1. The locomotor category was then further broken up to better reflect the diversity of mammal locomotor modes. Ground dwelling species locomotor categories were reassigned based on the ankle posture associated with their taxonomic group, as described in Table 4.2 [38]. Ankle posture was assumed uniform for all species within a taxonomic group except for those species assigned a non-ground dwelling locomotor category by the PBDB. All species for which it was possible to assign a locomotor category had one assigned, including species

for which post-crania are unknown but for which a taxonomic grouping is known. Ground dwelling species which were unable to be reassigned based on ankle posture were excluded from analysis. Finally, ecotype categories with less than 10 total species were excluded, yielding a total of 18 observed ecotypes out of a possible 24.

Table 4.1: Species trait assignments in this study are a coarser version of the information available in the PBDB. Information was coarsened to improve per category sample size.

This study		PBDB categories
Diet	Carnivore	Carnivore
	Herbivore	Browser, folivore, granivore, grazer, herbivore.
	Insectivore	Insectivore.
	Omnivore	Frugivore, omnivore.
Locomotor	Arboreal	Arboreal.
	Ground dwelling	Fossorial, ground dwelling, semifossorial, saltatorial.
	Scansorial	Scansorial.

Table 4.2: Ankle posture assignment as based on taxonomy. Assignments are based on [38]. Taxonomic groups are presented alphabetically and without reference for the nestedness of families in orders.

Order	Family	Stance
	Ailuridae	plantigrade
	Allomyidae	plantigrade
	Amphicyonidae	plantigrade
	Amphilemuridae	plantigrade
	Anthracotheriidae	digitigrade
	Antilocapridae	unguligrade
	Apheliscidae	plantigrade
	Aplodontidae	plantigrade
	Apternodontidae	scansorial
	Arctocyonidae	unguligrade
	Barbourofelidae	digitigrade
	Barylambdidae	plantigrade
	Bovidae	unguligrade
	Camelidae	unguligrade
	Canidae	digitigrade
	Cervidae	unguligrade
	Cimolodontidae	scansorial
	Coryphodontidae	plantigrade
	Cricetidae	plantigrade
	Cylindrodontidae	plantigrade
	Cyriacotheriidae	plantigrade

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Table 4.2 – continued from previous page

Order	Family	Stance
	Dichobunidae	unguligrade
Dinocerata		unguligrade
	Dipodidae	digitigrade
	Elephantidae	digitigrade
	Entelodontidae	unguligrade
	Eomyidae	plantigrade
	Erethizontidae	plantigrade
	Erinaceidae	plantigrade
	Esthonychidae	plantigrade
	Eutypomyidae	plantigrade
	Felidae	digitigrade
	Florentiamyidae	plantigrade
	Gelocidae	unguligrade
	Geolabididae	plantigrade
	Glyptodontidae	plantigrade
	Gomphotheriidae	unguligrade
	Hapalodectidae	plantigrade
	Heteromyidae	digitigrade
	Hyaenidae	digitigrade
	Hyaenodontidae	digitigrade
	Hypertragulidae	unguligrade
	Ischyromyidae	plantigrade
	Jimomyidae	plantigrade

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Table 4.2 – continued from previous page

Order	Family	Stance
Lagomorpha		digitigrade
	Leptictidae	plantigrade
	Leptochoeridae	unguligrade
	Leptomerycidae	unguligrade
	Mammutidae	unguligrade
	Megalonychidae	plantigrade
	Megatheriidae	plantigrade
	Mephitidae	plantigrade
	Merycoidodontidae	digitigrade
Mesonychia		unguligrade
	Mesonychidae	digitigrade
	Micropternodontidae	plantigrade
	Mixodectidae	plantigrade
	Moschidae	unguligrade
	Muridae	plantigrade
	Mustelidae	plantigrade
	Mylagaulidae	fossorial
	Mylodontidae	plantigrade
	Nimravidae	digitigrade
	Nothrotheriidae	plantigrade
Notoungulata		unguligrade
	Oromerycidae	unguligrade
	Oxyaenidae	digitigrade

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Table 4.2 – continued from previous page

Order	Family	Stance
	Palaeomerycidae	unguligrade
	Palaeoryctidae	plantigrade
	Pampatheriidae	plantigrade
	Pantolambdidae	plantigrade
	Peritychidae	digitigrade
Perissodactyla		unguligrade
	Phenacodontidae	unguligrade
Primates		plantigrade
	Procyonidae	plantigrade
	Proscalopidae	plantigrade
	Protoceratidae	unguligrade
	Reithroparamyidae	plantigrade
	Sciuravidae	plantigrade
	Sciuridae	plantigrade
	Simimyidae	plantigrade
	Soricidae	plantigrade
	Suidae	digitigrade
	Talpidae	fossorial
	Tayassuidae	unguligrade
	Tenrecidae	plantigrade
	Titanoideidae	plantigrade
	Ursidae	plantigrade
	Viverravidae	plantigrade

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Table 4.2 – continued from previous page

Order	Family	Stance
	Zapodidae	plantigrade

Estimates of species mass used in this study were sourced from multiple databases and papers, especially those focusing on similar macroevolutionary or macroecological questions [329, 30, 94, 198, 248, 305]; this is similar to what has been done before [309]. When species mass was not available, proxy measures were used and then used to estimate average species mass. For example, given a measurement of a mammal tooth size, it is possible and routine to estimate its mass given some regression equation (Table 4.3). The PBDB has one or more body part measures for many species. These were used as body size proxies for many species. Mass was log-transformed and then rescaled by first subtracting mean log-mass from all mass estimates, then dividing by two-times its standard deviation; this insures that the magnitude of effects for both continuous and discrete covariates are directly comparable [102, 100].

In total, 1400 mammal species occurrence histories were included in this study after applying all of the restrictions above.

All fossil occurrences from 64 to 2 million years ago (Mya) were binned into 31 two-million year (My) bins. This temporal length was chosen because it is approximately the resolution of the North American mammal fossil record [5, 10, 191, 6].

Table 4.3: Regression equations used in this study for estimating body size. Equations are presented with reference to taxonomic grouping, part name, and reference.

Group	Equation	log(Measurement)	Source
General	$\log(m) = 1.827x + 1.81$	lower m1 area	[165]
General	$\log(m) = 2.9677x - 5.6712$	mandible length	[91]
General	$\log(m) = 3.68x - 3.83$	skull length	[183]
Carnivores	$\log(m) = 2.97x + 1.681$	lower m1 length	[337]
Insectivores	$\log(m) = 1.628x + 1.726$	lower m1 area	[26]
Insectivores	$\log(m) = 1.714x + 0.886$	upper M1 area	[26]
Lagomorph	$\log(m) = 2.671x - 2.671$	lower toothrow area	[329]
Lagomorph	$\log(m) = 4.468x - 3.002$	lower m1 length	[329]
Marsupials	$\log(m) = 3.284x + 1.83$	upper M1 length	[108]
Marsupials	$\log(m) = 1.733x + 1.571$	upper M1 area	[108]
Rodentia	$\log(m) = 1.767x + 2.172$	lower m1 area	[165]
Ungulates	$\log(m) = 1.516x + 3.757$	lower m1 area	[201]
Ungulates	$\log(m) = 3.076x + 2.366$	lower m2 length	[201]
Ungulates	$\log(m) = 1.518x + 2.792$	lower m2 area	[201]
Ungulates	$\log(m) = 3.113x - 1.374$	lower toothrow length	[201]

4.2.2 Environmental and temporal covariates

The environmental covariates used in this study are collectively referred to as group-level covariates because they predict the response of a “group” of individual-level observations (i.e. species occurrences of an ecotype). Additionally, these covariates are defined for temporal bins and not the species themselves; as such they predict the parts of each species occurrence history. The group-level covariates in this study are two global temperature estimates and the Cenozoic “plant phases” defined by Graham [109].

Global temperature across most of the Cenozoic was calculated from Mg/Ca isotope record from deep sea carbonates [53]. Mg/Ca based temperature estimates are preferable to the frequently used $\delta^{18}\text{O}$ temperature proxy [368, 367, 10, 72] because Mg/Ca estimates do not conflate temperature with ice sheet volume and depth/stratification changes. The former is particularly important to this analysis as the current polar ice-caps appeared and grew during the second half of the Cenozoic. These properties make Mg/Ca based temperature

Table 4.4: Definitions of the start and stop times of the three plant phases used this study as defined by Graham [109].

Plant phase	Phase number	Start	Stop
Paleocene-Eocene	1	66	50
Eocene-Miocene	2	50	16
Miocene-Pleistocene	3	16	2

estimates preferable for macroevolutionary and macroecological studies [69]. Two aspects of the Mg/Ca-based temperature curve were included in this analysis: mean and range. Both were calculated as the mean of all respective estimates for each 2 My temporal bins. The distributions of the temperature mean and range estimates were then rescaled by subtracting their respective means from all values and then dividing by twice their respective standard deviations.

The second set of environmental factors included in this study are the Cenozoic plant phases defined by Graham [109]. Graham’s plant phases are holistic descriptors of the taxonomic composition of 12 ecosystem types, which plants are present at a given time, and the relative modernity of those plant groups with younger phases representing increasingly modern taxa [109]. Graham [109] defines four intervals from the Cretaceous to the Pliocene, though only three of these intervals take place during the time frame being analyzed. Graham’s plant phases was included as a series of “dummy variables” encoding the three phases included in this analysis [102]; this means that the first phase is synonymous with the intercept and subsequent phases are defined by their differences from the first phase. The temporal boundaries of these plant phases are defined in Table 4.4.

4.2.3 Modeling species occurrence

Two different models were used in this study: a pure-presence model and a birth-death model. Both models at their core are hidden Markov models where the latent process has an

State at $t + 1$				State at $t + 1$					
		0_{never}	1	$0_{extinct}$			0_{never}	1	$0_{extinct}$
State at t	0_{never}	$1 - \theta$	θ	0	State at t	0_{never}	$1 - \pi$	π	0
	1	0	θ	$1 - \theta$		1	0	ϕ	$1 - \phi$
	$0_{extinct}$	0	0	1		$0_{extinct}$	0	0	1

(a) Pure-presence

(b) Birth-death

Table 4.5: Transition matrices for the pure-presence (4.5a) and birth-death (4.5b) models. Both of these models share the core machinery of discrete-time birth-death processes but make distinct assumptions about the equality of originating and surviving (Eq. 4.2, and 4.3). Note also that while there are only two state “codes” (0, 1), there are in fact three states: never having originated 0_{never} , present 1, extinct $0_{extinct}$ [4].

absorbing state [4]. The difference between these two models lies in whether the probabilities of a species originating or surviving are considered equal or different (Table 4.5). While there are only two state “codes” in a presence-absence matrix (i.e. 0/1), there are in fact three states in a birth-death model: not having originated yet, extant, and extinct. The last of these is the absorbing state, as once a species has gone extinct it cannot re-originate [4]. Thus, in the transition matrices the probability of an extinct species changing states is 0 (Table 4.5). See below for parameter explanations (Tables 4.6, 4.7, and 4.8).

Observation process

The type of hidden Markov model used in this study has three characteristic probabilities: probability p of observing a species given that it is present, probability ϕ of a species surviving from one time to another, and probability π of a species first appearing [271]. In this formulation, the probability of a species becoming extinct is $1 - \phi$. For the pure-presence model $\phi = \pi$, while for the birth-death model $\phi \neq \pi$.

The probability p of observing a species that is present is modeled as a logistic regression with a time-varying intercept and species mass as a covariate. The effect of species mass on p

Table 4.6: Parameters for the observation process part of the hidden Markov model.

Parameter	dimensions	explanation
y	$N \times T$	observed species presence/absence
z	$N \times T$	“true” species presence/absence
p	T	probability of observing a species that is present at time t
m	N	species log mass, rescaled
α_0	1	average log-odds of p
α_1	1	change in average log-odds of p per change mass
r	T	difference from α_0 associated with time t
σ	1	standard deviation of r

was assumed linear and constant over time. These assumptions are reflected in the structure of this part of the model being presented here:

$$\begin{aligned} y_{i,t} &\sim \text{Bernoulli}(p_{i,t} z_{i,t}) \\ p_{i,t} &= \text{logit}^{-1}(\alpha_0 + \alpha_1 m_i + r_t) \\ r_t &\sim \mathcal{N}(0, \sigma). \end{aligned} \tag{4.1}$$

The parameters associated with Equation 4.1 are described in Table 4.6.

Pure-presence process

For the pure-presence model there is only a single probability dealing with the presence of a species θ (Table 4.5a). This probability was modeled as multi-level logistic regression with both species-level and group-level covariates [102, 101]. The parameters associated with the pure-presence model are presented in Table 4.7, and the full sampling statement in Equation 4.2.

Species mass was included as a covariate with two regression coefficients allowing for a quadratic relationship with log-odds of occurrence. Because the distribution of mammal species body mass is unimodal and approximately log-normal [305], I assume that species of

Table 4.7: Parameters for the model of occurrence in the pure-presence model

Parameter	dimensions	explanation
z	$N \times T$	“true” species presence/absence
θ	$N \times T - 1$	probability of $z = 1$
a	$T - 1 \times D$	ecotype-varying intercept; mean value of log-odds of θ
m	N	species log mass, rescaled
b_1	1	effect of species mass on log-odds of θ
b_2	1	effect of species mass, squared, on log-odds of θ
U	$T \times D$	matrix of group-level covariates
γ	$U \times D$	matrix of group-level regression coefficients
Σ	$D \times D$	covariance matrix of a
Ω	$D \times D$	correlation matrix of a
τ	D	vector of standard deviations for each ecotype a_d

intermediate body size will be more common than species of very large or very small mass. These assumptions are also reflected in the choice of priors for b_1 and b_2 where the latter is given a weakly informative prior with most of its density below 0 (Eq. 4.2).

The values of each ecotype’s intercept are themselves modeled as regressions using the group-level covariates associated with environmental context. Each of these regressions has an associated variance of possible values of each ecotype’s intercept [102]. In addition, the covariances between ecotype intercepts, given this group-level regression, are modeled [102]. The prior choice for the covariance matrix separates it into a vector of scales τ and a correlation matrix Ω . The elements of the former are given weakly informative, independent half-Normal priors while the latter was given a weakly informative LKJ prior as recommended in the Stan manual [319].

All parameters not modeled elsewhere were given weakly informative priors [101, 195, 319]. Weakly informative means that priors do not necessarily encode actual prior information but instead help regularize or weakly constrain posterior estimates. These priors have a concentrated probability density around and near zero; this has the effect of tempering our estimates and help prevent overfitting the model to the data [101, 195, 319]. The general line

of thinking behind this approach is that a result of 0 or “no effect” is more preferable to a wrong or extremely weak result.

$$\begin{aligned}
y_{i,t} &\sim \text{Bernoulli}(p_{i,t} z_{i,t}) & \alpha_0 &\sim \mathcal{N}(0, 1) \\
p_{i,t} &= \text{logit}^{-1}(\alpha_0 + \alpha_1 m_i + r_t) & \alpha_1 &\sim \mathcal{N}(1, 1) \\
r_t &\sim \mathcal{N}(0, \sigma) & \sigma &\sim \mathcal{N}^+(1) \\
z_{i,1} &\sim \text{Bernoulli}(\rho) & b_1 &\sim \mathcal{N}(0, 1) \\
z_{i,t} &\sim \text{Bernoulli}(\theta_{i,t}) & b_2 &\sim \mathcal{N}(-1, 1) \\
\theta_{i,t} &= \text{logit}^{-1}(a_{t,j[i]} + b_1 m_i + b_2 m_i^2) & \gamma &\sim \mathcal{N}(0, 1) \\
a &\sim \text{MVN}(u\gamma, \Sigma) & \tau &\sim \mathcal{N}^+(1) \\
\Sigma &= \text{diag}(\tau)\Omega\text{diag}(\tau) & \Omega &\sim \text{LKJ}(2)
\end{aligned} \tag{4.2}$$

Birth-death process

In the birth-death version of the model, $\phi \neq \pi$ and so each of these probabilities is modeled separately but each is handled in a similar manner to how θ is modeled in the pure-presence model (Eq. 4.2, Table 4.5b). The parameters associated with the birth-death presence model are presented in Table 4.8 and the full sampling statement, including observation (Eq. 4.1),

is described in Equation 4.3:

$$\begin{aligned}
y_{i,t} &\sim \text{Bernoulli}(p_{i,t} z_{i,t}) & \Sigma^\phi &= \text{diag}(\tau^\phi) \Omega^\phi \text{diag}(\tau^\phi) \\
p_{i,t} &= \text{logit}^{-1}(\alpha_0 + \alpha_1 m_i + r_t) & \Sigma^\pi &= \text{diag}(\tau^\pi) \Omega^\pi \text{diag}(\tau^\pi) \\
r_t &\sim \mathcal{N}(0, \sigma) & \rho &\sim U(0, 1) \\
\alpha_0 &\sim \mathcal{N}(0, 1) & b_1^\phi &\sim \mathcal{N}(0, 1) \\
\alpha_1 &\sim \mathcal{N}(1, 1) & b_1^\pi &\sim \mathcal{N}(0, 1) \\
\sigma &\sim \mathcal{N}^+(1) & b_2^\phi &\sim \mathcal{N}(-1, 1) \\
z_{i,1} &\sim \text{Bernoulli}(\phi_{i,1}) & b_2^\pi &\sim \mathcal{N}(-1, 1) \\
z_{i,t} &\sim \text{Bernoulli} \left(z_{i,t-1} \pi_{i,t} + \sum_{x=1}^t (1 - z_{i,x}) \phi_{i,t} \right) & \gamma^\phi &\sim \mathcal{N}(0, 1) \\
&& \gamma^\pi &\sim \mathcal{N}(0, 1) \\
\phi_{i,t} &= \text{logit}^{-1}(a_{t,j[i]}^\phi + b_1^\phi m_i + b_2^\phi m_i^2) & \tau^\phi &\sim \mathcal{N}^+(1) \\
\pi_{i,t} &= \text{logit}^{-1}(a_{t,j[i]}^\pi + b_1^\pi m_i + b_2^\pi m_i^2) & \tau^\pi &\sim \mathcal{N}^+(1) \\
a^\phi &\sim \text{MVN}(U \gamma^\phi, \Sigma^\phi) & \Omega^\phi &\sim \text{LKJ}(2) \\
a^\pi &\sim \text{MVN}(U \gamma^\pi, \Sigma^\pi) & \Omega^\pi &\sim \text{LKJ}(2).
\end{aligned} \tag{4.3}$$

Similar to the pure-presence model, both ϕ and π are modeled as logistic regressions with varying intercept and one covariate associated with two parameters. The possible relationships between mass and both ϕ and π are reflected in the parameterization of the model and choice of priors (Eq. 4.3).

The intercepts of ϕ and π both vary by species ecotype and those values are themselves the product of group-level regression using environmental factors as covariates (Eq. 4.3); this is identical to the pure presence model (Eq. 4.2).

Table 4.8: Parameters for the model of presence in the pure-presence model

Parameter	dimensions	explanation
z	$N \times T$	“true” species presence/absence
ϕ	$N \times T$	probability of $z_{-,t} = 1 z_{-,t-1} = 0$; origination
π	$N \times T - 1$	probability of $z_{-,t} = 1 z_{-,t-1} = 1$; survival
a^ϕ	$T - 1 \times D$	ecotype-varying intercept; mean value of log-odds of θ
a^π	$T - 1 \times D$	ecotype-varying intercept; mean value of log-odds of θ
m	N	species log mass, rescaled
b_1^ϕ	1	effect of species mass on log-odds of ϕ
b_1^π	1	effect of species mass on log-odds of π
b_2^ϕ	1	effect of species mass, squared, on log-odds of ϕ
b_2^π	1	effect of species mass, squared, on log-odds of π
U	$T \times D$	matrix of group-level covariates
γ^ϕ	$U \times D$	matrix of group-level regression coefficients
γ^π	$U \times D$	matrix of group-level regression coefficients
Σ^ϕ	$D \times D$	covariance matrix of a^ϕ
Σ^π	$D \times D$	covariance matrix of a^π
Ω^ϕ	$D \times D$	correlation matrix of a^ϕ
Ω^π	$D \times D$	correlation matrix of a^π
τ^ϕ	D	vector of standard deviations for each ecotype a_d^ϕ
τ^π	D	vector of standard deviations for each ecotype a_d^π

4.2.4 Posterior inference and model adequacy

Computer programs that implement joint posterior inference for the above models (Eqs. 4.2, 4.3) were written in the probabilistic programming language Stan [319]. Both models feature a large matrix of latent discrete parameters z (Tables 4.6, 4.7, 4.8; Eqs. 4.1, 4.2, 4.3). All methods for posterior inference implemented in Stan are derivative-based; this causes complications for actually implementing the above models, because integers do not have derivatives. Instead of implementing a latent discrete parameterization, the log posterior probabilities of all possible states of the latent parameters z were calculated and summed (i.e. marginalized).

Species durations at minimum range through from a species first appearance to their last appearance in the fossil record, but the incompleteness of all observations means that the actual times of origination and extinction are unknown. The marginalization approach used here means that the probabilities of all possible histories for a species are calculated, from the end members of the species having existed for the entire study interval and the species having only existed between the directly observed first and last appearances to all possible intermediaries (Fig 4.2) [319]. This process is identical, language-wise, to assuming range-through and then estimating the possibility of all possible range extension due to incomplete sampling.

The combined size of the dataset and large number of parameters in both models (Eqs. 4.2, 4.3), specifically the total number of latent parameters that are the matrix z , means that stochastic approximate posterior inference is computationally very slow even using NUTS based HMC as implemented in Stan [319]. Instead, an approximate Bayesian approach was used: variational inference. A recently developed automatic variational inference algorithm called “automatic differentiation variational inference” (ADVI) is implemented in Stan and was used here [163, 319]. ADVI assumes that the posterior is Gaussian but still yields a true

	Time Bin							
	1	2	3	4	5	6	7	8
Observed	0	0	0	1	0	1	1	0
-----	-----	-----	-----	-----	-----	-----	-----	-----
Certain	?	?	?	1	1	1	1	?
.....
Potential	0	0	0	1	1	1	1	0
Potential	0	0	1	1	1	1	1	0
Potential	0	1	1	1	1	1	1	0
Potential	0	0	0	1	1	1	1	1
Potential	0	0	1	1	1	1	1	1
Potential	0	1	1	1	1	1	1	1
Potential	1	1	1	1	1	1	1	1

Figure 4.2: Conceptual figure of all possible occurrence histories for an observed species. The first row represents the observed presence/absence pattern for a single species at eight time points. The second row corresponds to the known aspects of the “true” occurrence history of that species. The remaining rows correspond to all possible occurrence histories that are consistent with the observed data. By marginalizing over all possible occurrence histories, the probability of each potential history is estimated. The process of parameter marginalization is described in the text.

Bayesian posterior; this assumption is similar to quadratic approximation of the likelihood function commonly used in maximum likelihood based inference [195]. The principal limitation of assuming the joint posterior is Gaussian is that the true topology of the log-posterior isn’t estimated; this is a particular burden for scale parameters which are bounded to be positive (e.g. standard deviation).

Of additional concern for posterior inference is the partial identifiability of observation parameters $p_{t=1}$ and $p_{t=T}$ [271]. This issue means that the estimates of sampling probabilities at the “edges” of the time series cannot fully be estimated because there are no known “gaps” in species occurrence histories that are guaranteed to be filled. Instead, the values of the first and final columns of the “true” presence-absence matrix z for those observations that do

not already have presences in the observed presence-absence matrix y cannot be estimated [271]. The hierarchical modeling approach used here helps mitigate this problem by pulling the values of $p_{t=1}$ and $p_{t=T}$ towards the overall mean of p [101], and in fact this approach might be more analytically sound than the more ad-hoc approaches that are occasionally used to overcome this hurdle [271]. Additionally, because $p_{t=1}$ and $p_{t=T}$ are only partially identifiable, estimates of occurrence θ and origination ϕ at $t = 1$ and estimates of θ , ϕ and survival π at $t = T$ may suffer from similar edge effects. Again, the hierarchical modeling approach used here may help correct for this reality by drawing these estimates towards the overall means of those parameters.

After fitting both models (Eqs. 4.2, 4.3) using ADVI, model adequacy and quality of fit were assessed using a posterior predictive check [101]. By simulating 100 theoretical data sets from the posterior estimates of the model parameters and the observed covariate information the congruence between predictions made by the model and the observed empirical data can be assessed. These datasets are simulated by starting with the observed states of the presence-absence matrix at $t = 1$; from there, the time series roll forward as stochastic processes with covariate information given from the empirical observations. Importantly, this is fundamentally different from observing the posterior estimates of the “true” presence-absence matrix z . The posterior predictive check used in this study is to compare the observed average number of observations per species to a distribution of simulated averages; if the empirically observed value sits in the middle of the distribution then the model can be considered adequate in reproducing the observed number of occurrences per species.

The ADVI assumption of a purely Gaussian posterior limits the utility and accuracy of the posterior predictive checks because parameter estimates do not reflect the true posterior distribution and are instead just an approximation [101]. Because of this, posterior predictive estimates are themselves only approximate checks of model adequacy. The posterior predictive

check that is used in this study focuses on mean occurrence and not to any scale parameters that might be most affected by the ADVI assumptions.

Given parameter estimates, diversity and diversification rates are estimated through posterior predictive simulations. Given the observed presence-absence matrix y , estimates of the true presence-absence matrix z can be simulated and the distribution of possible occurrence histories can be analyzed. This is conceptually similar to marginalization where the probability of each possible occurrence history is estimated (Fig. 4.2), but now these occurrence histories are generated relative to their estimated probabilities.

The posterior distribution of z gives the estimate of standing diversity N_t^{stand} for all time points as

$$N_t^{stand} = \sum_{i=1}^M z_{i,t}. \quad (4.4)$$

Given estimates of N_t^{stand} for all time points, the estimated number of originations O_t is estimated as

$$O_t = \sum_{i=1}^M z_{i,t} = 1 | z_{i,t-1} = 0 \quad (4.5)$$

and number of extinctions E_t estimated as

$$E_t = \sum_{i=1}^M z_{i,t} = 0 | z_{i,t-1} = 1. \quad (4.6)$$

Per-capita growth D^{rate} , origination O^{rate} and extinction E^{rate} rates are then calculated as

$$\begin{aligned} O_t^{rate} &= \frac{O_t}{N_{t-1}^{stand}} \\ E_t^{rate} &= \frac{E_t}{N_{t-1}^{stand}} \\ D_t^{rate} &= O_t^{rate} - E_t^{rate}. \end{aligned} \quad (4.7)$$

4.3 Results

The results of the analyses described above take one of two forms: direct inspection of parameter posterior estimates from both models, and downstream estimates of diversity and diversification rates based on posterior predictive simulations from the birth-death model because this model has a better fit to the observed occurrence information.

4.3.1 Comparison of estimates from the pure-presence and birth-death models

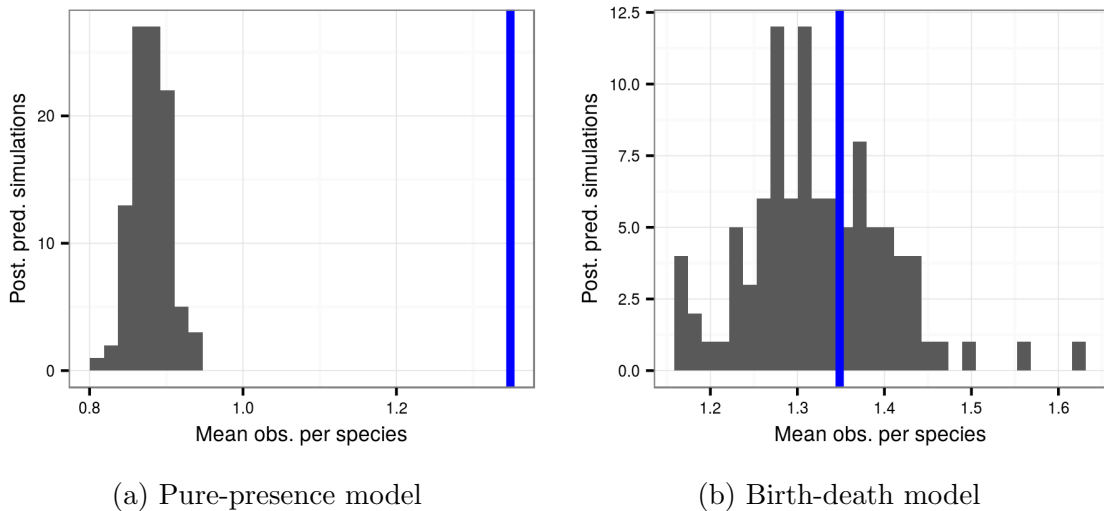


Figure 4.3: Comparison of the average observed number of occurrences per species (blue line) to the average number of occurrences from 100 posterior predictive datasets using the posterior estimates from the pure-presence and birth-death models.

Comparison of the posterior predictive results from the pure-presence and birth-death models reveals a striking difference in performance of either model to predict the structure of the underlying data (Fig. 4.3). The simulated datasets generated from the birth-death model are clearly able to better reproduce the observed average number of occurrence than the pure-presence model which underestimates the observed average number of occurrences.

This result means that inferences based on the birth-death model are more likely to be representative of the underlying data than inferences based on the pure-presence model. Further inspection of the posterior parameter estimates from both models gives further insight into the reasons for this difference in posterior predictive results [101].

Increases in the occurrence probability of an ecotype is interpreted as an increase in the commonness of that ecotype in the species pool. In turn, decreases in the occurrence probability of an ecotype are interpreted to a decrease in the commonness of that ecotype in the species pool. Additionally, when the uncertainty surrounding a probability estimate is very high, as with arboreal insectivores, this is interpreted as complete separation which means that that ecotype has most likely all but disappeared from the species pool [102]. In logistic regression, high uncertainty in the estimates of the underlying log-odds of occurrence, origination, or survival tends to indicate extreme rarity or complete absence of the specific ecotype. The latter is called complete separation and occurs when there is no uncertainty in the effect of a covariate on presence/absence. The problem of complete separation is mitigated by the hierarchical modeling strategy used here [101, 102, 195].

Estimates of occurrence probability estimated from the pure-presence model and estimates of origination probability from the birth-death model are broadly similar (Fig. 4.4, 4.5); this is not the case for the survival probability estimates (Fig. 4.6). This result supports the idea that changes to the North American regional species pool is more likely due to changes in origination than extinction, a result to which I will return to later in the discussion of per-capita diversification, origination, and extinction rates. For most ecotypes, occurrence and origination probability estimates increase with time (Fig. 4.5). This makes sense given that, over time, all species that have at least one observed occurrence must have had that occurrence by the last time point, so our certainty in a species occurring must increase with time. Notably, ecotypes with arboreal components do not appear to follow the same pattern

as most other ecotypes; instead, origination probabilities appear relatively flat with high posterior variance for most of the Cenozoic. For most ecotypes, occurrence or origination probability is estimated with less uncertainty than its estimate of survival probability (Fig. 4.4, 4.5, 4.6).

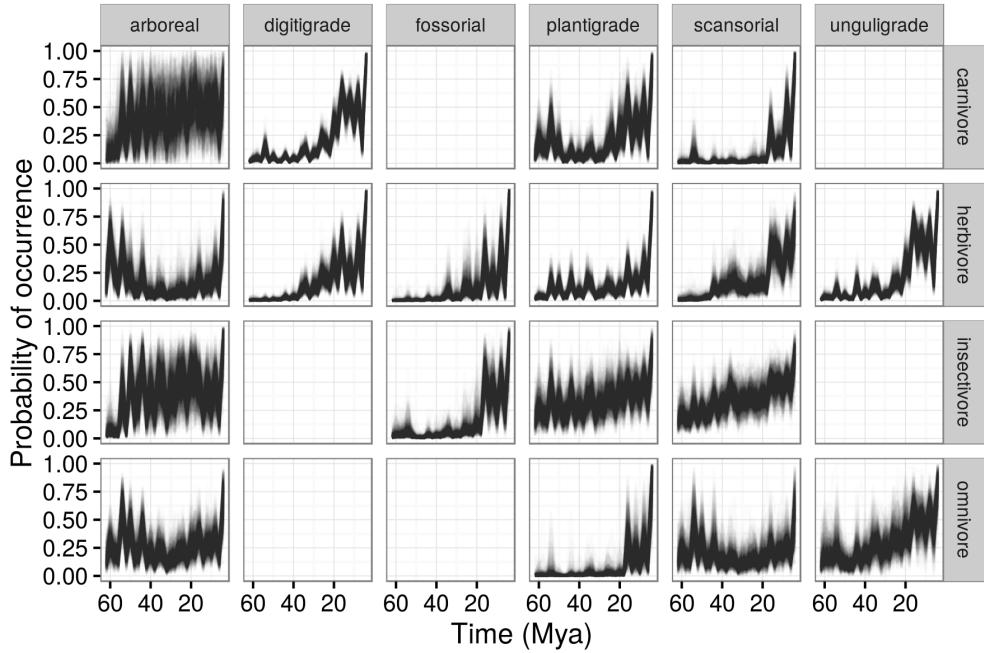


Figure 4.4: Probability of a mammal ecotype occurring over time as estimated from the pure-presence model. Each panel depicts 100 random samples from the model’s posterior. The columns are by locomotor category and rows by dietary category; their intersections are the observed and analyzed ecotypes. Panels with no lines are ecotypes not observed in the dataset.

The pure-presence and birth-death models have similar estimates of the relationship between species mass and the probability of sampling a species that is present (Fig. 4.7). For both models this relationship is at least weakly positive, which means that as species body mass increases it is expected that they are more likely to be sampled if present. The estimated relationship from the pure-presence model is with greater uncertainty than that from the birth-death model (Fig. 4.7). These results are consistent with the intuition that larger fossils are easier to sample because they are more visible to the eye. In turn, this means that the observed occurrence histories of small bodied species are more likely to have gaps, where

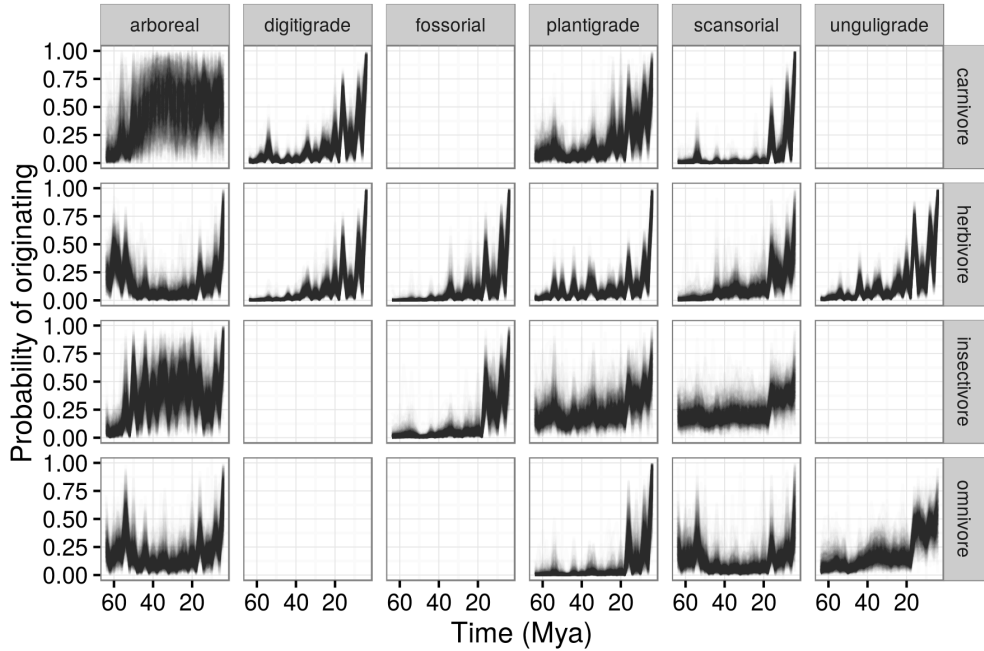


Figure 4.5: Probability of a mammal ecotype origination probabilities at each time point as estimated from the birth-death model. Each panel depicts 100 random samples from the model’s posterior. The columns are by locomotor category and rows by dietary category; their intersections are the observed and analyzed ecotypes. Panels with no lines are ecotypes not observed in the dataset.

$y = 0$ for that species the true state z is 1.

There is broad congruence between the estimated effect of body mass on occurrence probability (Fig. 4.8) and the effect of species mass on body mass on origination probability (Fig. 4.9). The striking pattern is higher probability of origination for species with body sizes closer to the mean than either extremes. This result is consistent with the canonically normal distribution of mammal body sizes [305]; it is then expected that the most likely to occur species would be those from the middle of the distribution, and that species originating will on average be of average mass, especially considering species shared common ancestry [70]. All variation in estimates between ecotypes (Fig. 4.8, 4.9) is due to differences in ecotype-specific origination probabilities and the associated effects of plant phase; the effect of mass was considered constant for all ecotypes.

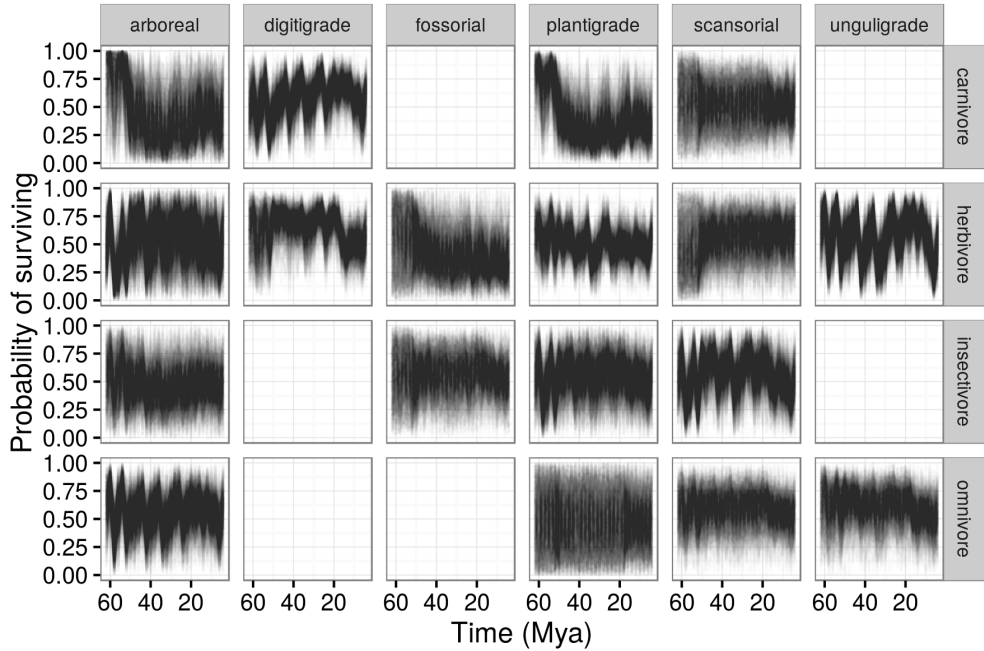


Figure 4.6: Probability of a mammal ectype survival probabilities at each time point as estimated from the birth-death model. Each panel depicts 100 random samples from the model’s posterior. The columns are by locomotor category and rows by dietary category; their intersections are the observed and analyzed ecotypes. Panels with no lines are ecotypes not observed in the dataset.

In contrast, the effect of species mass on probability of survival as estimated from the birth-death model (Fig. 4.10) is consistent with previous findings that there is little effect of mass on extinction for North American mammals for the Cenozoic [309, 329]. Note that all variation between ecotypes depicted in Figure 4.10 is due to differences in ectype-specific survival probability and the associated effects of plant phase; the effect of mass was considered constant for all ecotypes (Eqs. 4.2, 4.3).

Similarities in parameter estimates between ecotypes may be due to a similar response to environmental factors (Fig. 4.11, 4.12, and 4.13). The estimated group-level effects on ectype occurrence, origination, or survival are all very different from each other. At best, the effects of temperature on occurrence and origination can be considered congruent (Fig. 4.11, 4.12). As demonstrated in the comparisons of the effect of body mass on occurrence

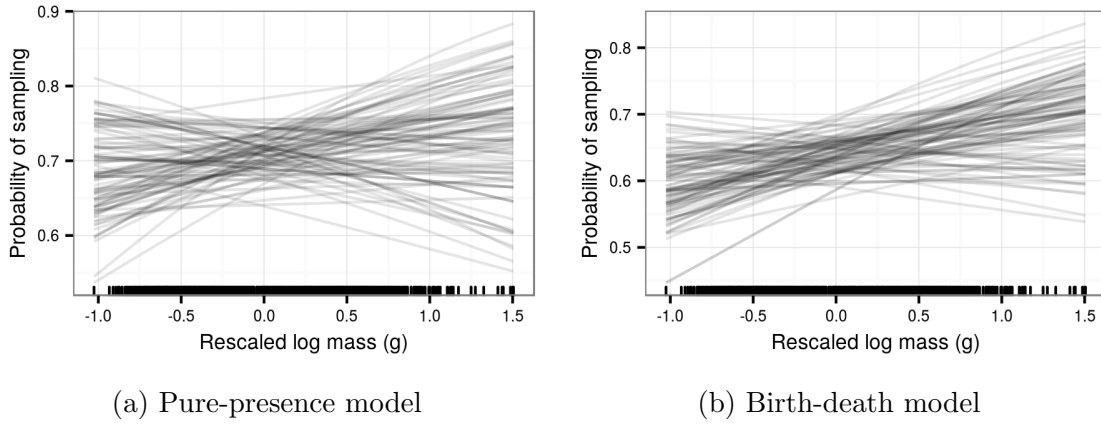


Figure 4.7: Estimates of the effect of species mass on probability of sampling a present species (p). Mass has been log-transformed, centered, and rescaled; this means that a mass of 0 corresponds to the mean of log-mass of all observed species and that mass is in standard deviation units. Estimates are from both the pure-presence and birth-death models.

from the pure-presence model (Fig. 4.8) with the effect of body mass on origination and survival from the birth-death model (Fig. 4.9, and 4.10), there is considerable variation in the effect of plant phases on ecotype-specific estimates.

An association between plant phase and differences in the log-odds of occurrence (Fig. 4.11), origination (Fig. 4.12), or extinction (Fig. 4.13) is interpreted to mean that the set of possible mammal-plant interactions was relatively more favorable (positive association) or less so (negative association) to those ecotypes. In the case of species origination, for example, more favorable conditions for an ecotype may indicate an increasing number of possible and available mammal-plant interactions (e.g. ecological opportunity; 366, 181, 182); while adverse conditions may translate to a decreasing set of interactions or loss of appropriate environmental context. Remember that favorable versus adverse condition of a plant phase is definitionally relative to the other two plant phases.

One of the limitations to this interpretation is the almost deterministic increase in probability of occurrence and origination for most ecotypes (Fig. 4.4, 4.5). This “pull of the Recent” means that interpreting the biological meaning of differences between the final plant phase

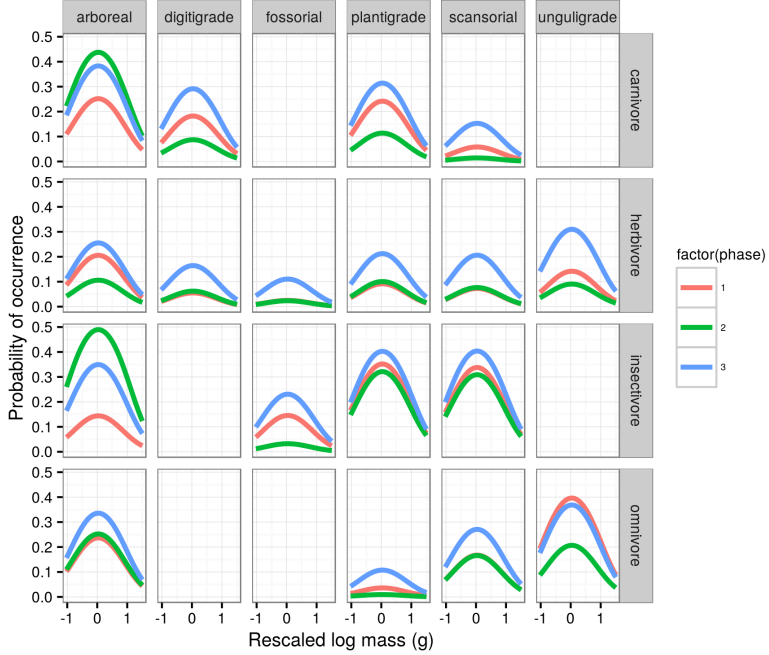


Figure 4.8: Mean estimate of the effect of species mass on the probability of a species occurrence for each of the three plant phases. The effect of mass is considered constant over time and that the only aspect of the model that changes with plant phase is the intercept of the relationship between mass and occurrence. The three plant phases are indicated by the color of the line. Mass has been log-transformed, centered, and rescaled; this means that a mass of 0 corresponds to the mean of log-mass of all observed species and that mass is in standard deviation units. For clarity, only the mean estimates of the effects of mass and plant phase are plotted.

and the two previous phases is difficult as the guaranteed occurrence of the later taxa increases the average probability for that phase, which in turn affects the other time bins in that phase.

Plant phases are associated with large differences in log-odds for occurrence and origination probabilities (Tables 4.9, 4.10), though there is little evidence of plant phase being an important distinguishing factor in species survival, as only a few ecotypes demonstrate strong affinities with some plant phases (Table 4.11).

The effects of plant phase on occurrence and origination probabilities are broadly congruent with each other (Fig. 4.11, 4.12). The almost universal pattern of the effect of plant phase on ecotype origination is that during first and last plant phases ecotypes have a greater

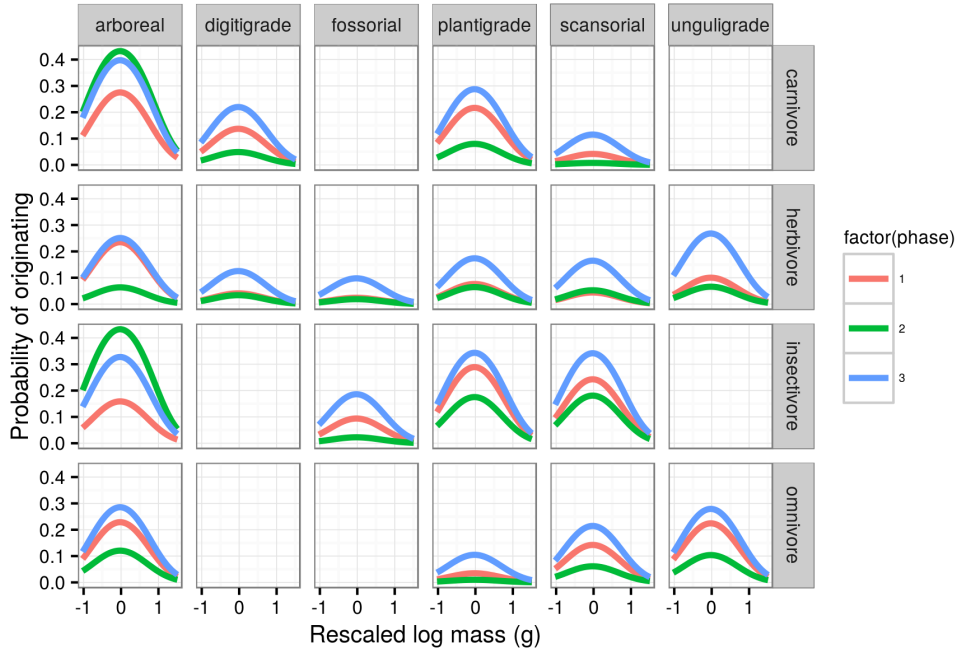


Figure 4.9: Mean estimate of the effect of species mass on the probability of a species originating for each of the three plant phases. The effect of mass is considered constant over time and that the only aspect of the model that changes with plant phase is the intercept of the relationship between mass and origination. The three plant phases are indicated by the color of the line. Mass has been log-transformed, centered, and rescaled; this means that a mass of 0 corresponds to the mean of log-mass of all observed species and that mass is in standard deviation units. For clarity, only the mean estimates of the effects of mass and plant phase are plotted.

log-odds of occurrence or origination than the second plant phase (Fig. 4.4, 4.5). The three ecotypes that do not follow this pattern are fossorial herbivores, scansorial herbivores, and arboreal insectivores.

Both aspects of global temperature analyzed here are estimated to have strong effects on species occurrence and origination for most mammal ecotypes (Tables 4.12, 4.13). Similarly, the probability that temperature has a large effect on species extinction is very low for all ecotypes (Table 4.14). The effects of the temperature covariates on ecotype occurrence and origination are estimated to be negative, which means that as temperature decreases, occurrence and origination are expected to increase. In the case of survival, the only strong

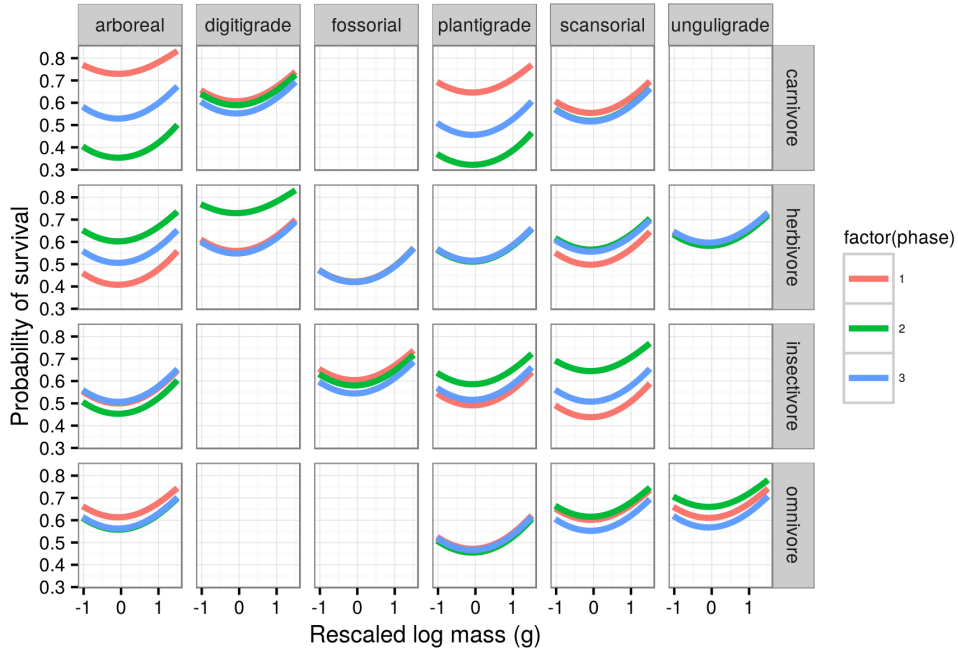


Figure 4.10: Mean estimate of the effect of species mass on the probability of a species survival for each of the three plant phases. The effect of mass is considered constant over time and that the only aspect of the model that changes with plant phase is the intercept of the relationship between mass and survival. The three plant phases are indicated by the color of the line. Mass has been log-transformed, centered, and rescaled; this means that a mass of 0 corresponds to the mean of log-mass of all observed species and that mass is in standard deviation units. For clarity, only the mean estimates of the effects of mass and plant plant are plotted.

ecotype association for either of the temperature covariates is a positive relationship between temperature range and occurrence probabilities of with plantigrade herbivores (Tab. 4.14).

The apparent similarities in origination rate of digitigrade carnivores, digitigrade herbivores, plantigrade herbivores, and unguilgrade herbivores (Fig. 4.5 can be tested by inspecting the estimates of the two correlation matrices Ω^ϕ and Ω^π . The elements of these matrices are estimates of the correlation in origination and survival probabilities, respectively, between each of the 18 observed ecotypes. However, because ADVI-based inference assumes that the joint posterior is Gaussian, estimates of scale and correlation parameters are very approximate as these parameters tend to have decidedly non-Gaussian true posterior distributions [101].

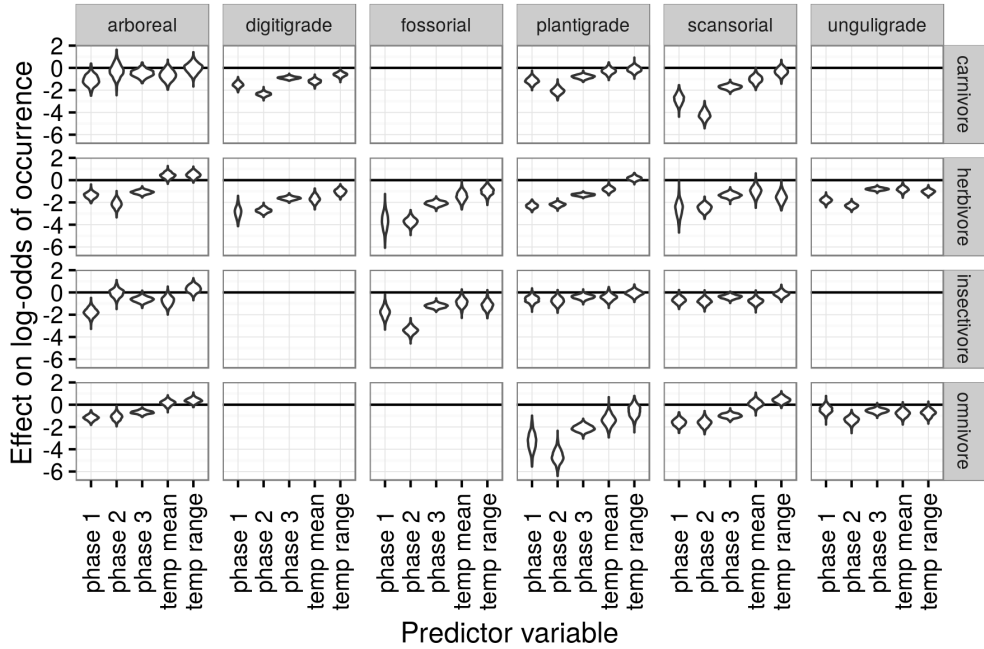


Figure 4.11: Estimated effects of the group-level covariates describing environmental context on log-odds of species occurrence. These estimates are from the pure-presence model. What is plotted is a violin of the distribution of 1000 samples from the approximate posterior. The effect of plant phase graphed here is calculated as Phase 1 = $\gamma_{phase\ 1}$, Phase 2 = $\gamma_{phase\ 1} + \gamma_{phase\ 2}$, and so on.

Because of this fundamental limitation, inference based on these correlation matrices are very approximate and subject to change.

Consistent with visual inspection of the ecotype origination probability time series, there are some strong positive correlations between a few ecotypes (Fig. 4.14). Given the posterior distribution of correlation estimates, the probability of these correlation estimates being greater than 0 can be estimated. Again, because of the assumed Gaussian posterior, these probability estimates are at best approximate and are subject to change given full posterior inference. To visualize these results, I've plotted an association graph of the correlations between ecotypes that are estimated to have a greater than 95% posterior probability of being greater than 0 (Fig. 4.15); in total there are 35 correlations that fit this criterion. The ecotypes correlated with the most number of other ecotypes, in order from most to least, are

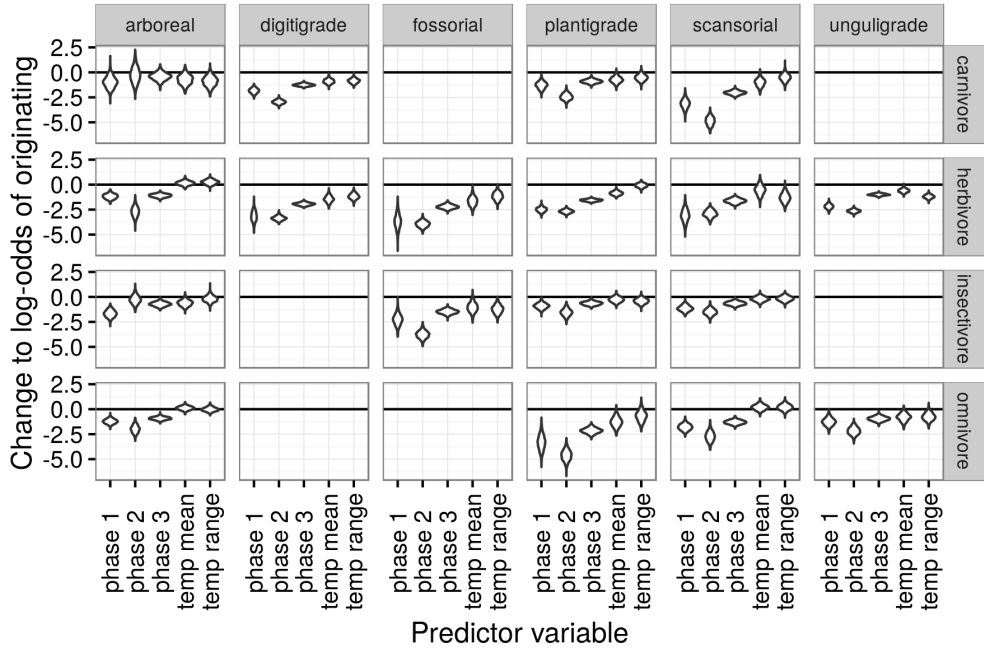


Figure 4.12: Estimated effects of the group-level covariates describing environmental context on log-odds of species origination. These estimates are from the birth-death model. What is plotted is a violin of the distribution of 1000 samples from the approximate posterior. The effect of plant phase graphed here is calculated as Phase 1 = $\gamma_{phase\ 1}$, Phase 2 = $\gamma_{phase\ 1} + \gamma_{phase\ 2}$, and so on.

unguligrade herbivores, plantigrade herbivores, digitigrade carnivores, scansorial carnivores, fossorial herbivores, and plantigrade omnivores. These results support the conclusion that the origination probabilities of many ecotypes are correlated which I interpret to mean that changes to the species pool's environment context can lead to the simultaneous enrichment of multiple ecotypes instead of each ecotype responding independently. This result is most obvious for digitigrade carnivores, scansorial carnivores, digitigrade herbivores, fossorial herbivores, plantigrade herbivores, and unguilgrade herbivores; these ecotypes are heavily cross-correlated with each other.

In contrast to the visually obvious correlations in ecotype origination probability, visual inspection of the ecotype-specific survival probabilities (Fig. 4.6) does not indicate that many strong correlations between ecotype survival probabilities. This conclusion is supported

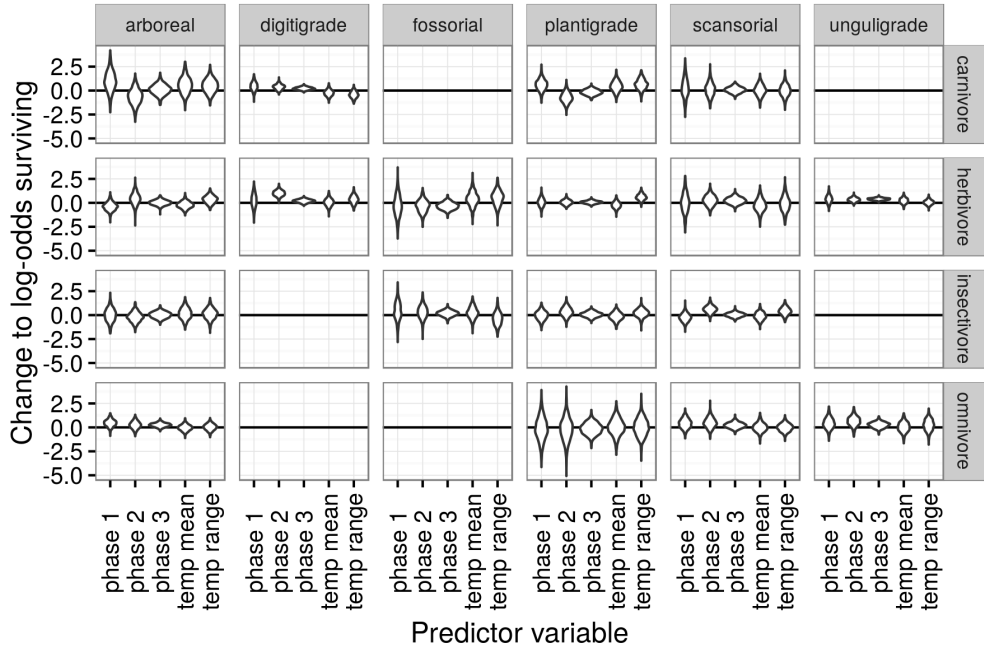


Figure 4.13: Estimated effects of the group-level covariates describing environmental context on log-odds of species survival. These estimates are from the birth-death model. What is plotted is a violin of the distribution of 1000 samples from the approximate posterior. The effect of plant phase graphed here is calculated as Phase 1 = $\gamma_{phase\ 1}$, Phase 2 = $\gamma_{phase\ 1} + \gamma_{phase\ 2}$, and so on.

by the estimated correlations in ecotype survival probability (Fig. 4.16). There are very few large magnitude estimates of correlation between any of the ecotypes. This result is further supported by the fact that only a single correlation, survival probability of digitigrade carnivores and unguiligrade herbivores, has a greater than 95% posterior probability of being positive (Fig. 4.17). This single correlation, however, adds more nuance to the interpretations from the origination probability correlations. In addition to correlation in enrichment, these ecotypes are correlated in their depletions. This result supports the conclusion that the diversity histories of digitigrade carnivores and unguiligrade herbivores are strongly related to each other both in terms of origination and survival, which stands in contrast to those ecotypes which are only correlated for origination.

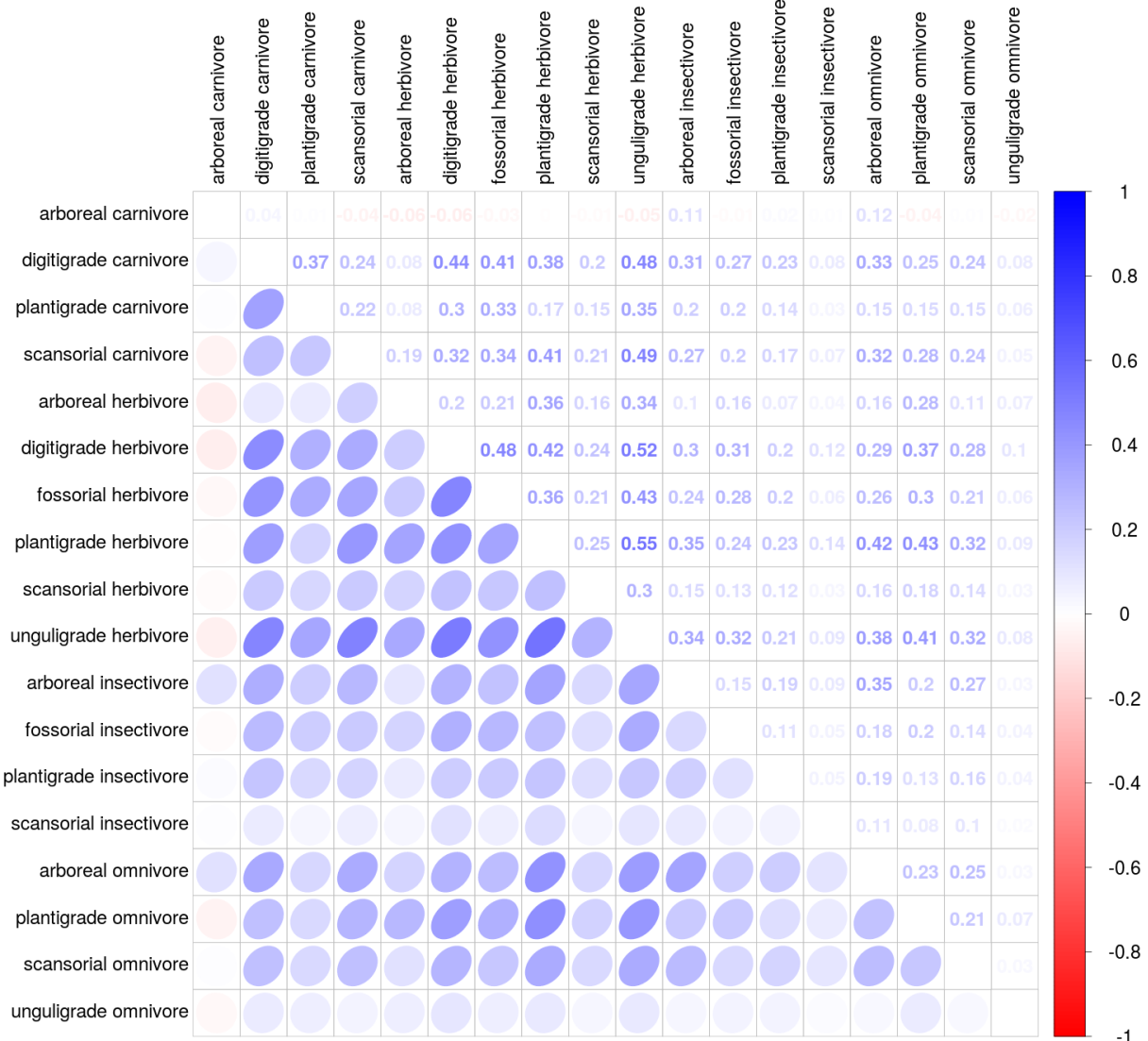


Figure 4.14: Posterior mean estimates of the correlations in origination probability between the mammal ecotypes. The lower triangle of the matrix is populated with ellipses corresponding to the level of correlation between the two ecotypes, while the upper triangle of the matrix corresponds to the mean estimated correlation between ecotypes. Darker values correspond to a greater magnitude of correlation with blue values corresponding to a positive correlation and red values a negative correlation.

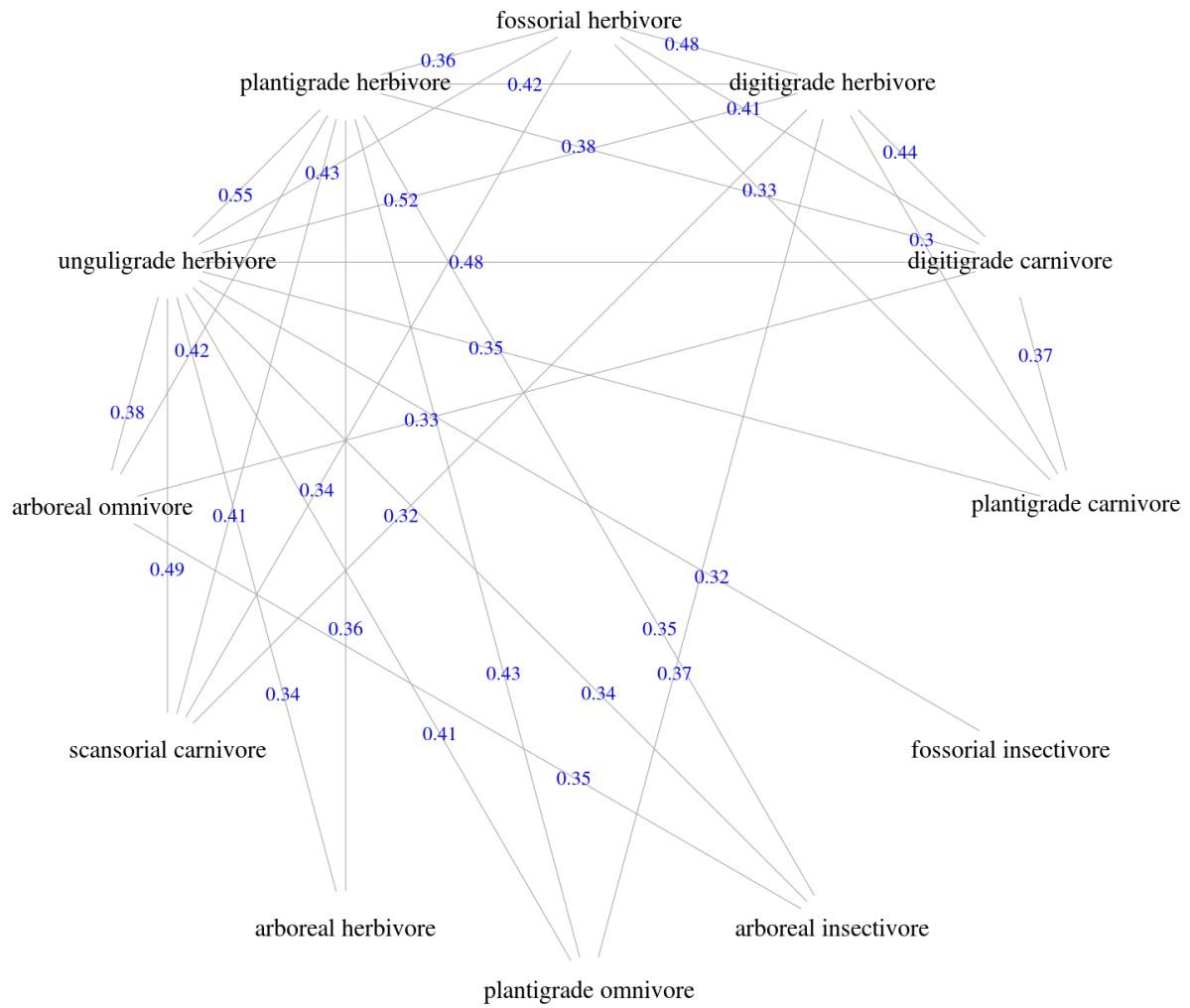


Figure 4.15: Ecotypes that have a strong correlation in origination probability. These ecotypes have a greater than 95% posterior probability of being positively correlated. The number plotted at the midpoint of each edge corresponds to the mean estimated correlation between those two ecotypes.

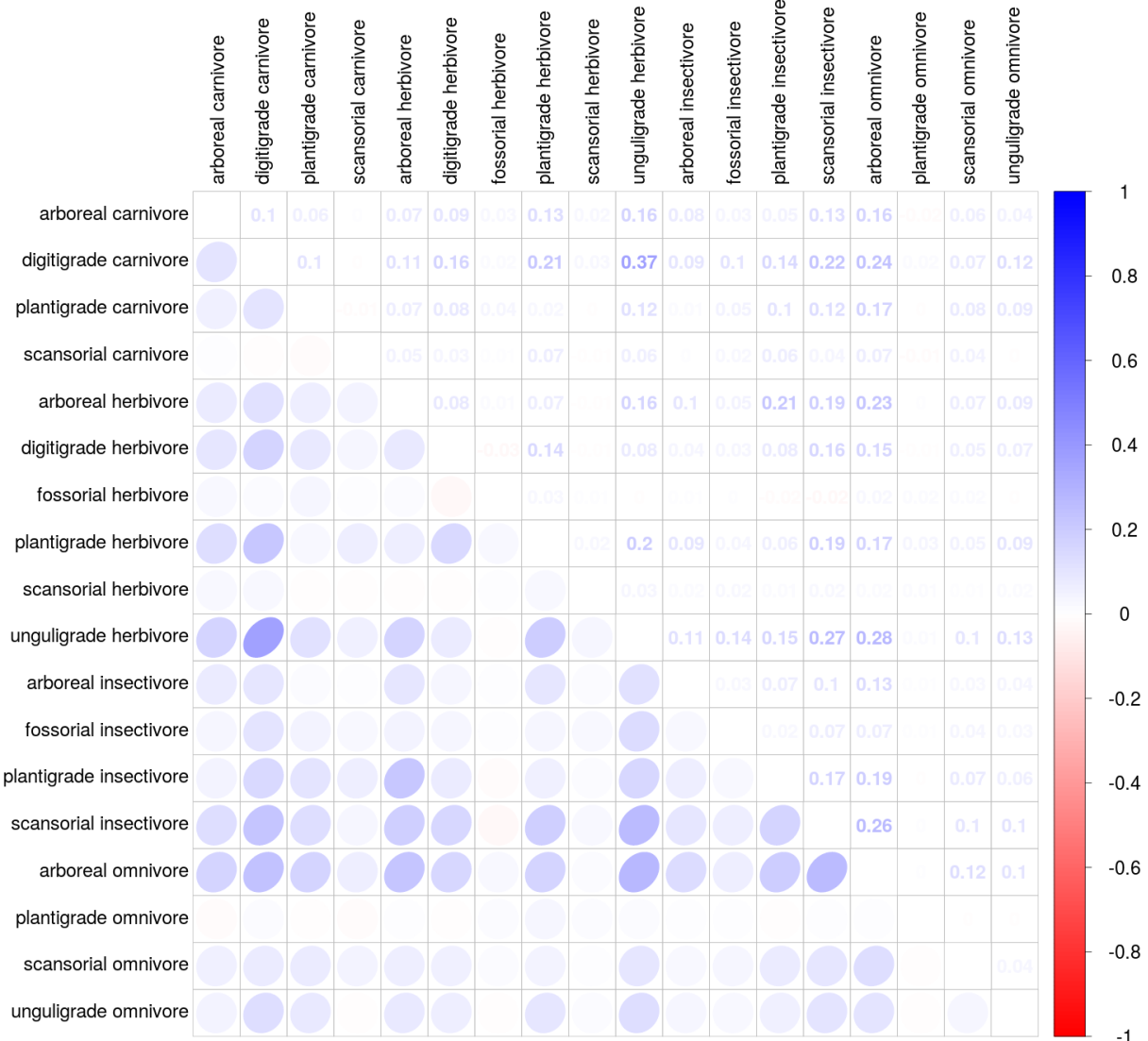


Figure 4.16: Posterior mean estimates of the correlations in survival probability between the mammal ecotypes. The lower triangle of the matrix is populated with ellipses corresponding to the level of correlation between the two ecotypes, while the upper triangle of the matrix corresponds to the mean estimated correlation between ecotypes. Darker values correspond to a greater magnitude of correlation with blue values corresponding to a positive correlation and red values a negative correlation.

Table 4.9: Posterior probability of the differences in the log-odds of an ecotype occurring based on plant phase. These probabilities are calculated as $P(\text{Phase 1} > 2) = (\sum \gamma_{\text{phase1}} > \gamma_{\text{phase1}} + \gamma_{\text{phase2}})/100$ and similarly for the other comparisons. These estimates are from the pure-presence model.

	$P(\text{Phase 1} > \text{Phase 2})$	$P(\text{Phase 2} > \text{Phase 3})$	$P(\text{Phase 1} > \text{Phase 3})$
arboreal carnivore	0.323	0.874	0.926
digitigrade carnivore	1.000	0.000	1.000
plantigrade carnivore	1.000	0.040	1.000
scansorial carnivore	1.000	0.015	1.000
arboreal herbivore	1.000	0.515	1.000
digitigrade herbivore	1.000	0.995	1.000
fossorial herbivore	1.000	0.923	1.000
plantigrade herbivore	1.000	0.995	1.000
scansorial herbivore	1.000	0.717	1.000
unguligrade herbivore	1.000	0.000	1.000
arboreal insectivore	0.024	0.999	0.997
fossorial insectivore	1.000	0.000	1.000
plantigrade insectivore	0.923	0.558	0.985
scansorial insectivore	0.982	0.483	0.992
arboreal omnivore	0.959	0.837	1.000
plantigrade omnivore	1.000	0.247	1.000
scansorial omnivore	0.983	0.861	1.000
unguligrade omnivore	1.000	0.189	0.998

Table 4.10: Posterior probability of the differences in the log-odds of an ecotype originating based on plant phase. These probabilities are calculated as $P(\text{Phase 1} > 2) = (\sum \gamma_{\text{phase1}} > \gamma_{\text{phase1}} + \gamma_{\text{phase2}})/100$ and similarly for the other comparisons. These estimates are from the birth-death model.

	$P(\text{Phase 1} > \text{Phase 2})$	$P(\text{Phase 2} > \text{Phase 3})$	$P(\text{Phase 1} > \text{Phase 3})$
arboreal carnivore	0.405	0.777	0.905
digitigrade carnivore	1.000	0.043	1.000
plantigrade carnivore	1.000	0.053	1.000
scansorial carnivore	1.000	0.035	1.000
arboreal herbivore	1.000	0.163	1.000
digitigrade herbivore	1.000	0.998	1.000
fossorial herbivore	1.000	0.896	1.000
plantigrade herbivore	1.000	0.996	1.000
scansorial herbivore	1.000	0.884	1.000
unguligrade herbivore	1.000	0.003	1.000
arboreal insectivore	0.088	0.994	1.000
fossorial insectivore	1.000	0.020	1.000
plantigrade insectivore	0.995	0.419	1.000
scansorial insectivore	0.999	0.360	1.000
arboreal omnivore	0.999	0.317	1.000
plantigrade omnivore	1.000	0.308	1.000
scansorial omnivore	0.999	0.418	1.000
unguligrade omnivore	1.000	0.219	1.000

Table 4.11: Posterior probability of the differences in the log-odds of an ecotype surviving based on plant phase. These probabilities are calculated as $P(\text{Phase 1} > 2) = (\sum \gamma_{\text{phase1}} > \gamma_{\text{phase1}} + \gamma_{\text{phase2}})/100$ and similarly for the other comparisons. These estimates are from the birth-death model.

	$P(\text{Phase 1} > \text{Phase 2})$	$P(\text{Phase 2} > \text{Phase 3})$	$P(\text{Phase 1} > \text{Phase 3})$
arboreal carnivore	0.849	0.127	0.313
digitigrade carnivore	0.292	0.411	0.162
plantigrade carnivore	0.926	0.243	0.790
scansorial carnivore	0.409	0.544	0.453
arboreal herbivore	0.250	0.733	0.480
digitigrade herbivore	0.000	0.990	0.246
fossorial herbivore	0.547	0.680	0.812
plantigrade herbivore	0.404	0.555	0.480
scansorial herbivore	0.534	0.277	0.204
unguligrade herbivore	0.600	0.046	0.003
arboreal insectivore	0.673	0.379	0.539
fossorial insectivore	0.464	0.442	0.308
plantigrade insectivore	0.216	0.714	0.446
scansorial insectivore	0.019	0.923	0.382
arboreal omnivore	0.394	0.440	0.242
plantigrade omnivore	0.582	0.542	0.677
scansorial omnivore	0.292	0.590	0.289
unguligrade omnivore	0.212	0.555	0.183

Table 4.12: Posterior probabilities that the effects of the two temperature covariates on the log-odds of an ecotype occurring are greater than 0. What is estimated is the probability that these estimates are greater than 0; high or low probabilities indicate the “strength” of the covariate in that direction (positive and negative, respectively). These estimates are from the pure-presence model.

	$P(\gamma_{temp\ mean} > 0)$	$P(\gamma_{temp\ range} > 0)$
arboreal carnivore	0.067	0.043
digitigrade carnivore	0.000	0.000
plantigrade carnivore	0.012	0.054
scansorial carnivore	0.007	0.086
arboreal herbivore	0.762	0.852
digitigrade herbivore	0.000	0.000
fossorial herbivore	0.000	0.002
plantigrade herbivore	0.000	0.364
scansorial herbivore	0.141	0.004
unguligrade herbivore	0.001	0.000
arboreal insectivore	0.031	0.251
fossorial insectivore	0.014	0.002
plantigrade insectivore	0.164	0.058
scansorial insectivore	0.197	0.250
arboreal omnivore	0.711	0.449
plantigrade omnivore	0.009	0.103
scansorial omnivore	0.754	0.732
unguligrade omnivore	0.015	0.022

Table 4.13: Posterior probability that the effects of the two temperature covariates on the log-odds of an ecotype origination are greater than 0. What is estimated is the probability that these estimates are greater than 0; high or low probabilities indicate the “strength” of the covariate in that direction (positive and negative, respectively). These estimates are from the birth-death model.

	$P(\gamma_{temp\ mean} > 0)$	$P(\gamma_{temp\ range} > 0)$
arboreal carnivore	0.072	0.031
digitigrade carnivore	0.000	0.001
plantigrade carnivore	0.010	0.094
scansorial carnivore	0.006	0.104
arboreal herbivore	0.785	0.881
digitigrade herbivore	0.000	0.000
fossorial herbivore	0.001	0.007
plantigrade herbivore	0.000	0.626
scansorial herbivore	0.097	0.002
unguligrade herbivore	0.010	0.000
arboreal insectivore	0.018	0.319
fossorial insectivore	0.008	0.003
plantigrade insectivore	0.223	0.084
scansorial insectivore	0.139	0.202
arboreal omnivore	0.646	0.628
plantigrade omnivore	0.013	0.120
scansorial omnivore	0.688	0.735
unguligrade omnivore	0.016	0.027

Table 4.14: Posterior probability that the effects of the two temperature covariates on the log-odds of an ecotype survival are greater than 0. What is estimated is the probability that these estimates are greater than 0; high or low probabilities indicate the “strength” of the covariate in that direction (positive and negative, respectively). These estimates are from the birth-death model.

	$P(\gamma_{temp\ mean} > 0)$	$P(\gamma_{temp\ range} > 0)$
arboreal carnivore	0.751	0.752
digitigrade carnivore	0.161	0.110
plantigrade carnivore	0.851	0.878
scansorial carnivore	0.598	0.510
arboreal herbivore	0.281	0.819
digitigrade herbivore	0.546	0.721
fossorial herbivore	0.751	0.766
plantigrade herbivore	0.316	0.960
scansorial herbivore	0.342	0.429
unguligrade herbivore	0.750	0.418
arboreal insectivore	0.572	0.579
fossorial insectivore	0.648	0.279
plantigrade insectivore	0.477	0.786
scansorial insectivore	0.303	0.845
arboreal omnivore	0.536	0.611
plantigrade omnivore	0.499	0.507
scansorial omnivore	0.544	0.488
unguligrade omnivore	0.520	0.688

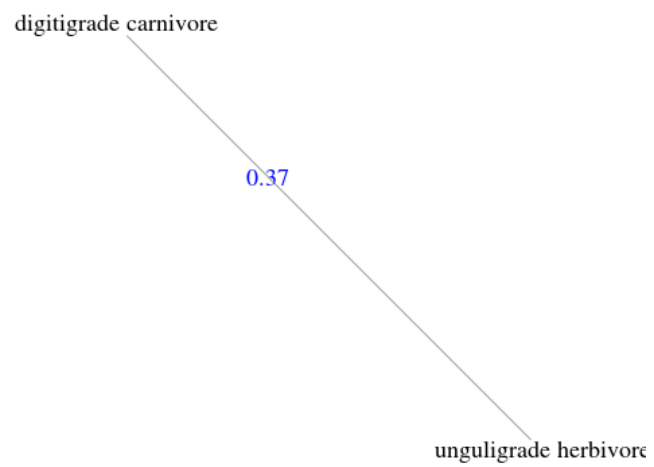


Figure 4.17: Ecotypes that have a strong correlation in survival probability. These ecotypes have a greater than 95% posterior probability of being positively correlated. The number plotted at the midpoint of each edge corresponds to the mean estimated correlation between those two ecotypes.

4.3.2 Analysis of diversity

All of the analyses of diversification and macroevolutionary rates has been done using only the birth-death model because of the model’s better posterior predictive check performance (Fig. 4.3).

The general pattern of the estimated North American total mammal diversity for the Cenozoic is “stable” in that diversity fluctuates around a constant mean standing diversity, does not fluctuate wildly and rapidly over the Cenozoic, and demonstrates no sustained directional trends (Fig. 4.18a). In broad strokes, the first 15 or so million years of the Cenozoic are characterized by first an increase and then a decline in standing diversity at approximately 45–50 Mya (early-middle Eocene). Following this decline, standing diversity is broadly constant from 45 to 18 Mya (early Miocene). After this, there is a rapid spike in diversity followed by a slight decline in diversity up to the Recent.

The pattern exhibited by the diversity history estimated in this study (Fig. 4.18a) has some major similarities with previous mammal diversity curves [6]: both diversity history estimates begin with an increase in diversity and most of the major increases in diversity are retained including the large diversity spike during the Miocene. Unlike subsampling based approaches to estimating diversity [7], I’m able to interpolate over unsampled/poorly sampled time periods because of how the hierarchical model can share information across the different units [101]; for cases like unsampled temporal bins, this may lead to estimates with high uncertainty, but that is preferable to no estimate at all. Finally, the Bayesian framework here gives a distribution of possible estimates of diversity allowing for direct inspection of the uncertainty of our inferences, something that is preferable to both traditional and resampling based confidence interval estimates [101]. Note that my time series of estimated diversity begins at a slightly different point than that used by Alroy [6] and that the time intervals used by Alroy [6] are slightly shorter than those used here, so this may cause some of the

minor differences between the curves. Also, please note that the diversity values are plotted at the “ceiling” of each temporal interval and not at the midpoint (Fig. 4.18a).

When viewed through the lens of diversification rate, some of the structure behind the estimated diversity history begins to take shape (Fig. 4.18b). For most of the Cenozoic, the diversification rate hovers around zero, punctuated by both positive and negative spikes. The largest spike in diversification rate is at 16 Mya, which is early Oligocene (Fig. 4.18b). Other notable increases in diversification rate occur at 56, 46, 22, 18, and 6 Mya (Table 4.15), though the last of these may be due to edge effects surrounding the partial-identifiability of $p_{t=T}$. Notable decreases in diversification rate occur at 54, 50, 48, 44, 40, 34, 30, 24, 20, 16, 12, and 8 Mya (Table 4.15), meaning that diversification rate has more major decreases than increases. While diversification rates significantly lower than average are more common than diversification rates greater than average, when positive diversification rates have a greater magnitude than most periods of low or negative diversification (Fig. 4.18b). Given that diversification rate more closely resembles origination rate than extinction rate (Fig. 4.18b, 4.18c, 4.18d), these decreases in diversification rate may be indicative of “depletions” (failure to replace extinct taxa) rather than pulses of extinction.

The estimates from this study of per capita origination and extinction rates for the entire species pool (Fig. 4.18c, 4.18d) are very different from the origination and extinction rates estimated by Alroy [6]. The two most striking differences are the very different estimates of extinction rate between the two studies and the very different scales of the origination rate estimates. This may be due to the fundamentally different way these rates are calculated, and how the diversification process was modeled. The per capita rates estimated in this study follow straight from the definition of a per capita rate (e.g. number of originations between time t and $t + 1$ divided by the diversity at time t) while the rates calculated by Alroy [6] are based on log ratios of standing diversity.

The comparison between per capita origination and extinction rate estimates reveals how diversification rate is formed (Fig. 4.18c, 4.18d). As expected given previous inspection of the ecotype specific estimates of origination and survival probabilities from the birth-death model, diversification rate seems most driven by changes in origination rate as opposed to extinction rate. Extinction rate, on the other hand, demonstrates an almost saw-toothed pattern around a constant mean (Fig. 4.18d). These results are broadly consistent with those from previous analyses of North American mammals diversity and diversification [5, 10, 6].

Diversity partitioned by ecotype reveals a lot of the complexity behind the pattern of mammal diversity for the Cenozoic (Fig. 4.19).

Arboreal ecotypes obtain peak diversity early in the Cenozoic and then decline for the rest of the time series, becoming increasingly rare or absent as time approaches the Recent (Fig. 4.19). Arboreal herbivores and omnivores obtain peak diversity at the beginning of the Cenozoic then go into decline while remaining a small part of the species pool, while arboreal carnivores and insectivores obtain peak diversity 52-50 Mya and then quickly decline and become extremely rare or entirely absent from the species pool. This is consistent with increasing extinction risk in the Neogene compared to the Paleogene as proposed by Smits [309].

The diversity of digitigrade and unguligrade herbivores increases over the Cenozoic (Fig. 4.19). In contrast, plantigrade herbivore diversity does not have a single, broad-strokes pattern; instead, diversity increases, decreases, and may have then increased till the Recent. In contrast, fossorial and scansorial herbivores demonstrate a much flatter history of diversity, with a slight increase in diversity that over time is more pronounced among fossorial taxa than scansorial taxa. The expansion of digitigrade and unguligrade herbivores over the Cenozoic is consistent with the gradual expansion of grasslands which these ecotypes are better adapted to than closed environments [28, 326].

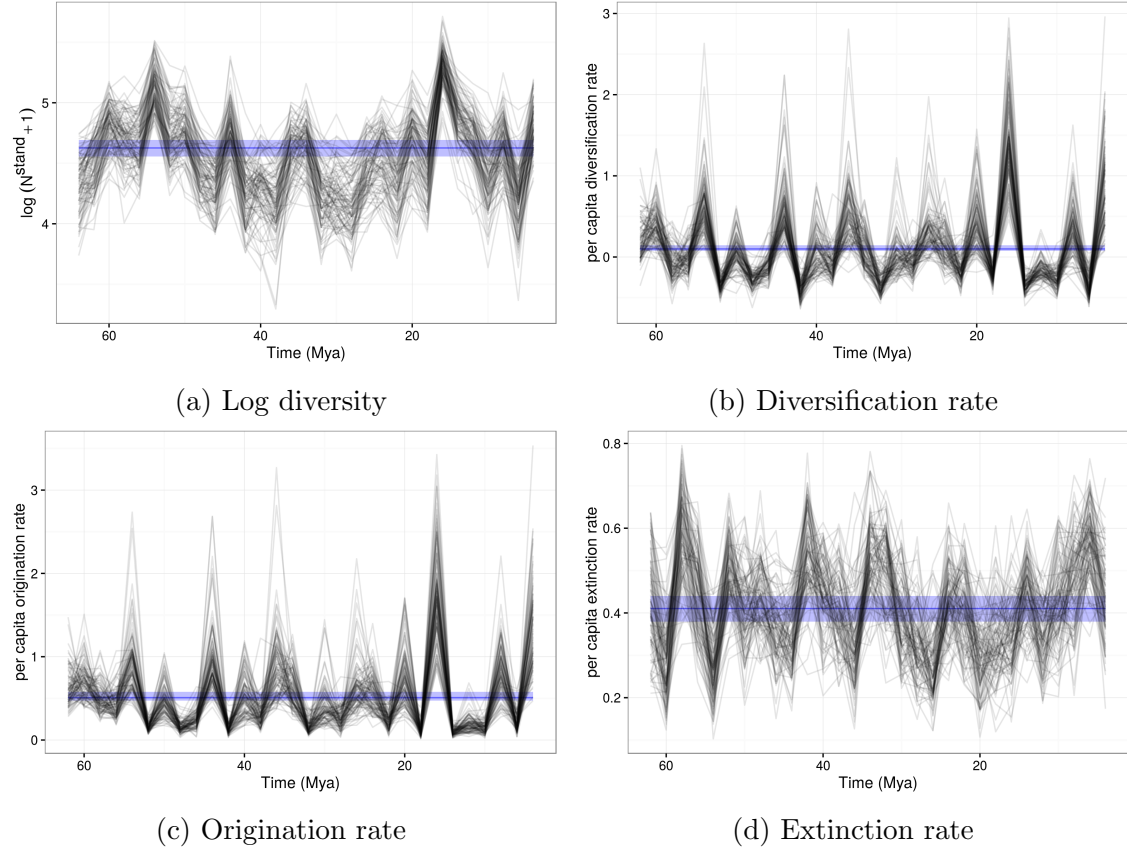


Figure 4.18: Posterior estimates of the time series of Cenozoic North American mammal diversity and its characteristic macroevolutionary rates; all estimates are from the birth-death model and 100 posterior draws are plotted to indicate the uncertainty in these estimates. The blue horizontal strip corresponds to the 80% credible interval of estimated mean standing diversity, diversification rate, origination rate, and extinction rate respectively; the median estimate is also indicated. What is also plotted is the The dramatic differences between diversity estimates at the first and second time points and the penultimate and last time points in this series are caused by well known edge effects in discrete-time birth-death models caused by $p_{-,t=1}$ and $p_{-,t=T}$ being partially unidentifiable [271]; the hierarchical modeling strategy used here helps mitigate these effects but they are still present [101, 271]. Diversification rate is in units of species gained per species present per time unit (2 My), origination rate is in units of species originating per species present per time unit, and extinction rate is in units of species becoming extinct per species present per time unit.

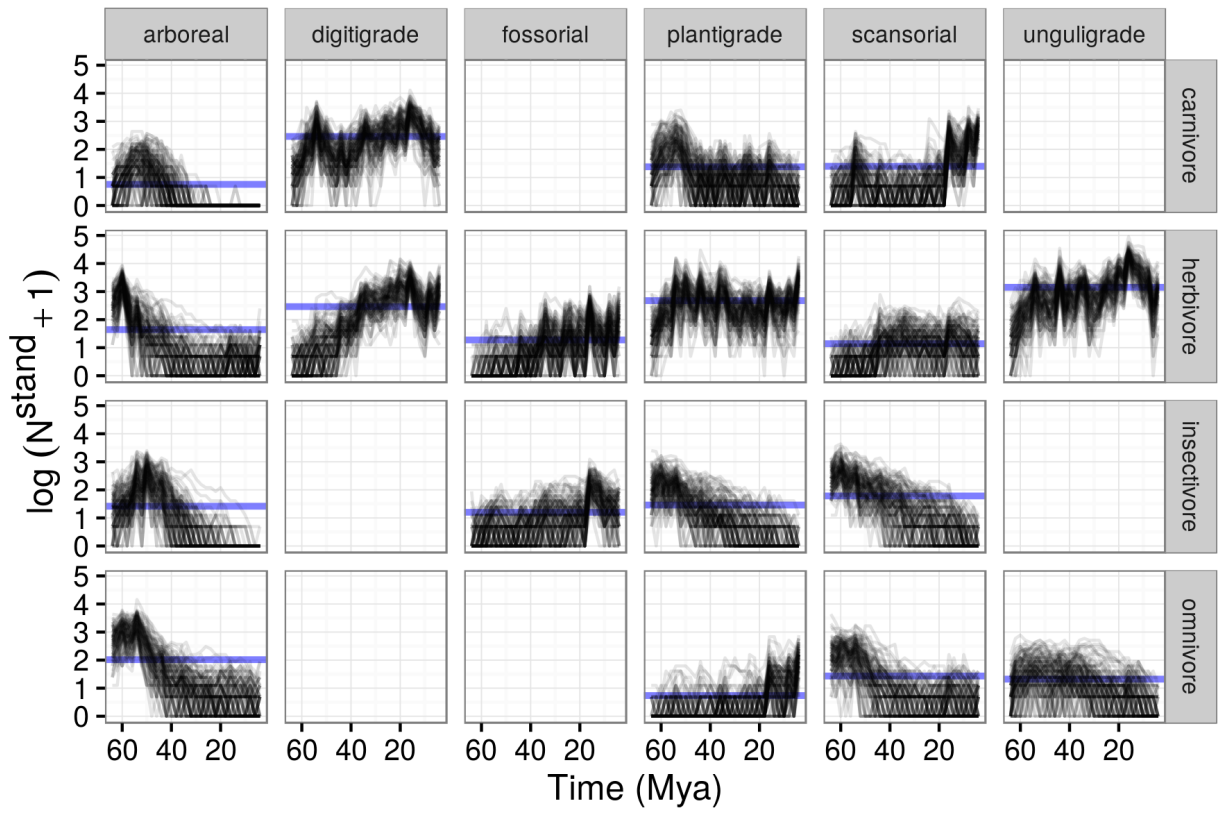


Figure 4.19: Posterior of standing log-diversity of North American mammals by ecotype for the Cenozoic as estimated from the birth-death model; 100 posterior draws are plotted to indicate the uncertainty in these estimates and what is technically plotted is log of diversity plus 1.

Table 4.15: Posterior probabilities of diversity N_t^{stand} or diversification rate D_t^{rate} being greater than average standing diversity $\overline{N^{stand}}$ or average diversification rate $\overline{D^{rate}}$ for the whole Cenozoic. The “Time” column corresponds to the top of each of the temporal bins. Diversification rate can not be estimated for the last time point because it is unknown how many more species originated or went extinct following this temporal bin. The estimates are from the birth-death model.

Time (Mya)	$P(N_t^{stand} > \overline{N^{stand}})$	$P(D_t^{rate} > \overline{D^{rate}})$
64.00	0.07	0.63
62.00	0.28	0.94
60.00	0.86	0.13
58.00	0.68	0.18
56.00	0.62	0.99
54.00	1.00	0.00
52.00	0.68	0.41
50.00	0.80	0.00
48.00	0.12	0.04
46.00	0.01	0.98
44.00	0.64	0.00
42.00	0.02	0.47
40.00	0.03	0.08
38.00	0.00	0.89
36.00	0.40	0.46
34.00	0.52	0.00
32.00	0.02	0.27
30.00	0.06	0.09
28.00	0.02	0.88
26.00	0.22	0.39
24.00	0.38	0.03
22.00	0.09	0.96
20.00	0.81	0.00
18.00	0.29	1.00
16.00	1.00	0.00
14.00	0.95	0.02
12.00	0.80	0.01
10.00	0.13	0.83
8.00	0.67	0.00
6.00	0.02	1.00
4.00	0.91	

Digitigrade carnivores have a multi-modal diversity history, with peaks at 54-52 and 12-10 Mya (Fig. 4.19). Between these two peaks digitigrade carnivore diversity dips below average diversity following the first peak and then grows slowly until the second peak. Plantigrade carnivores obtain peak diversity in the early Cenozoic and then maintain a relatively stable diversity until another peak at the end of the Cenozoic. The generally flat diversity history of digitigrade carnivores lacks any sustained temporal trends and seems to reflect previous findings of limited diversity in spite of constant turnover and morphological evolution [338, 290, 304]

There are some broad similarities in diversity histories of insectivorous and omnivorous taxa. The diversity histories of arboreal, plantigrade, and scansorial insectivorous taxa all demonstrate a decreasing pattern with time, while fossorial insectivores have a flat diversity history with a peak approximately 10 Mya (Fig. 4.19). Arboreal and scansorial omnivores decrease in diversity from their initial peaks early in the Cenozoic, and plantigrade omnivores have a generally flat diversity history with a sudden peak in diversity late in the Cenozoic (Fig. 4.19). Unguligrade omnivores also demonstrate a possible decrease in diversity over the Cenozoic, but not as clearly as arboreal and scansorial omnivores.

The waxing and waning of the mammal ecotypes is obvious when comparing changes to estimated relative log-mean of diversity (Fig. 4.20). While ectype diversity does appear to change gradually, there are definite changes to the relative contributions of the ecotypes to the regional species pool. All arboreal ecotypes clearly decrease in relative diversity over the Cenozoic. In contrast the digitigrade herbivore, fossorial herbivore, scansorial herbivore, and unguligrade herbivore ecotypes which increase in relative diversity over the Cenozoic. The digitigrade carnivore ectype increases in relative diversity until approximately the start of the Neogene, after which it maintains a generally constant relative diversity; this is consistent with previous observations of constant or density-dependent diversity of the canid guild for the Neogene [338, 290, 304], a guild that overlaps with the digitigrade carnivore

ecotype. Plantigrade herbivores remain a constant relative contribution to ecotypic diversity. These results support the hypothesis of a gradual transition from the early Paleogene with a region with more available habitat for arboreal taxa and less available habitat for many digitigrade and unguligrade taxa, to an environment where arboreal taxa are absent from the species pool and digitigrade and unguligrade taxa are much more dominant (Fig. 4.20). It is the relative contributions of digitigrade carnivores, digitigrade herbivores, and unguligrade herbivores which really shape the regional species pool of the Neogene.

Many of the estimated ecotype-specific diversity histories share a similar increase in diversity in the late Cenozoic, 16-14 Mya (Fig. 4.19). These increases are either sustained or temporary and are seen in digitigrade carnivores, plantigrade carnivores, scansorial carnivores, unguligrade herbivores, fossorial insectivores, and plantigrade omnivores.

When ecotype diversity is decomposed into per capita origination (Fig. 4.21) and per capita extinction rates (Fig. 4.22) the way in which their diversity developed can be explained. For ecotype-specific origination and extinction rates, the number of origination or extinction events for each ecotype was calculated and that number was divided by the total standing diversity of all mammals at the time.

As should be expected from the comparisons between estimates of origination and survival probability, origination rates have a much greater range of values with a few very large spikes that line up with the spikes in over all diversification rate (Fig. 4.18b). Importantly, the source of the massive increase in diversification rate at 16 Mya can be attributed almost solely to the origination of unguligrade herbivores (Fig. 4.21). Additionally, by decomposing origination rate by ecotype, it is possible to identify a few possible cross-ecotype increases in origination rate. For example, digitigrade carnivores, digitigrade herbivores, and plantigrade herbivores share a lot of increases in origination rate with unguligrade herbivores; these are all ecotypes that demonstrate an obvious increase in diversity during the Paleogene and then

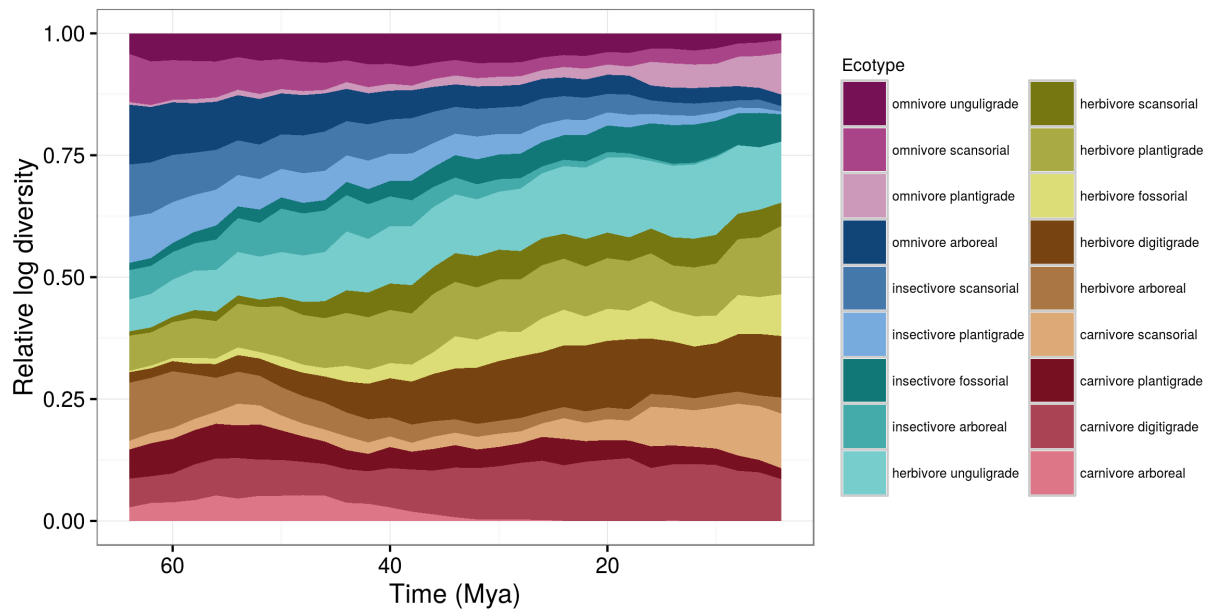


Figure 4.20: Mean posterior estimate of relative log standing diversity of 18 North American mammal ecotypes for the Cenozoic. These estimates are calculated from 100 posterior estimates of the true occurrence matrix z as estimated from the birth-death model.

maintain relatively high diversity through out the Neogene (Fig. 4.19).

In contrast to ecotype-specific per capita origination rates which demonstrate distinct peaks, the estimates of ecotype-specific per capita rates are more of a smear (Fig. 4.22). There are few increases in extinction rate that are shared across ecotypes. The per capita extinction rates of digitigrade, plantigrade, and unguligrade herbivores are lower in Paleogene than the Neogene. This result is interpreted to mean that as the diversity of these three ecotypes was increasing, the number of extinction events was also increasing. Also, the per capita extinction rate of arboreal taxa is higher in the Paleogene than the Neogene. While this result may seem odd considering the observed diversity pattern for these ecotypes (Fig. 4.19), I argue that this result is actually extremely intuitive: if there are no species of that ecotype originating or present, than there can be extinctions. This result highlights the distinction between extinction risk and extinction rate; an ecotype can have a high extinction risk, but if that ecotype is not present in the species pool in the first place than it has no associated extinction rate.

4.4 Discussion

Both the composition of a species pool and its environmental context change over time, though not necessarily at the same rate or concurrently. Local communities, whose species are drawn from the regional species pool, have “roles” in their communities defined by their interactions with a host of biotic and abiotic interactors (i.e. a species’ niche). For higher level ecological characterizations like ecotypes and guilds, these roles are broad and not defined by specific interactions but by the genre of interactions species within that grouping participate in. The diversity of species within an ecotype or guild can be stable over millions of years despite constant species turnover [152, 304, 338]. This implies that the size and scope of the

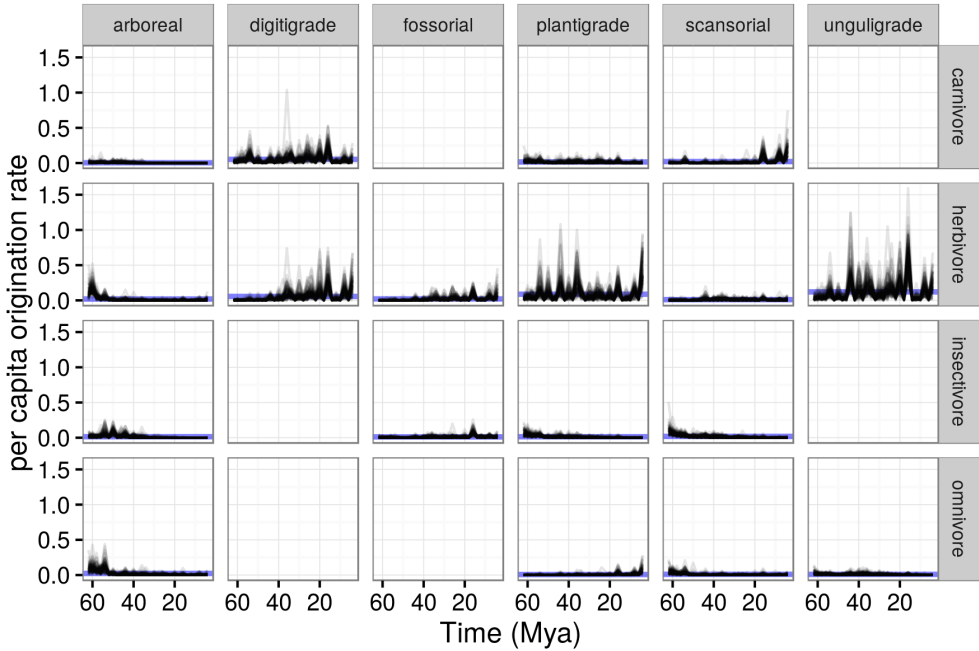


Figure 4.21: Posterior estimates of the per capita origination rates for each ecotype, plotted at the bin they originate from. These rates are calculated as the number of origination events for that ecotype from one time point to the next, divided by the standing diversity of all mammals at the initial time point. 100 posterior draws are plotted to indicate the uncertainty in these estimates.

role of an ecotype or guild in local communities, and the regional species pool as a whole, is preserved even as the individual interactors change. This also implies that the structure of regional species pools can be constant over time despite a constantly changing set of “players.” There is even evidence that functional groups are at least partially self-organizing and truly emergent [274].

Comparison of the results from the posterior predictive checks for the pure-presence and birth-death models supports the conclusion that regional species pool dynamics cannot simply be described by a single occurrence probability and are instead the result of the interplay between the origination and extinction processes. Additionally, changes to the ecotypic composition and diversification rate of the North American regional species pool are driven primarily by variation in origination and not extinction (Fig. 4.18). These aspects

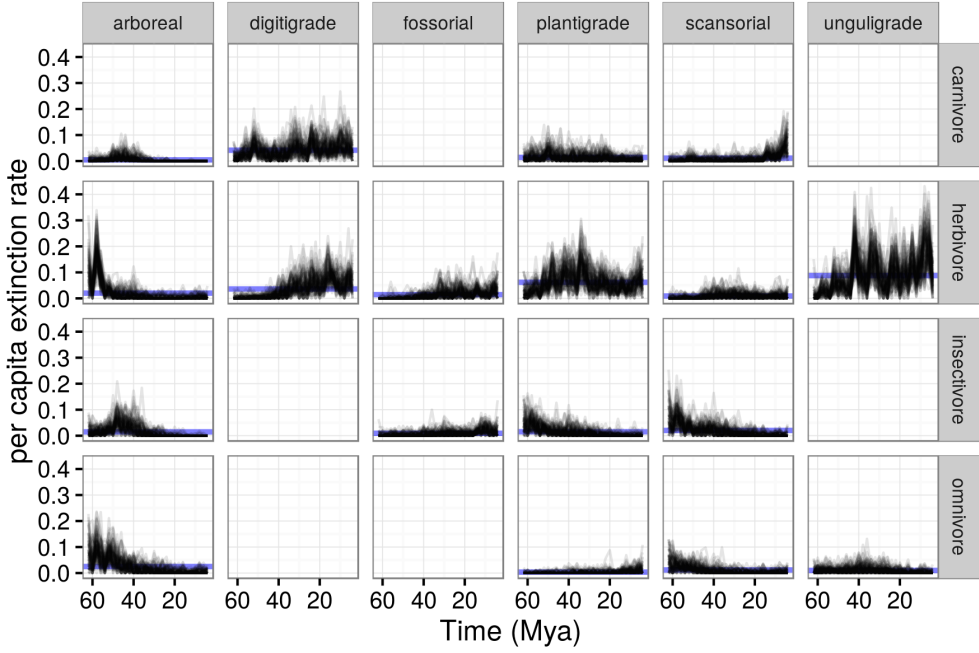


Figure 4.22: Posterior estimates of the per capita extinction rates for each ecotype, plotted at the bin they go extinct from. These rates are calculated as the number of extinction events for that ecotype from one time point to the next, divided by the standing diversity of all mammals at the initial time point. 100 posterior draws are plotted to indicate the uncertainty in these estimates.

of how regional species pool diversity is shaped are not directly observable in studies of the Recent where time scales are short and macroevolutionary dynamics are inferable solely from phylogeny [97, 236].

Extinction rate for the entire regional species pool through time is highly variable and demonstrates a saw-toothed pattern with no obvious temporal trends. While a constant mean extinction rate is consistent with previous observation [5, 10], the degree to which mammal extinction rate is actually variable may not have been equally appreciated as it has been for the marine invertebrate record [79, 80, 83, 84]. What is most consistent with previous observations, however, is that diversity seems to be most structured by changes to origination rather than changes to extinction [5, 10].

Comparison of the ecotype specific diversity histories adds a considerable degree of nuance to

broad narrative of shifts in functional composition of the North American mammal species pool as being gradual (Fig. 4.19). Origination and extinction rates do not suddenly shift through time for most ecotypes (Fig. 4.21, 4.22). As with the diversification rate of the entire species pool, the diversification of individual ecotypes seem principally driven by origination and not extinction. Instead, while species seem to originate in waves (Fig. 4.21), they appear to leave the regional species pool in an uncoordinated and individual manner (Fig. 4.22) which could be considered consistent with the maxim that all species respond differently to environmental change [28]. Note, however, this result characterizes the entire North American mammal regional species pool and thus may not reflect the dynamics of specific local communities but instead represent the average community.

The few large-magnitude, but temporary, increases in ecotype-specific origination rate occur in digitigrade carnivores, digitigrade herbivores, plantigrade herbivores, and unguligrade herbivores. Importantly, the large peak in diversification and origination rates 16 Mya (Fig. 4.18) appears driven almost entirely by a massive increase in the origination rate of unguligrade herbivores (Fig. 4.21). Additionally, there is some evidence that the origination probabilities of these ecotypes are correlated (Fig. 4.14, 4.15). While this result does not mean that there are large and sudden cross-ecotype changes to the regional species pool, it does suggest that additions to the species pool do not occur in individual ecotypes idiosyncratically.

Arboreal taxa disappear from the regional species pool over the Cenozoic, with long term decline over the Paleogene leading to the disappearance by start of Neogene ~22 Mya. This is partially consistent with one of the two possible patterns presented here and by Smits [309] that would result in arboreal taxa having a greater extinction risk than other ecotypes: the Paleogene and Neogene were different selective regimes and, while the earliest Cenozoic may have been neutral with respect to arboreal taxa, they disappeared quickly over the Cenozoic which may account for their higher extinction risk. However, these result add some nuance to

this scenario as arboreal taxa were declining throughout the Paleogene instead of maintaining a flat diversity as hypothesized [309]. I interpret the decline of arboreal taxa throughout the Paleogene to mean that the shift from closed to open environments began in the Paleogene and led to increasingly hostile environments for arboreal taxa as opposed to being a sudden change in selective regime between the Paleogene and Neogene. In addition to all arboreal taxa, the diversity of plantigrade and scansorial insectivores decreases with time (Fig. 4.19).

Digitigrade carnivores have a relatively stable diversity history through the Cenozoic and can be characterized as varying around a constant mean diversity. This ecotype has a large amount of overlap with the carnivore guild which has been the focus of much research [304, 338, 230, 147]. This result is consistent with some form of “control” on the diversity of this ecotype, such as diversity-dependent diversification [304, 290, 338].

Both digitigrade and unguligrade herbivores increase in diversity over the Cenozoic. The increase of these cursorial forms is consistent with the gradual opening up of the North American landscape [28, 326, 109] and may indicate a long-term shift in the interactors associated with those ecotypes leading to increased contribution to the regional species pool. This result may be comparable to the increasing percentage of hypsodont (high-crowned teeth) mammals in the Neogene of Europe being due to an enrichment of hyposodont taxa and not a depletion of non-hypsodont taxa. Smaller scale increases in fossorial herbivore species, and a lesser extent plantigrade herbivores, suggests that the increase of interactors may be associated mostly with the herbivore dietary category with locomotor category tempering that relationship. These results support the conclusion that the increase in digitigrade and unguligrade herbivores is the result of an enrichment of these ecotypes as opposed to being caused by the depletion of other herbivorous ecotypes; this is further supported by the lack of major changes to the number of extinctions of all herbivore ecotypes (Fig. 4.22).

An association between plant phase and differences in ecotype occurrence or origination-

extinction probabilities is interpreted to mean that an ecotype enrichment or depletion is due to associations between that ecotype and whatever plants are dominant at that time. Plant phase clearly structures the occurrence and origination probability time series (Fig. 4.4, 4.5). These differences in occurrence or origination translate to the estimates of diversity and diversification rate; the largest spike in diversity, diversification rate, and origination rate all correspond to the onset of the last plant phase (Fig. 4.18). The clearest example of the diversity of an ecotype increasing at this particular transition is in scansorial carnivores (Fig. 4.19); similar shifts in other ecotypes are much more subtle, as was previously noted for fossorial insectivores.

Interestingly, for all of the ecotypes with sudden changes in diversity at this transition the change is an increase, even if only temporarily. There are two interpretations of these results. A biological interpretation of this result is that, because plant phase associations are only with occurrence or origination probabilities and not survival, these ecotypes were well suited for the newly available mammal-plant interactions due to the increased modernization of taxa characteristic of the younger plant phases [109]. Alternatively, the increase in diversity associated with the third plant phase may be caused by the edge effect in origination probability that is artificially inflating the number of origination events (Fig. 4.5). However, the estimated number of origination events does not have a tremendous spike at this transition, nor is a major increase in the number of origination events sustained (Fig. 4.21).

There are fewer, less obvious shifts in diversity surrounding the transition from the first to second plant phase, with the following ecotypes having apparent shifts in diversity at 50 My: plantigrade carnivores (down), arboreal omnivores (down), and scansorial omnivores (down). Arboreal insectivore peak diversity also occurs 50 Mya, and is then followed by a steep decline in diversity till 30 Mya when this ecotype is lost from the species pool. Because plant phase has been found to structure occurrence/origination (Fig. 4.4, 4.5), but not survival (Fig. 4.6),

my interpretation of these results is that new species were not entering the system because there were fewer available interactors available for those ecotypes. Instead, these ecotypes were poorly suited for the newly available mammal-plant interactions brought upon by the changing environmental context [109].

The temperature covariates are found to have similar effects on occurrence and origination probabilities (Tables 4.12, 4.13). Temperature is found to more often affect ecotype occurrence probabilities than origination probabilities. In most cases, there is a negative association between temperature and probability of occurring or first originating; this means that if temperature decreases, we would then expect an increase in the probability of occurring or first originating. In contrast, temperature range is estimated to be a good predictor of survival in only two mammal ecotypes and only marginally for one of those (Table 4.14). Additionally, both of these cases have positive relationships, meaning that if temperature decreases it is expected that species survival will also decrease.

The result that temperature does not affect the survival probability of most ecotypes is consistent with previous analysis of mammal diversity [10]. The result that temperature affects origination probability, on the other hand, is in strong contrast to previous results [10]. An important difference between the analyses presented here and those obtained by Alroy [10] is I am considering the effect of temperature on the probability of a species originating, assuming it hasn't originated yet while Alroy [10] analyzed the correlation between the first differences of the origination and extinction rates with an oxygen isotope curve [368]. Origination or extinction rates have very different properties than the origination probabilities estimated here brought upon by the difference both in definition and units. Origination probability is the expected probability that a species that has never been present and is not present at time t will be present at time $t + 1$; origination probability is defined for a single species. In contrast, per capita rates are defined (for origination) as the expected number of

new species to have originated between time t and $t + 1$ given the total number of species present at time t ; per capita rates are defined for the standing diversity. It is also important to note that even though there is an edge effect at the last time interval that causes an increase in the occurrence and origination probabilities of some ecotypes (Fig. 4.4, 4.5, the corresponding rates and population level birth/death dynamics do not share that pattern (Fig. 4.18, 4.21, 4.22). However, it is still possible that the finding that temperature has an effect on origination may simply be because as time approaches the present the number of species which have originated increases and not because of climatic forcing of origination.

All environmental factors are found to affect the occurrence and origination probabilities for most, but not all, mammal ecotypes (Fig. 4.11, 4.12). In contrast, the environmental factors probably do not affect differences in ecotype survival probability (Fig. 4.13). The focus in previous research on temperature and major climatic or geological events without other measures of environmental context may have led to confusion in discussions of how the “environment” affects mammal diversity and diversification [10, 72]. The environment or climate are more than just global or regional temperature, it is also the set of all possible biotic and abiotic interactions that can be experienced by a member of the species pool. By including more descriptors of species’ environmental context than simple an estimate of global temperature a more complete “picture” of the diversification process is inferred.

Analysis of relationship between temperature and origination rate is probably better suited for a continuous-time birth-death or multilevel stochastic differential equation model instead of a discrete-time model because the continuous models estimate rates while discrete time models estimate probabilities [4]. For example, the PyRate model(s) are based on a continuous-time birth-death process [291, 290]. Unfortunately, a continuous-time model may be unsuited for most paleontological data as the fossil record is naturally discrete; fossils are assigned to temporal units, such as stages, which have age ranges. Individual fossils are not assigned

individual numeric ages. This reality was in fact my one of motivations for using discrete-time birth-death model instead of one in continuous-time. There are of course exceptions to this characterization; the fossil record of graptolites from the Ordovician and Silurian [55] and the fossil record of some mammal orders from Neogene are of high enough resolution that the application of continuous-time models is appropriate and less fraught.

The effect of species mass on either occurrence or origination and extinction was not allowed to vary by ecotype or environmental context. The primary reason for this modeling choice was that this study focuses on ecotypic based differences in either occurrence, or origination and extinction. Allowing the effect of body size to vary by ecotype, time, and environmental factors would increase the overall complexity of the model beyond the scope of the study. Instead, body size was included in order to control for its possible underlying effects [195]. A control means that if there is variation due to body mass, having a term to “absorb” that effect is better than ignoring it.

The only covariate allowed to affect sampling probability was mass and only as a linear predictor. Other covariates, such as the environmental factors considered here, could have affected the underlying preservation process that limits sampling probability; their exclusion as covariates of sampling/observation was the product of a few key decisions: model complexity, model interpretability, the scope of this study, and a lack of good hypotheses related to these covariates to warrant their inclusion.

4.4.1 Conclusions

These results add a considerable degree of nuance to the narrative of changes to North American diversity being gradual. My results support the conclusions that ecotypic diversity is shaped more by changes to origination than extinction and that major changes to total

diversification rate can be attributed to increases in origination of only some ecotypes. There are a number of interesting estimated ecotype diversity patterns. While arboreal ecotypes are diverse in the Paleogene, by the Neogene all arboreal ecotypes had dramatically decreased in diversity and became either rare or absent in the regional species pool. The other ecotypes that decrease in diversity over the Cenozoic are plantigrade and scansorial insectivores and scansorial omnivores. The only ecotypes that demonstrate a sustained pattern of increasing diversity are digitigrade and unguligrade herbivores. When the environmental covariates analyzed here are inferred to affect the diversification of an ecotype, this effect is virtually always on origination and not survival. This analysis provides a much more complete picture of North American mammal diversity and diversification, specifically the dynamics of the ecotypic composition of that diversity. By increasing the complexity of analysis while precisely translating research questions into a statistical model, the context of the results is much better understood. Future studies of diversity and diversification should incorporate as much information as possible into their analyses in order to better understand or at least contextualize the complex processes underlying that diversity.

CHAPTER 5

CONCLUSION

Macroevolution and macroecology are disciplines devoted to explaining emergent patterns in evolutionary and ecological data. These disciplines are linked through the analysis of the distribution of trait values across time, space, and/or species [196, 358]. Emergent evolutionary and ecological patterns in time can require at least a million of years to observe [332]. Paleontological and phylogenetic data preserve aspects of these large scale temporal patterns. Paleontological data is unique however in being empirical observations of these dynamics while phylogenetic data only preserves the branching history leading to the diversity pattern exhibited by the tips. Additionally, phylogenies of only extant taxa are most likely unrepresentative of the evolutionary history of that clade; for example, multiple rounds of extinction and radiation can obscure the actual diversification process from possibly being inferred [205, 204, 174, 239].

In the studies presented as a part of this dissertation, I have emphasized functional traits. These are traits which directly relate to the way in which a taxon interacts with its environment [196]. When these traits are defined for a species, they are called species traits [196]; examples include species geographic range, average body size, trophic level, environmental preference, etc. Functional traits are an excellent window in macroevolution and macroecology because of their obvious selective importance: an organism which cannot interact with its environment is by definition maladapted. Additionally, the simultaneous analysis of multiple functional traits improves overall process-based inference because it more accurately accounts for the ways in which species interact.

5.1 Summary

All of the studies conducted here were analyses of fundamentally emergent patterns which are not reducible to the properties of their constituents. In each of these studies, hypotheses and analysis were framed in terms of how a species functional ecology can be associated with or shape these emergent patterns. Because of the complexity of processes which shape these emergent patterns, as well as the vagaries of the fossil record, each of these studies required the development of specific inference devices (e.g. statistical models) which attempt to estimate the actual quantities of interest to that analysis.

The emergent pattern at the heart of both the first and second studies (Chapters 2, 3) is species duration. A species endures because of the continued success of individuals of that species but the duration of that species is only knowable by integrating across all individuals. The third study (Chapter 4) deals with a fundamentally different emergent pattern: the functional composition of a regional species pool. A regional species pool is the set of species present in all communities. While the functional composition of a community depends on the set of interactors at that locality, the functional composition of a regional species pool depends on the possibility of those interactors being present in at least one constituent community.

In Chapter 2 I tested two long standing hypotheses of how species durations are structured: the survival of the unspecialized hypothesis [298], and the Law of Constant Extinction [335]. I analyzed how the distribution of mammal species durations is affected by differences in multiple species traits, species' phylogenetic relatedness, and species' origination cohort. My results supported the conclusion that generalist mammal species will, on average, have a greater duration than more specialized mammal species. I also found that phylogeny and origination cohort contribute sub-equally to variation in species duration. Additionally, I found that only some of factors associated with extinction risk for Modern mammals could

be considered risk factors for mammals from the rest of the Cenozoic. For example, while there is an increased risk of extinction with increasing body size amongst Modern fauna, this was not the normal case for Cenozoic. Finally, I found evidence of species extinction risk increasing with species duration, a result that is counter the Law of Constant Extinction.

Chapter 3 also deals with the survival of the unspecialized hypothesis as well as the Law of Constant Extinction, this time with the global record of post-Cambrian Paleozoic brachiopods. In addition to these hypotheses, I also analyzed the relationship between extinction intensity and the strength of trait selection, namely “do the selective differences between traits increase or decrease with average fitness increases or decreases?” I found a negative correlation between intensity of extinction and the effects of geographic range and environmental preference. As with Chapter 2, I also found support for greater survival among environmental generalists than specialists. These results supported the conclusion that, at least for Paleozoic brachiopods, as extinction intensity increases, the selective difference of traits increases. In this analysis this means that when average duration decreases (e.g. intensity is high) the effect of genus geographic range increases in magnitude and taxa which favor epicontinental environments are expected to have a greater duration than those which favor open-ocean environments. I also find that the change in magnitude of effect is expected to be greater for environmental preference than for geographic range as the overall effect of former have a much greater variance than that of the later.

The final study of this dissertation (Chapter 4) was an analysis of how the functional composition of the North American mammal regional species pool changed over time and in response to multiple environmental factors. The goals of this analysis were to understand when are different ecotypes enriched or depleted in the regional species pool, and to understand how changes to environmental context may affect changes to the functional diversity of the regional species pool. By focusing on functional groups instead of taxonomic groups, the

results from this study are phrased in terms of species interactions and not differences in clade diversity. My results add considerable nuance to the taxon-focused narrative of North American environmental change. There are many results and conclusions from this analysis, so I focus here on a few key results. I found that mammal diversity is more strongly shaped by changes to origination rate among a few ecotypes rather than being driven by selective extinction one or more ecotypes. I also found that all arboreal ecotypes decline throughout the Paleogene and disappear from the species pool by the Neogene. Additionally, I found that most herbivore ecotypes expand their relative contribution to functional diversity over time. Finally, I found that the environmental factors analyzed here structure differences in ecotype origination probability but not survival probability.

5.2 Synthesis

These three chapters are united by their analysis of species functional traits. Analysis of species traits is the easiest way to unite macroevolutionary and macroecological inference by using this commonality to develop and test integrated hypotheses [196, 358]. All inferences lead to more hypotheses as we realize what we do and do not know. Inferences from analysis of macroevolutionary patterns may lead to hypotheses about macroecological patterns, such as the relationship between extinction risk and diversity in arboreal mammals (Chapter 2, and 4).

The first two studies, when considered together, add a considerable degree of nuance to our understanding of multiple macroevolutionary hypotheses such as the Law of Constant Extinction and the survival of the unspecialized.

First and foremost, the results of neither study support the Law of Constant Extinction [335]. I found evidence for extinction risk increasing with taxon duration; these results are

consistent with those of a “nearly-neutral” theoretical model of macroevolution [267]. While the dynamics of this model are described as “Red Queen” [267], this is not strictly true as Red Queen dynamics as described by Van Valen [335] require that extinction does not increase with taxon age. Instead, the dynamics of the nearly-neutral model are “Red Queen” in the sense that while all species increase in expected fitness, their relative fitnesses do not change. Interestingly, the decrease in absolute extinction risk towards the Modern [258, 82], which translates to an increase in expected species duration, may reflect similar dynamics to the nearly-neutral model. One of the innovations behind this result may be allowing species survival to decay non-linearly with duration by modeling durations as observations from a Weibull distribution as opposed to an exponential distribution (Chapter 2, and 3), a modeling decision that is becoming more commonplace [55, 68].

These two studies also provide broad support for the hypothesis of the survival of the unspecialized [298], at least with respect to the covariates included in either model. The survival of the unspecialized appears to be a nearly universal pattern in macroevolution [298, 169, 170, 217, 218, 21, 250]. I did find, however, that for post-Cambrian Paleozoic brachiopods when extinction intensity increases, the relationship with environmental preference and duration changes from a pattern where intermediate environmental preference are are expected to have the greatest durations to a pattern where more specialized, epicontinental-favoring taxa are expected to have the greatest durations. This result adds a degree of nuance to the survival of the unspecialized, specifically with regards to when it is expected to “hold.” My conclusion is that during periods of low intensity extinction risk or “background extinction” [134, 87], the survival of the unspecialized hypothesis will hold. However, as extinction intensity increases, this hypothesis may not accurately describe differences in extinction risk across taxa. As such, the survival of the unspecialized may serve as an excellent default for studies of trait selection and taxon extinction. This conclusion can be considered evidence for the qualitative difference between background and mass extinction [134].

In the third study I analyzed how the functional composition of a regional species pool changes over millions of years (Chapter 4). While this pattern is macroecological, many of the hypotheses and questions encompassed by this study were generated from macroevolutionary analyses. For example, the result from the first study that arboreal taxa have a greater extinction risk than other mammal locomotor groups did not have an unambiguous explanation for if and how extinction risk could have changed over the Cenozoic: always present but high risk, or higher extinction risk in the Neogene compared to the Paleogene (Chapter 2). The third study, in its analysis of the relative diversity of mammal ecotypes, was able to more fully resolve the previous results as I found that diversity of arboreal ecotypes declined through the Paleogene, becoming extremely rare or entirely absent from the species pool by the Neogene (Chapter 4). This result is much more nuanced than the either of two proposed processes (Chapter 2) and speaks to the improved insights and inferences a unified macroevolutionary and macroecological research program can provide.

The third study also, methodologically, represents the strongest unification of macroevolutionary and macroecological approaches to inference (Chapter 4). The question at the heart of this study is “when are mammal ecotypes enriched or depleted relative to their average diversity” is both macroevolutionary and macroecological, any statistical inferences needed to represent this. The model at the center of this study was a combination of a birth-death process with a fourth-corner model from ecology. The birth-death process provided a mechanism for changes to species diversity over time while the fourth-corner model recast diversification in terms of the relative contribution of functional groups to the regional species pool. The results of this study could then be phrased in how well adapted the functional groups are to their changing environmental context.

5.3 Future

There are many possible ways to expand on the analyses presented in this dissertation. There are also many unanswered questions which have been raised by each of these analyses which require future study.

A major limitation the fossil record of North American mammals is poor spatial resolution for the entire Cenozoic, something that restricts macroecological analyses (Chapter 4). While a regional species pool represents the set of all species present in a region, the individual dynamics communities give a much more complete idea of why species pool diversity changes. For example, I've been unable to estimate how changes to functional diversity of local communities scales up to the functional diversity of the regional species pool these communities are drawn from [117]. Given a fossil record with a high resolution spatial record, the types of analyses presented in Chapter 4 could be expanded to incorporate the spatial dynamics of functional diversity. The Bayesian modeling framework used throughout this dissertation makes this imminently possible given the right dataset [16]. The results of such an analysis would shed a lot of light on how the functional diversity of communities can vary spatially and how those communities respond differently to environmental change. A restricted analysis, both temporally and spatially, that focuses on a high resolution fossil record would be an appropriate place to start for this analysis; for example, the late Neogene of North America is extensively sampled as evidenced by the MioMap, Neotoma, and FaunMap projects (<http://www.ucmp.berkeley.edu/neomap/use.html>) and may be an ideal starting point for this type of analysis.

Preservation of the fossil record is a pervasive issue in paleontology [89, 78, 76, 81, 177, 352]. The first two studies gloss over issues of preservation by either ignoring it (Chapter 2) or including a sampling proxy as a covariate (Chapter 3); these decisions were partially due to

limitations of the models underlying both those studies. In contrast, in Chapter 4 I directly modeled the preservation process and allowed preservation probability to vary with time and allowed species body size to potentially affect differences in preservation probability across taxa. However, I ignored the possible differences in preservation over time due to species functional group or changes to environmental context. I chose to limit the possible covariates which could affect of preservation because this type of analysis was beyond the scope of Chapter 4.

All of three of these studies could be improved by incorporating more information about fossil preservation such as how functional groups and environmental context can shape differences in preservation probability. For example, when studying marine invertebrates covariates such as sea-level or shell composition (e.g. aragonitic vs calcitic) are all potentially very important for better understanding the differential preservation of species [86, 227, 228, 112]. Given a more representative model of preservation, even more complete and accurate macroevolutionary and macroecological inferences can be made.

5.4 Final thoughts

The three studies presented in this dissertation are all representative of my rhetoric championing integrated macroevolutionary and macroecological study (Chapter 1). Each study emphasizes the importance of framing hypotheses in terms of ecological interactions in order to make strong inferences. By using an extremely flexible and expressive modeling strategy, I have been able to translate precise scientific questions into statistical models. The scope of inferences that can be made for each study are clear and conditioned on the explicit modeling assumptions made for each analysis. The first step to building a dialogue is agreeing on a common language and my strategy has been to emphasize a common statistical framework

with which to phrase our analyses. My hope is that this approach to using paleontological data for answering questions about the processes underlying emergent patterns in evolutionary and ecological data fosters a stronger relationship between the disciplines of macroevolution and macroecology as well as paleontologists and neontologists.

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APPENDIX A

TABLE OF SPECIES MASS ESTIMATES

Table A.1: Body mass estimates in grams for all mammal species included in this study. Also included is the source of this measurement. PBDB = Paleobiology Database, NOW = Neogene Old World Database, EOL = Encyclopedia of Life, ADW = Animal Diversity Web. PBDB + regression indicates that body mass was estimated from a measurement on the PBDB using one of the many regression equations listed in Table 2.3.

Species	Mass (g)	Source
<i>Aaptoryctes ivyi</i>	215.70	PBDB + regression
<i>Absarokius abbotti</i>	176.39	PBDB + regression
<i>Absarokius metoecus</i>	166.29	PBDB + regression
<i>Acarictis ryani</i>	291.90	PBDB + regression
<i>Achaenodon robustus</i>	492900.83	PBDB + regression
<i>Achlyoscapter longirostris</i>	11.94	[329]
<i>Acmeodon secans</i>	325.36	PBDB + regression
<i>Acritohippus isonesus</i>	135944.23	[329]
<i>Acritohippus quinni</i>	178082.11	[329]
<i>Acritoparamys atwateri</i>	190.07	PBDB + regression
<i>Acritoparamys francesi</i>	125.17	PBDB + regression
<i>Acritoparamys pattersoni</i>	432.28	PBDB
<i>Acritoparamys wyomingensis</i>	274.75	PBDB
<i>Adeloblarina berklandi</i>	12.68	[329]
<i>Adelphailurus kansensis</i>	33189.87	[329]
<i>Adilophontes brachykolos</i>	119372.01	[329]
<i>Adjidaumo burkei</i>	4.91	PBDB
<i>Adjidaumo craigi</i>	7.29	PBDB + regression
<i>Adjidaumo intermedius</i>	10.39	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Adjidaumo maximus</i>	20.14	PBDB + regression
<i>Adjidaumo minimus</i>	7.29	PBDB + regression
<i>Adjidaumo minutus</i>	15.90	PBDB + regression
<i>Adunator ladae</i>	17.36	PBDB + regression
<i>Aelurodon asthenostylus</i>	22026.47	[329]
<i>Aelurodon ferox</i>	26370.47	[329]
<i>Aelurodon mcgrewi</i>	22247.84	[329]
<i>Aelurodon montanensis</i>	27722.51	[329]
<i>Aelurodon stirtoni</i>	20537.34	[329]
<i>Aelurodon taxoides</i>	29436.77	[329]
<i>Aepinacodon americanus</i>	423622.49	[20]
<i>Aepycamelus bradyi</i>	516464.33	[59]
<i>Aepycamelus giraffinus</i>	499050.27	PBDB
<i>Aepycamelus robustus</i>	420836.64	[329]
<i>Aepycamelus stocki</i>	348014.70	[329]
<i>Aethomylos simplicidens</i>	30.03	PBDB + regression
<i>Ageitodendron matthewi</i>	331.60	PBDB + regression
<i>Agriochoerus antiquus</i>	56387.34	[329]
<i>Agriochoerus guyotianus</i>	59000.00	[198]
<i>Agriotherium schneideri</i>	355045.06	[329]
<i>Alagomys russelli</i>	4.67	PBDB + regression
<i>Aletodon conardae</i>	576.81	PBDB + regression
<i>Aletodon gunnelli</i>	615.51	PBDB + regression
<i>Aletodon quadravus</i>	365.67	PBDB + regression
<i>Aletomeryx gracilis</i>	27173.57	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Alforjas taylori</i>	408399.03	[329]
<i>Alilepus vagus</i>	225.12	PBDB + regression
<i>Alilepus wilsoni</i>	111.45	PBDB + regression
<i>Allomys cristabrevis</i>	239.72	PBDB + regression
<i>Allomys simplicidens</i>	122.00	[198]
<i>Allomys storeri</i>	92.85	PBDB + regression
<i>Alluviorex arcadentes</i>	5.70	[329]
<i>Alphagaulus pristinus</i>	521.00	[198]
<i>Alphagaulus vetus</i>	523.22	[329]
<i>Alticonus gazini</i>	1140.52	PBDB + regression
<i>Alveojunctus minutus</i>	25.80	PBDB + regression
<i>Alveugena carbonensis</i>	815.39	PBDB
<i>Alwoodia harkseni</i>	294.29	PBDB + regression
<i>Alwoodia magna</i>	226.00	[198]
<i>Amebelodon floridanus</i>	37020787.90	PBDB + regression
<i>Amelotabes simpsoni</i>	57.16	PBDB + regression
<i>Ammospermophilus hanfordi</i>	58.00	[198]
<i>Ammospermophilus junturensis</i>	53.52	[329]
<i>Amphechinus horncloudi</i>	175.91	[329]
<i>Amphicaenopus platycephalus</i>	2397650.84	[329]
<i>Amphicyon frendens</i>	245241.81	[329]
<i>Amphicyon galushai</i>	138690.48	[329]
<i>Amphicyon ingens</i>	600000.00	[314]
<i>Amphicyon longiramus</i>	113550.16	[329]
<i>Amphicyon riggsi</i>	418398.16	PBDB

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Amphimachairodus coloradensis</i>	157147.46	PBDB + regression
<i>Ampliconus antoni</i>	2906.21	PBDB + regression
<i>Amynodon advenus</i>	1987010.17	PBDB + regression
<i>Amynodon reedi</i>	579004.85	PBDB
<i>Amynodontopsis bodei</i>	608129.44	PBDB + regression
<i>Anaptomorphus aemulus</i>	95.39	PBDB + regression
<i>Anaptomorphus westi</i>	186.75	PBDB + regression
<i>Anasazia williamsoni</i>	186.23	[361]
<i>Anchitherium clarencei</i>	230960.04	[329]
<i>Anconodon cochranensis</i>	57.00	[362]
<i>Anemorhysis natronensis</i>	32.59	PBDB + regression
<i>Anemorhysis pattersoni</i>	57.16	PBDB + regression
<i>Anemorhysis pearcei</i>	32.00	[2]
<i>Anemorhysis wortmani</i>	42.74	PBDB + regression
<i>Angustidens vireti</i>	18.17	[329]
<i>Anisonchus oligistus</i>	388.07	PBDB + regression
<i>Anisonchus onostus</i>	616.78	PBDB + regression
<i>Anisonchus sectorius</i>	1619.16	PBDB + regression
<i>Ankalagon saurognathus</i>	28250.27	PBDB + regression
<i>Ankyloodon annectens</i>	30.27	[329]
<i>Anomoemys lewisi</i>	207.64	[293]
<i>Ansomys hepburnensis</i>	44.70	[329]
<i>Ansomys nevadensis</i>	51.94	[329]
<i>Ansomys nexodens</i>	92.85	PBDB + regression
<i>Antecalomys phthanus</i>	22.87	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Antecalomys valensis</i>	13.60	[329]
<i>Antecalomys vasquezi</i>	10.70	PBDB
<i>Antiacodon pygmaeus</i>	464.26	PBDB + regression
<i>Antiacodon venustus</i>	2156.43	PBDB + regression
<i>Apataelurus kayi</i>	10833.16	PBDB + regression
<i>Apatemys bellulus</i>	17.36	PBDB + regression
<i>Apatemys bellus</i>	21.80	PBDB + regression
<i>Apatemys chardini</i>	14.63	PBDB + regression
<i>Apatemys downsi</i>	65.02	PBDB + regression
<i>Apatemys hendryi</i>	12.08	PBDB + regression
<i>Apatemys uintensis</i>	28.30	PBDB + regression
<i>Apatosciuravus bifax</i>	35.35	PBDB + regression
<i>Apatosciuravus jacobsi</i>	29.87	PBDB + regression
<i>Apheliscus chydaeus</i>	47.27	PBDB + regression
<i>Apheliscus insidiosus</i>	60.38	PBDB + regression
<i>Apheliscus nitidus</i>	77.18	PBDB + regression
<i>Apheliscus wapitiensis</i>	26.62	PBDB + regression
<i>Aphelops malacorhinus</i>	3541284.24	[329]
<i>Aphelops megalodus</i>	1689595.99	[329]
<i>Aphelops mutilus</i>	4325334.34	[329]
<i>Aphronorus fraudator</i>	112.44	PBDB + regression
<i>Aphronorus orieli</i>	427.05	PBDB + regression
<i>Aphronorus ratatoski</i>	176.14	PBDB + regression
<i>Apletotomeus crassus</i>	8.78	PBDB + regression
<i>Apternodus gregoryi</i>	90.14	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Apternodus iliffensis</i>	87.48	PBDB + regression
<i>Arapahovius advena</i>	42.74	PBDB + regression
<i>Archaeocyon falkenbachi</i>	2116.88	[322]
<i>Archaeocyon leptodus</i>	3533.34	[329]
<i>Archaeocyon pavidus</i>	2275.60	[329]
<i>Archaeohippus blackbergi</i>	33189.87	[329]
<i>Archaeohippus mannulus</i>	48917.80	PBDB + regression
<i>Archaeohippus mourningi</i>	54176.36	[329]
<i>Archaeohippus penultimus</i>	71682.36	[329]
<i>Archaeohippus stenolophus</i>	63678.98	PBDB
<i>Archaeolagus acaricolus</i>	578.25	[329]
<i>Archaeolagus emeraldensis</i>	2344.90	[329]
<i>Archaeolagus ennisianus</i>	1064.22	[329]
<i>Archaeolagus macrocephalus</i>	1826.21	[329]
<i>Archaeolagus primigenius</i>	1540.71	[329]
<i>Archaeotherium lemleyi</i>	1361176.27	PBDB + regression
<i>Archaeotherium mortoni</i>	240808.54	PBDB + regression
<i>Archaeotherium trippensis</i>	3353590.86	PBDB + regression
<i>Arctocyon montanensis</i>	14771.98	PBDB + regression
<i>Arctocyon mumak</i>	57080.05	PBDB
<i>Arctodontomys nuptus</i>	318.11	PBDB + regression
<i>Arctodontomys simplicidens</i>	219.44	PBDB + regression
<i>Arctodontomys wilsoni</i>	236.68	PBDB + regression
<i>Arctodus pristinus</i>	299916.25	[305]
<i>Arctonasua eurybates</i>	15994.50	[329]

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Arctonasua gracilis</i>	8866.19	[329]
<i>Arctonasua minima</i>	7044.48	[329]
<i>Arctostylops steini</i>	512.92	PBDB + regression
<i>Ardynomys occidentalis</i>	202.01	PBDB + regression
<i>Arfia junnei</i>	1188.60	PBDB + regression
<i>Arfia opisthotoma</i>	11400.50	PBDB + regression
<i>Arfia shoshoniensis</i>	9550.20	PBDB + regression
<i>Arfia zele</i>	2236.53	PBDB + regression
<i>Arretotherium acridens</i>	179871.86	[329]
<i>Arretotherium fricki</i>	138690.48	[329]
<i>Arretotherium leptodus</i>	252848.32	PBDB
<i>Artimonius australis</i>	133.09	PBDB + regression
<i>Artimonius nocerae</i>	66.71	PBDB + regression
<i>Artimonius witteri</i>	91.55	PBDB + regression
<i>Astrohippus stockii</i>	134877.06	PBDB
<i>Aulolithomys bounites</i>	68.53	PBDB + regression
<i>Aulolithomys vexilliaris</i>	17.97	PBDB + regression
<i>Australocamelus orarius</i>	100709.96	[329]
<i>Auxodonton pattersoni</i>	6261.76	PBDB + regression
<i>Avunculus didelphodonti</i>	60.38	PBDB + regression
<i>Aycrossia lovei</i>	124.23	PBDB + regression
<i>Aztlanolagus agilis</i>	1999.86	[305]
<i>Azygonyx ancylion</i>	15969.85	PBDB + regression
<i>Azygonyx grangeri</i>	24602.41	PBDB + regression
<i>Azygonyx xenicus</i>	7316.20	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Baiocodon denverensis</i>	4195.64	PBDB + regression
<i>Baiocodon nordicus</i>	1675.41	PBDB + regression
<i>Baiomys rexroadi</i>	7.29	PBDB + regression
<i>Baiotomeus douglassi</i>	192.03	PBDB + regression
<i>Baiotomeus rhothonion</i>	12.82	PBDB + regression
<i>Barbourofelis fricki</i>	255250.32	[329]
<i>Barbourofelis loveorum</i>	160091.76	PBDB + regression
<i>Barbourofelis morrisi</i>	90219.42	[329]
<i>Barbourofelis osborni</i>	162309.29	[192]
<i>Barbourofelis whitfordi</i>	77652.58	[329]
<i>Barbouromeryx trigonocorneus</i>	33860.35	[329]
<i>Barylambda faberi</i>	255787.00	PBDB + regression
<i>Barylambda jackwilsoni</i>	43914.63	PBDB + regression
<i>Basirepomys pliocenicus</i>	72.36	PBDB + regression
<i>Basirepomys robertsi</i>	54.13	PBDB + regression
<i>Bassariscus antiquus</i>	1881.83	[329]
<i>Bassariscus casei</i>	1652.43	[329]
<i>Bassariscus minimus</i>	1183.06	[260]
<i>Bassariscus ogallalae</i>	1724.26	[103]
<i>Bassariscus parvus</i>	1685.81	[329]
<i>Bathygenys alpha</i>	6155.81	PBDB + regression
<i>Bathygenys reevesi</i>	5297.18	PBDB + regression
<i>Batodonoides powayensis</i>	3.14	PBDB + regression
<i>Batodonoides vanhouteni</i>	1.26	PBDB + regression
<i>Beckiasorex hibbardi</i>	6.56	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Bensonomys arizonae</i>	23.19	PBDB
<i>Bensonomys baskini</i>	26.95	PBDB
<i>Bensonomys elachys</i>	8.78	[159]
<i>Bensonomys gidleyi</i>	13.90	PBDB
<i>Bensonomys lindsayi</i>	8.78	PBDB + regression
<i>Bensonomys meadensis</i>	19.00	PBDB
<i>Bensonomys winklerorum</i>	9.09	PBDB
<i>Bensonomys yazhi</i>	9.98	[266]
<i>Bettonnia tsosia</i>	50.22	[45]
<i>Bisonalveus browni</i>	90.14	PBDB + regression
<i>Bisonalveus holtzmani</i>	146.12	PBDB + regression
<i>Blacktops latidens</i>	344.57	PBDB
<i>Blacktops longinaires</i>	255.48	PBDB
<i>Blarina brevicauda</i>	16.40	PBDB
<i>Blarina carolinensis</i>	13.49	[305]
<i>Blastomeryx gemmifer</i>	10938.02	[329]
<i>Blastomeryx pristinus</i>	38955.58	PBDB
<i>Blickomylus galushai</i>	16983.54	[329]
<i>Boreameryx braskerudi</i>	9643.88	PBDB + regression
<i>Borophagus diversidens</i>	34891.55	[329]
<i>Borophagus dudleyi</i>	94433.22	[57]
<i>Borophagus hilli</i>	29143.87	[329]
<i>Borophagus littoralis</i>	23388.51	[329]
<i>Borophagus orc</i>	16814.55	[329]
<i>Borophagus parvus</i>	19341.34	[329]

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Borophagus pugnator</i>	24100.79	[329]
<i>Borophagus secundus</i>	24100.79	[329]
<i>Bothriodon rostratus</i>	865704.50	[40]
<i>Bouromeryx americanus</i>	68186.37	[329]
<i>Bouromeryx submilleri</i>	50011.09	[329]
<i>Brachycrus buwaldi</i>	250196.03	[329]
<i>Brachycrus laticeps</i>	359952.66	PBDB
<i>Brachycrus rusticus</i>	117343.54	PBDB
<i>Brachycrus siouense</i>	145801.30	[329]
<i>Brachyerix hibbardi</i>	78.19	[44]
<i>Brachyerix incertis</i>	79.84	[329]
<i>Brachyerix macrotis</i>	131.63	[329]
<i>Brachyerix richi</i>	340.36	[329]
<i>Brachyhyops viensis</i>	202519.16	PBDB + regression
<i>Brachyhyops wyomingensis</i>	97111.99	PBDB + regression
<i>Brachypsalis modicus</i>	512.86	[329]
<i>Brachypsalis obliquidens</i>	63169.09	PBDB
<i>Brachypsalis pachycephalus</i>	487.85	[329]
<i>Brachyrhynchocyon dodgei</i>	13906.88	PBDB + regression
<i>Brachyrhynchocyon montanus</i>	7397.01	PBDB + regression
<i>Brontops tyleri</i>	571500.00	PBDB
<i>Buisnictis brevirostris</i>	849.04	PBDB + regression
<i>Buisnictis burrowsi</i>	817.07	PBDB
<i>Buisnictis schoffi</i>	22.42	[329]
<i>Bunomeryx montanus</i>	3196.73	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Bunophorus etsagicus</i>	13323.34	PBDB + regression
<i>Bunophorus grangeri</i>	13969.78	PBDB + regression
<i>Bunophorus macropternus</i>	10378.10	PBDB + regression
<i>Bunophorus pattersoni</i>	6332.74	[187]
<i>Bunophorus robustus</i>	5163.91	PBDB + regression
<i>Bunophorus sinclairi</i>	16312.82	PBDB + regression
<i>Caenolambda jepseni</i>	40906.63	PBDB + regression
<i>Calippus cerasinus</i>	81633.91	[329]
<i>Calippus elachistus</i>	43044.94	[329]
<i>Calippus hondurensis</i>	71682.36	[329]
<i>Calippus martini</i>	119372.01	[329]
<i>Calippus placidus</i>	79221.26	[329]
<i>Calippus proplacidus</i>	64860.88	[329]
<i>Calippus regulus</i>	45251.90	[329]
<i>Camelops hesternus</i>	1099005.84	[305]
<i>Camelops traviswhitei</i>	1027000.94	PBDB + regression
<i>Campestrallomys annectens</i>	80.29	PBDB + regression
<i>Campestrallomys dawsonae</i>	298.87	[329]
<i>Campestrallomys siouxensis</i>	159.17	[329]
<i>Canis armbrusteri</i>	30333.26	[329]
<i>Canis edwardii</i>	79873.95	PBDB + regression
<i>Canis latrans</i>	11765.00	PBDB
<i>Canis lepophagus</i>	14617.87	[329]
<i>Canis rufus</i>	15566.00	PBDB
<i>Cantius abditus</i>	2798.04	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Cantius angulatus</i>	417.80	PBDB + regression
<i>Cantius frugivorus</i>	913.71	PBDB + regression
<i>Cantius mckennai</i>	902.97	PBDB + regression
<i>Cantius nunienus</i>	1212.97	PBDB + regression
<i>Cantius ralstoni</i>	738.98	PBDB + regression
<i>Cantius simonsi</i>	3858.47	PBDB + regression
<i>Cantius torresi</i>	410.27	PBDB + regression
<i>Cantius trigonodus</i>	2000.00	[312]
<i>Capricamelus gettyi</i>	361557.35	PBDB + regression
<i>Capromeryx tauntonensis</i>	15835.35	[329]
<i>Cardiolophus radinskyi</i>	32305.76	PBDB + regression
<i>Cardiolophus semihians</i>	33364.14	PBDB + regression
<i>Carpocristes cygneus</i>	25.80	PBDB + regression
<i>Carpocristes hobackensis</i>	33.00	[312]
<i>Carpocyon compressus</i>	15214.44	[329]
<i>Carpocyon robustus</i>	19341.34	[329]
<i>Carpocyon webbi</i>	20537.34	[329]
<i>Carpodaptes hazelae</i>	51.17	PBDB + regression
<i>Carpodaptes stonleyi</i>	32.59	PBDB + regression
<i>Carpolestes nigridens</i>	87.00	[277]
<i>Carpolestes simpsoni</i>	27.99	PBDB + regression
<i>Carpomegodon jepseni</i>	166.29	PBDB + regression
<i>Catopsalis alexanderi</i>	3415.85	PBDB + regression
<i>Catopsalis calgariensis</i>	22815.78	PBDB + regression
<i>Catopsalis foliatus</i>	7371.53	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Catopsalis joyneri</i>	2435.00	[362]
<i>Cedromus wardi</i>	246.28	PBDB + regression
<i>Centetodon aztecus</i>	17.36	PBDB + regression
<i>Centetodon bembicophagus</i>	10.87	PBDB + regression
<i>Centetodon chadronensis</i>	23.37	PBDB + regression
<i>Centetodon divaricatus</i>	30.57	[329]
<i>Centetodon hendryi</i>	17.36	PBDB + regression
<i>Centetodon kuenzii</i>	26.62	PBDB + regression
<i>Centetodon magnus</i>	33.45	[329]
<i>Centetodon neashami</i>	53.67	PBDB + regression
<i>Centetodon pulcher</i>	30.03	PBDB + regression
<i>Centimanomys major</i>	239.72	PBDB + regression
<i>Ceratogaulus hatcheri</i>	1490.63	[40]
<i>Cernictis hesperus</i>	177.68	[329]
<i>Cernictis repenningi</i>	8800.69	[111]
<i>Chacomylus sladei</i>	151.65	PBDB + regression
<i>Chadrolagus emryi</i>	63.45	PBDB + regression
<i>Chalicomomys willwoodensis</i>	4.06	PBDB + regression
<i>Chasmaporthes ossifragus</i>	107821.85	PBDB + regression
<i>Chipetaia lamporea</i>	417.80	PBDB + regression
<i>Chiromyoides caesor</i>	219.44	PBDB + regression
<i>Chiromyoides minor</i>	70.04	PBDB + regression
<i>Chiromyoides potior</i>	202.80	PBDB + regression
<i>Chriacus badgleyi</i>	1200.76	PBDB + regression
<i>Chriacus baldwini</i>	1550.07	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Chriacus gallinae</i>	1468.97	PBDB + regression
<i>Chriacus pelvidens</i>	5047.26	PBDB + regression
<i>Chumashius balchi</i>	111.45	PBDB + regression
<i>Churcheria baroni</i>	50.76	PBDB + regression
<i>Cimexomys minor</i>	27.99	PBDB + regression
<i>Cimolestes incisus</i>	204.60	PBDB + regression
<i>Colodon cingulatus</i>	222816.55	PBDB
<i>Colodon kayi</i>	49060.48	PBDB + regression
<i>Colodon occidentalis</i>	110573.45	PBDB + regression
<i>Colodon stovalli</i>	57498.66	PBDB + regression
<i>Colodon woodi</i>	85748.60	PBDB + regression
<i>Conacodon cophater</i>	236.68	PBDB + regression
<i>Conacodon delphae</i>	3318.67	PBDB + regression
<i>Conacodon entoconus</i>	2268.93	PBDB + regression
<i>Conacodon kohlbergeri</i>	590.00	PBDB + regression
<i>Conoryctes comma</i>	16724.87	PBDB + regression
<i>Copecion brachypternus</i>	4501.11	PBDB + regression
<i>Copecion davisi</i>	2798.04	PBDB + regression
<i>Copedelphys innominata</i>	23.70	PBDB + regression
<i>Copedelphys titanelix</i>	6.11	PBDB + regression
<i>Copelemur australotutus</i>	1633.14	PBDB + regression
<i>Copelemur tutus</i>	1864.20	PBDB + regression
<i>Copemys barstowensis</i>	32.14	[329]
<i>Copemys esmeraldensis</i>	27.94	[329]
<i>Copemys lindsayi</i>	14.88	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Copemys longidens</i>	26.84	[329]
<i>Copemys loxodon</i>	28.79	[329]
<i>Copemys mariae</i>	31.50	[329]
<i>Copemys pagei</i>	15.18	[329]
<i>Copemys russelli</i>	24.29	[329]
<i>Copemys shotwelli</i>	10.39	PBDB + regression
<i>Copemys tenuis</i>	23.34	[329]
<i>Coriphagus encinensis</i>	254.29	PBDB + regression
<i>Coriphagus montanus</i>	130.39	PBDB + regression
<i>Cormocyon copei</i>	4817.45	[329]
<i>Cormocyon haydeni</i>	4359.01	[329]
<i>Coryphodon armatus</i>	351055.71	PBDB + regression
<i>Coryphodon eocaenus</i>	218518.57	PBDB + regression
<i>Coryphodon lobatus</i>	570461.25	PBDB + regression
<i>Coryphodon proterus</i>	479331.19	PBDB + regression
<i>Coryphodon radians</i>	339220.51	PBDB + regression
<i>Cosoryx cerroensis</i>	16814.55	[329]
<i>Cosoryx furcatus</i>	13678.75	PBDB
<i>Cranioceras clarendonensis</i>	89936.76	PBDB + regression
<i>Cranioceras teres</i>	96761.07	[329]
<i>Cranioceras unicornis</i>	134591.56	[329]
<i>Cratogeomys sansimoniensis</i>	140.29	PBDB + regression
<i>Crucimys milleri</i>	36.60	[329]
<i>Crypholestes vaughni</i>	18.68	PBDB
<i>Cryptotis adamsi</i>	13.33	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Cryptotis kansasensis</i>	33.60	PBDB + regression
<i>Cryptotis parva</i>	4.10	PBDB
<i>Cupidinimus avawatzensis</i>	20.09	[329]
<i>Cupidinimus bidahochiensis</i>	27.39	[329]
<i>Cupidinimus boronensis</i>	14.59	[329]
<i>Cupidinimus eurekensis</i>	2.98	PBDB
<i>Cupidinimus halli</i>	10.39	PBDB + regression
<i>Cupidinimus lindsayi</i>	7.61	[329]
<i>Cupidinimus madisonensis</i>	10.56	PBDB
<i>Cupidinimus magnus</i>	22.41	PBDB + regression
<i>Cupidinimus nebraskensis</i>	9.30	[329]
<i>Cupidinimus prattensis</i>	21.33	[329]
<i>Cupidinimus tertius</i>	16.44	[329]
<i>Cupidinimus whitlocki</i>	16.78	[329]
<i>Cuvieronius tropicus</i>	56818220.68	PBDB + regression
<i>Cuyamalagus dawsoni</i>	639.06	[329]
<i>Cylindrodon fontis</i>	88.57	PBDB + regression
<i>Cylindrodon nebraskensis</i>	167.14	PBDB + regression
<i>Cynarctoides acridens</i>	2921.93	[329]
<i>Cynarctoides gawnae</i>	2643.87	[329]
<i>Cynarctoides harlowi</i>	1863.11	[329]
<i>Cynarctoides lemur</i>	2321.57	[329]
<i>Cynarctoides luskensis</i>	2392.27	[329]
<i>Cynarctoides roii</i>	1826.21	[329]
<i>Cynarctus crucidens</i>	4964.16	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Cynarctus galushai</i>	9228.02	[329]
<i>Cynarctus saxatilis</i>	10097.06	[329]
<i>Cynelos caroniavorus</i>	16647.24	[329]
<i>Cynelos idoneus</i>	105873.47	[329]
<i>Cynelos sinapius</i>	213202.99	[329]
<i>Cynodesmus martini</i>	14185.85	[329]
<i>Cynodesmus thoooides</i>	9228.02	[329]
<i>Cynorca occidentale</i>	20537.34	[329]
<i>Cynorca sociale</i>	34624.09	PBDB + regression
<i>Cyriacotherium psamminum</i>	3551.95	PBDB
<i>Dakotallomys lillegraveni</i>	196.53	PBDB
<i>Dakotallomys pelycomyoides</i>	244.01	[202]
<i>Daphoenodon falkenbachi</i>	137310.49	[329]
<i>Daphoenodon notionastes</i>	43477.55	[329]
<i>Daphoenodon superbus</i>	77652.58	[329]
<i>Daphoenus hartshornianus</i>	13329.17	PBDB + regression
<i>Daphoenus lambei</i>	7397.01	PBDB + regression
<i>Daphoenus ruber</i>	12767.60	PBDB + regression
<i>Daphoenus socialis</i>	13000.00	[198]
<i>Daphoenus vetus</i>	19535.72	[329]
<i>Dartonius jepseni</i>	10.52	[119]
<i>Dasypus bellus</i>	44977.99	[305]
<i>Delahomeryx browni</i>	38948.67	[329]
<i>Desmatippus avus</i>	179871.86	[329]
<i>Desmatippus texanus</i>	102452.28	PBDB

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Desmatochoerus hesperus</i>	239142.20	[197]
<i>Desmatochoerus megalodon</i>	335100.00	[198]
<i>Desmatoclaenus hermaeus</i>	11400.50	PBDB + regression
<i>Desmatolagus schizopetrus</i>	198.46	PBDB
<i>Desmocyon matthewi</i>	8103.08	[329]
<i>Desmocyon thomsoni</i>	6974.39	[329]
<i>Diacocherus meizon</i>	23.70	PBDB + regression
<i>Diacocherus minutus</i>	45.48	PBDB + regression
<i>Diacodexis gracilis</i>	788.63	PBDB + regression
<i>Diacodexis kelleyi</i>	1550.07	PBDB + regression
<i>Diacodexis metsiacus</i>	1176.50	PBDB + regression
<i>Diacodexis minutus</i>	555.14	PBDB + regression
<i>Diacodexis primus</i>	1389.84	PBDB + regression
<i>Diacodexis secans</i>	2124.77	PBDB + regression
<i>Diacodon alticuspis</i>	324.82	PBDB + regression
<i>Diceratherium annectens</i>	864580.76	[329]
<i>Diceratherium armatum</i>	3541284.24	[329]
<i>Diceratherium gregorii</i>	946518.90	PBDB + regression
<i>Diceratherium niobrarense</i>	2105366.25	[329]
<i>Diceratherium tridactylum</i>	965112.54	[329]
<i>Didelphodus absarokae</i>	361.10	PBDB + regression
<i>Didelphodus altidens</i>	246.37	PBDB + regression
<i>Didelphodus rheos</i>	139.74	PBDB + regression
<i>Didelphodus serus</i>	133.48	PBDB + regression
<i>Didymictis altidens</i>	30566.06	PBDB

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Didymictis leptomylus</i>	3213.72	PBDB + regression
<i>Didymictis protenus</i>	7205.96	PBDB + regression
<i>Didymictis proteus</i>	3605.48	PBDB + regression
<i>Didymictis vancleveae</i>	65421.88	[276]
<i>Dikkomys matthewi</i>	47.47	[329]
<i>Dillerlemur pagei</i>	133.09	PBDB + regression
<i>Dilophodon destitutus</i>	369866.51	[300]
<i>Dilophodon minusculus</i>	27074.75	PBDB + regression
<i>Dinaelurus crassus</i>	37320.00	[198]
<i>Dinofelis palaeoonca</i>	70032.50	PBDB + regression
<i>Dinohippus interpolatus</i>	257815.63	[329]
<i>Dinohippus leardi</i>	392385.48	[329]
<i>Dinohippus leidyanus</i>	229900.00	[188]
<i>Dinohippus mexicanus</i>	609259.77	[329]
<i>Dinohippus spectans</i>	536500.00	[198]
<i>Dinohyus hollandi</i>	2032979.00	PBDB + regression
<i>Diplobunops matthewi</i>	100751.34	PBDB + regression
<i>Dipodomys compactus</i>	49.20	[305]
<i>Dipodomys gidleyi</i>	17.97	PBDB + regression
<i>Dipodomys hibbardi</i>	17.81	[329]
<i>Diprionomys agrarius</i>	29.87	PBDB + regression
<i>Diprionomys parvus</i>	11.00	[198]
<i>Dipsalidictis aequidens</i>	42263.45	PBDB + regression
<i>Dipsalidictis platypus</i>	8936.27	PBDB + regression
<i>Dipsalidictis transiens</i>	22310.00	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Dipsalodon churchillorum</i>	26647.76	PBDB + regression
<i>Dipsalodon matthewi</i>	66505.41	PBDB + regression
<i>Dissacus navajovius</i>	6299.67	PBDB + regression
<i>Dissacus praenuntius</i>	14804.22	PBDB + regression
<i>Domnina dakotensis</i>	28.50	[329]
<i>Domnina gradata</i>	36.60	[329]
<i>Domnina greeni</i>	33.78	[329]
<i>Domnina thompsoni</i>	20.28	PBDB + regression
<i>Domninoides hessei</i>	149.90	[329]
<i>Domninoides knoxjonesi</i>	31.40	PBDB
<i>Domninoides mimicus</i>	135.64	[329]
<i>Domninoides riparensis</i>	56.83	[329]
<i>Dorraletes diminutivus</i>	37.32	PBDB + regression
<i>Douglassciurus jeffersoni</i>	360.63	[202]
<i>Downsimus chadwicki</i>	56.26	[329]
<i>Drepanomeryx falciformis</i>	90219.42	[329]
<i>Dromomeryx borealis</i>	144350.55	[329]
<i>Dryomomys dulcifer</i>	4.58	[215]
<i>Dryomomys szalayi</i>	6.07	[215]
<i>Duchesneodus uintensis</i>	2406372.12	PBDB + regression
<i>Dyseohyus fricki</i>	21807.30	[329]
<i>Dyseolemur pacificus</i>	70.04	PBDB + regression
<i>Ectocion cedrus</i>	4839.14	PBDB + regression
<i>Ectocion collinus</i>	5869.11	PBDB + regression
<i>Ectocion major</i>	12433.55	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Ectocion mediotuber</i>	5451.18	PBDB + regression
<i>Ectocion osbornianus</i>	6800.01	PBDB + regression
<i>Ectocion parvus</i>	3109.34	PBDB + regression
<i>Ectocion superstes</i>	13800.02	PBDB + regression
<i>Ectoconus ditrigonus</i>	22355.77	PBDB + regression
<i>Ectoganus gliriformes</i>	51205.04	[215]
<i>Ectoganus gliriformis</i>	51205.04	PBDB + regression
<i>Ectopocynus antiquus</i>	8266.78	[329]
<i>Ectopocynus intermedius</i>	12456.53	[329]
<i>Ectypodus aphronorus</i>	18.00	[362]
<i>Ectypodus childei</i>	14.42	PBDB + regression
<i>Ectypodus lovei</i>	14.42	PBDB + regression
<i>Ectypodus musculus</i>	30.25	PBDB + regression
<i>Ectypodus powelli</i>	19.74	PBDB + regression
<i>Ectypodus szalayi</i>	17.88	PBDB + regression
<i>Ectypodus tardus</i>	16.11	PBDB + regression
<i>Edworthia lerbekmoi</i>	23.70	PBDB + regression
<i>Ekgmowechashala philotau</i>	2079.74	[329]
<i>Elomeryx armatus</i>	157944.66	[329]
<i>Elphidotarsius florencae</i>	17.88	PBDB + regression
<i>Elphidotarsius russelli</i>	23.70	PBDB + regression
<i>Elphidotarsius wightoni</i>	27.99	PBDB + regression
<i>Elpidophorus elegans</i>	829.40	PBDB + regression
<i>Elpidophorus minor</i>	151.65	PBDB + regression
<i>Elwynella oreas</i>	99.30	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Elymys complexus</i>	5.92	PBDB + regression
<i>Enhydritherium terraenovae</i>	22180.76	PBDB + regression
<i>Enhydrocyon basilatus</i>	20332.99	[329]
<i>Enhydrocyon crassidens</i>	18582.95	[329]
<i>Enhydrocyon pahinsintewakpa</i>	14764.78	[329]
<i>Enhydrocyon stenocephalus</i>	14044.69	[329]
<i>Entoptychus grandiplanus</i>	59.74	[329]
<i>Entoptychus individens</i>	84.00	[198]
<i>Entoptychus planifrons</i>	134.29	[329]
<i>Entoptychus sheppardi</i>	94.63	[329]
<i>Entoptychus wheelerensis</i>	84.00	[198]
<i>Eoconodon gaudrianus</i>	3093.31	[371]
<i>Eoconodon hutchisoni</i>	9450.73	PBDB
<i>Eoconodon nidhoggii</i>	1665.06	PBDB + regression
<i>Eohaplomys matutinus</i>	1536.44	PBDB + regression
<i>Eohaplomys serus</i>	953.63	PBDB + regression
<i>Eohaplomys tradux</i>	772.04	PBDB + regression
<i>Eomoropus amarorum</i>	67651.40	PBDB + regression
<i>Eoryctes melanurus</i>	53.67	PBDB + regression
<i>Eotitanops borealis</i>	240072.90	PBDB + regression
<i>Eotitanops minimus</i>	43277.45	PBDB + regression
<i>Eotitanotherium osborni</i>	1484895.52	PBDB
<i>Eotylopus reedi</i>	38563.77	PBDB + regression
<i>Epeiromys spanius</i>	101.65	PBDB + regression
<i>Epicyon haydeni</i>	41772.77	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Epicyon saevus</i>	27722.51	[329]
<i>Epihippus gracilis</i>	18366.29	PBDB + regression
<i>Epihippus intermedius</i>	21835.02	PBDB
<i>Epitriplopus uintensis</i>	32600.00	[188]
<i>Eporeodon occidentalis</i>	118300.00	[198]
<i>Equus complicatus</i>	399944.75	[305]
<i>Equus conversidens</i>	306196.34	[305]
<i>Equus cumminsii</i>	314354.39	PBDB + regression
<i>Equus francisci</i>	217178.22	PBDB + regression
<i>Equus fromanius</i>	172311.00	[198]
<i>Equus giganteus</i>	399944.75	[305]
<i>Equus idahoensis</i>	629100.30	PBDB + regression
<i>Equus leidyi</i>	291150.23	PBDB + regression
<i>Equus occidentalis</i>	574116.46	[305]
<i>Equus scotti</i>	554625.71	[305]
<i>Equus simplicidens</i>	296558.57	[329]
<i>Eremotherium eomigrans</i>	2584400.00	[194]
<i>Eremotherium laurillardi</i>	799834.26	[305]
<i>Erythizon bathygnathum</i>	9163.63	PBDB + regression
<i>Erythizon kleini</i>	3854.02	PBDB + regression
<i>Escavadodon zygus</i>	137.63	PBDB + regression
<i>Esthonyx acutidens</i>	7159.09	PBDB + regression
<i>Esthonyx bisulcatus</i>	3066.57	PBDB + regression
<i>Esthonyx spatularius</i>	4505.96	PBDB + regression
<i>Eucyon davisii</i>	10509.13	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Eudaemonema cuspidata</i>	529.62	PBDB + regression
<i>Eumys brachyodus</i>	126.47	[329]
<i>Eumys elegans</i>	106.19	PBDB + regression
<i>Euoploctyon brachygnathus</i>	11271.13	[329]
<i>Euoploctyon spissidens</i>	9798.65	[329]
<i>Eusmilus cerebralis</i>	804.32	[329]
<i>Eusmilus sicarius</i>	34789.94	PBDB + regression
<i>Eutypomys acares</i>	84.39	PBDB + regression
<i>Eutypomys hibernodus</i>	729.27	PBDB + regression
<i>Eutypomys inexpectatus</i>	708.27	PBDB + regression
<i>Eutypomys montanensis</i>	943.88	[329]
<i>Eutypomys obliquidens</i>	202.01	PBDB + regression
<i>Eutypomys parvus</i>	273.26	PBDB + regression
<i>Eutypomys thomsoni</i>	578.37	PBDB + regression
<i>Fanimus clasoni</i>	87.14	PBDB
<i>Fanimus ultimus</i>	164.02	[329]
<i>Felis rexroadensis</i>	30333.26	[329]
<i>Ferinestrix vorax</i>	29333.85	PBDB
<i>Florentiamys agnewi</i>	84.77	[329]
<i>Florentiamys kinseyi</i>	157.59	[329]
<i>Florentiamys loomisi</i>	151.41	[329]
<i>Florentiamys tiptoni</i>	117.92	[329]
<i>Floridachoerus olsenii</i>	35596.41	[329]
<i>Floridameryx floridanus</i>	8184.52	[329]
<i>Floridatragulus dolichanthereus</i>	43477.55	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Floridatragulus texanus</i>	61083.68	[329]
<i>Fouchia elyensis</i>	10126.74	PBDB + regression
<i>Galbreathia bettae</i>	165.67	[329]
<i>Galbreathia novellus</i>	271.54	PBDB
<i>Galecyon mordax</i>	1984.90	PBDB + regression
<i>Gazinius amplus</i>	875.00	[312]
<i>Gazinocyon vulpeculus</i>	5070.63	PBDB + regression
<i>Gelastops joni</i>	183.10	PBDB + regression
<i>Gelastops parcus</i>	149.35	PBDB + regression
<i>Geomys carrranzai</i>	29.87	PBDB + regression
<i>Geringia gloveri</i>	26.84	[329]
<i>Geringia mcgregori</i>	41.26	[329]
<i>Gigantocamelus spatulus</i>	2615036.35	PBDB + regression
<i>Glossotherium chapadmalense</i>	310540.00	[194]
<i>Glyptotherium arizonae</i>	789680.00	[194]
<i>Gomphotherium obscurum</i>	50047676.55	PBDB
<i>Gomphotherium osborni</i>	24506452.87	[354]
<i>Goniodontomys disjunctus</i>	44.26	[329]
<i>Gracilicyon winkleri</i>	690.80	PBDB + regression
<i>Grangeria anarsius</i>	65434.40	PBDB + regression
<i>Gregorymys curtus</i>	106.70	[329]
<i>Gregorymys formosus</i>	77.48	[329]
<i>Gregorymys riograndensis</i>	43.82	[329]
<i>Gripholagomys lavocati</i>	507.76	[329]
<i>Griphomys alecer</i>	15.90	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Griphomys toltecus</i>	25.61	[202]
<i>Guanajuatomys hibbardi</i>	167.14	PBDB + regression
<i>Guildayomys hibbardi</i>	82.27	[329]
<i>Hapalodectes anthracinus</i>	352.26	PBDB + regression
<i>Hapalodectes leptognathus</i>	617.19	PBDB + regression
<i>Haplaletes disceptatrix</i>	84.86	PBDB + regression
<i>Haplaletes pelicatus</i>	130.39	PBDB + regression
<i>Hapoconus angustus</i>	1200.76	PBDB + regression
<i>Haplohippus texanus</i>	23265.89	PBDB + regression
<i>Haplolambda quinni</i>	53092.09	PBDB
<i>Haplolambda simpsoni</i>	138271.36	[328]
<i>Haplomylus bozemanensis</i>	87.48	PBDB + regression
<i>Haplomylus palustris</i>	72.22	PBDB + regression
<i>Haplomylus scottianus</i>	115.36	PBDB + regression
<i>Haplomylus simpsoni</i>	162.53	PBDB + regression
<i>Haplomylus speirianus</i>	98.28	PBDB + regression
<i>Haplomys galbreathi</i>	38.23	PBDB + regression
<i>Haplomys liolophus</i>	39.00	[198]
<i>Harpagolestes leotensis</i>	138880.62	PBDB + regression
<i>Harpagolestes uintensis</i>	106686.23	PBDB + regression
<i>Harrymys irvini</i>	83.93	[329]
<i>Harrymys magnus</i>	50.40	[329]
<i>Harrymys woodi</i>	43.52	[159]
<i>Heliscomys hatcheri</i>	7.19	[362]
<i>Heliscomys ostranderi</i>	3.56	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Heliscomys senex</i>	4.67	PBDB + regression
<i>Heliscomys vetus</i>	4.67	PBDB + regression
<i>Heliscomys woodi</i>	10.39	PBDB + regression
<i>Hemiacodon engardae</i>	538.07	PBDB + regression
<i>Hemiacodon gracilis</i>	563.76	PBDB + regression
<i>Hemiauchenia gracilis</i>	157598.86	PBDB + regression
<i>Hemiauchenia macrocephala</i>	109900.58	[305]
<i>Hemiauchenia minima</i>	85819.37	[329]
<i>Hemiauchenia vera</i>	243143.10	PBDB + regression
<i>Hemipsalodon grandis</i>	436751.47	[280]
<i>Hemithlaeus harbourae</i>	1283.98	[180]
<i>Hendryomeryx defordi</i>	5742.28	PBDB
<i>Hendryomeryx esulcatus</i>	6619.36	PBDB + regression
<i>Hendryomeryx wilsoni</i>	3001.58	PBDB + regression
<i>Heptacodon pellionis</i>	88438.62	PBDB + regression
<i>Heptodon calciculus</i>	26551.76	PBDB + regression
<i>Herpetotherium fugax</i>	19.74	PBDB + regression
<i>Herpetotherium knighti</i>	51.17	PBDB + regression
<i>Herpetotherium merriami</i>	25.80	PBDB + regression
<i>Herpetotherium valens</i>	57.16	PBDB + regression
<i>Herpetotherium youngi</i>	40.09	PBDB + regression
<i>Hesperhys pinensis</i>	68186.37	[329]
<i>Hesperhys vagrans</i>	73865.41	[329]
<i>Hesperocyon gregarius</i>	3533.34	[329]
<i>Hesperolagomys fluviatilis</i>	169.02	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Hesperolagomys galbreathi</i>	149.90	[329]
<i>Hesperoscalops mcgrewi</i>	186.62	PBDB + regression
<i>Heteraletes leotanus</i>	14016.35	PBDB + regression
<i>Heteromeryx dispar</i>	51072.86	PBDB + regression
<i>Heteroplohippus hulberti</i>	302549.45	[329]
<i>Hexacodus pelodes</i>	625.82	PBDB + regression
<i>Hexameryx simpsoni</i>	30638.11	[329]
<i>Hexobelomeryx fricki</i>	59621.29	PBDB + regression
<i>Hibbarderix obfuscatus</i>	33.60	PBDB + regression
<i>Hibbardiomyx fayae</i>	61.14	PBDB + regression
<i>Hibbardiomyx marthae</i>	84.39	PBDB + regression
<i>Hibbardiomyx skinneri</i>	61.14	PBDB + regression
<i>Hibbardiomyx voorhiesi</i>	61.14	PBDB + regression
<i>Hipparion forcei</i>	194852.86	[329]
<i>Hipparion tehonense</i>	100709.96	[329]
<i>Hippotherium emsliei</i>	201622.66	PBDB + regression
<i>Hippotherium ingenuum</i>	109258.44	PBDB + regression
<i>Hippotherium plicatile</i>	165091.29	PBDB + regression
<i>Hippotherium quinni</i>	422193.86	PBDB + regression
<i>Hitonkala macdonaltau</i>	24.53	[329]
<i>Holmesina floridanus</i>	68696.59	PBDB + regression
<i>Homacodon vagans</i>	6691.69	PBDB + regression
<i>Homogalax protapirinus</i>	19815.58	PBDB + regression
<i>Homotherium crusafonti</i>	233968.19	PBDB
<i>Homotherium idahoensis</i>	257900.00	[198]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Homotherium johnstoni</i>	124571.99	PBDB + regression
<i>Hoplophoneus mentalis</i>	48420.73	PBDB + regression
<i>Hoplophoneus primaevus</i>	20243.49	PBDB + regression
<i>Huerfanodon polecatensis</i>	29749.42	PBDB + regression
<i>Huerfanodon torrejonius</i>	20823.53	PBDB + regression
<i>Hyaenodon brevirostrus</i>	37331.34	PBDB + regression
<i>Hyaenodon crucians</i>	14771.98	PBDB + regression
<i>Hyaenodon horridus</i>	91750.41	PBDB + regression
<i>Hyaenodon montanus</i>	28357.49	PBDB + regression
<i>Hyaenodon mustelinus</i>	8079.01	PBDB + regression
<i>Hyaenodon raineyi</i>	1093.34	PBDB + regression
<i>Hyaenodon venturae</i>	2552.88	PBDB + regression
<i>Hyaenodon vetus</i>	33333.28	PBDB + regression
<i>Hylomeryx quadricuspis</i>	4966.19	PBDB + regression
<i>Hyopsodus fastigatus</i>	1894.05	PBDB + regression
<i>Hyopsodus lepidus</i>	625.82	PBDB + regression
<i>Hyopsodus loomisi</i>	488.31	PBDB + regression
<i>Hyopsodus lysitensis</i>	924.51	PBDB + regression
<i>Hyopsodus mentalis</i>	1482.35	PBDB + regression
<i>Hyopsodus minor</i>	440.75	PBDB + regression
<i>Hyopsodus minusculus</i>	440.75	PBDB + regression
<i>Hyopsodus paulus</i>	768.59	PBDB + regression
<i>Hyopsodus pauxillus</i>	181.54	PBDB + regression
<i>Hyopsodus powellianus</i>	1416.00	PBDB + regression
<i>Hyopsodus simplex</i>	729.22	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Hyopsodus tonksii</i>	464.26	PBDB + regression
<i>Hyopsodus uintensis</i>	1495.78	PBDB + regression
<i>Hyopsodus walcottianus</i>	6172.52	PBDB + regression
<i>Hyopsodus wortmani</i>	798.73	PBDB + regression
<i>Hypertragulus calcaratus</i>	8880.00	[198]
<i>Hypertragulus hesperius</i>	4572.00	[198]
<i>Hypohippus affinis</i>	442413.39	[329]
<i>Hypohippus equinus</i>	271034.12	[329]
<i>Hypohippus osborni</i>	299539.03	[329]
<i>Hypolagus edensis</i>	665.14	[329]
<i>Hypolagus fontinalis</i>	1211.97	[329]
<i>Hypolagus furlongi</i>	678.58	[329]
<i>Hypolagus gidleyi</i>	1998.20	[329]
<i>Hypolagus oregonensis</i>	2344.90	[329]
<i>Hypolagus parviplicatus</i>	1702.75	[329]
<i>Hypolagus ringoldensis</i>	319.00	[198]
<i>Hypolagus vetus</i>	2892.86	[329]
<i>Hypolagus voorhiesi</i>	319.00	[198]
<i>Hypsipops bannackensis</i>	146895.45	[355]
<i>Hypsipops breviceps</i>	156373.08	[329]
<i>Hyrachyus affinis</i>	111138.67	PBDB + regression
<i>Hyrachyus eximius</i>	198513.08	PBDB + regression
<i>Hyrachyus modestus</i>	217770.66	PBDB + regression
<i>Hyracodon leidyanus</i>	459564.25	PBDB + regression
<i>Hyracodon mediocris</i>	102488.96	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Hyracodon nebrascensis</i>	211081.59	[329]
<i>Hyracodon petersoni</i>	313951.52	PBDB + regression
<i>Hyracodon priscidens</i>	351463.36	PBDB + regression
<i>Hyracotherium cristatum</i>	23898.63	PBDB
<i>Hyracotherium vasacciense</i>	24900.00	[188]
<i>Ictidopappus mustelinus</i>	354.83	PBDB + regression
<i>Ignacius fremontensis</i>	19.74	PBDB + regression
<i>Ignacius frugivorus</i>	73.44	PBDB + regression
<i>Ignacius graybullianus</i>	128.62	PBDB + regression
<i>Indarctos nevadensis</i>	376215.79	PBDB + regression
<i>Indarctos oregonensis</i>	302549.45	[329]
<i>Ischyrocyon gidleyi</i>	282095.23	[329]
<i>Ischyromys typus</i>	761.25	PBDB + regression
<i>Ischyromys veterior</i>	477.49	PBDB + regression
<i>Jacobsomys verdensis</i>	50.91	[329]
<i>Janimus dawsonae</i>	17.97	PBDB + regression
<i>Jaywilsonomys ojinagaensis</i>	895.10	PBDB + regression
<i>Jemezius szalayi</i>	95.39	PBDB + regression
<i>Jepsenella praepropera</i>	18.80	PBDB + regression
<i>Jimomys labaughii</i>	69.41	[329]
<i>Jimomys lulli</i>	28.00	[198]
<i>Kansasimys dubius</i>	135.64	[329]
<i>Kansasimys wilsoni</i>	228.43	[193]
<i>Kimbetohia mziae</i>	72.00	[362]
<i>Knightomys cremneus</i>	27.28	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Knightomys cuspidatus</i>	48.29	PBDB
<i>Knightomys depressus</i>	64.79	PBDB + regression
<i>Knightomys huerfanensis</i>	135.17	PBDB + regression
<i>Knightomys minor</i>	27.28	PBDB + regression
<i>Knightomys reginensis</i>	10.39	PBDB + regression
<i>Knightomys senior</i>	44.30	PBDB + regression
<i>Kolponomos clallamensis</i>	103303.51	[330]
<i>Kolponomos newportensis</i>	876418.72	[278]
<i>Kyptoceras amatorum</i>	369534.73	[329]
<i>Labidolemur kayi</i>	33.60	PBDB + regression
<i>Labidolemur serus</i>	67.38	PBDB + regression
<i>Labidolemur soricoides</i>	14.63	PBDB + regression
<i>Lambdotherium popoagicum</i>	29620.91	PBDB + regression
<i>Lambertocyon eximius</i>	6045.26	PBDB + regression
<i>Lambertocyon ischyrus</i>	2015.61	PBDB + regression
<i>Lanthanotherium sawini</i>	90.14	PBDB + regression
<i>Laredochoerus edwardsi</i>	5523.91	PBDB + regression
<i>Laredomys riograndensis</i>	5.92	PBDB + regression
<i>Leidymys cerasus</i>	41.68	[329]
<i>Leidymys korthi</i>	35.35	PBDB + regression
<i>Leipsanolestes siegfriedti</i>	26.62	PBDB + regression
<i>Lemoynea biradicularis</i>	45.60	[329]
<i>Lepoides lepoides</i>	3640.95	[329]
<i>Leptacodon munusculum</i>	6.56	PBDB + regression
<i>Leptacodon packi</i>	14.63	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Leptacodon tener</i>	8.61	PBDB + regression
<i>Leptarctus mummorum</i>	8673.78	PBDB + regression
<i>Leptarctus oregonensis</i>	2580.00	[198]
<i>Leptarctus primus</i>	149.90	[329]
<i>Leptarctus supremus</i>	7725.15	PBDB
<i>Leptauchenia decora</i>	20130.67	[329]
<i>Leptauchenia major</i>	30946.03	[329]
<i>Leptictis dakotensis</i>	644.52	PBDB + regression
<i>Leptochoerus elegans</i>	8623.90	PBDB + regression
<i>Leptochoerus spectabilis</i>	9021.94	PBDB + regression
<i>Leptochoerus supremus</i>	19397.68	PBDB + regression
<i>Leptocyon mollis</i>	3300.00	[198]
<i>Leptocyon vafer</i>	5377.61	[329]
<i>Leptodontomys douglassi</i>	4.67	PBDB + regression
<i>Leptodontomys stirtoni</i>	7.32	[323]
<i>Leptolambda schmidti</i>	90362.43	[369]
<i>Leptomeryx blacki</i>	4578.72	PBDB + regression
<i>Leptomeryx elissae</i>	6511.38	PBDB + regression
<i>Leptomeryx evansi</i>	9426.05	PBDB + regression
<i>Leptomeryx mammifer</i>	18828.04	PBDB + regression
<i>Leptomeryx speciosus</i>	9223.24	PBDB + regression
<i>Leptomeryx yoderi</i>	10716.49	PBDB + regression
<i>Leptoreodon edwardsi</i>	13277.57	PBDB + regression
<i>Leptoreodon leptolophus</i>	9223.24	PBDB + regression
<i>Leptoreodon major</i>	32615.84	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Leptoreodon marshi</i>	20660.42	PBDB + regression
<i>Leptoreodon pusillus</i>	3957.21	PBDB + regression
<i>Leptoreodon stocki</i>	20980.32	PBDB + regression
<i>Leptromys wilsoni</i>	110.81	PBDB + regression
<i>Leptotomus caryophilus</i>	872.14	PBDB + regression
<i>Leptotomus leptodus</i>	2852.74	PBDB + regression
<i>Leptotomus parvus</i>	941.80	PBDB + regression
<i>Leptotragulus clarki</i>	5364.24	PBDB + regression
<i>Leptotragulus medioides</i>	13323.34	PBDB + regression
<i>Leptotragulus proavus</i>	11797.49	PBDB + regression
<i>Lepus californicus</i>	2288.00	PBDB
<i>Lignimus austridakotensis</i>	14.88	[329]
<i>Lignimus montis</i>	38.23	PBDB + regression
<i>Limaconyssus habrus</i>	18.80	PBDB + regression
<i>Limnoecus niobrarensis</i>	7.17	[329]
<i>Limnoecus tricuspis</i>	5.10	[329]
<i>Liodontia alexandrae</i>	198.34	[329]
<i>Litaletes disjunctus</i>	681.34	PBDB + regression
<i>Litocherus lacunatus</i>	84.86	PBDB + regression
<i>Litocherus notissimus</i>	37.32	PBDB + regression
<i>Litocherus zygeus</i>	55.87	PBDB + regression
<i>Litolagus molidens</i>	103.28	PBDB + regression
<i>Litolestes ignotus</i>	18.80	PBDB + regression
<i>Litomylus dissentaneus</i>	124.29	PBDB + regression
<i>Litomylus orthronepius</i>	115.36	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Litoyoderimys auogoleus</i>	47.48	PBDB + regression
<i>Longirostromeryx clarendonensis</i>	13226.80	[329]
<i>Longirostromeryx wellsi</i>	17500.77	[329]
<i>Lophiparamys debequensis</i>	29.87	PBDB + regression
<i>Lophiparamys murinus</i>	88.57	PBDB + regression
<i>Loveina minuta</i>	45.48	PBDB + regression
<i>Loveina zephyri</i>	66.71	PBDB + regression
<i>Loxolophus criswelli</i>	2140.57	PBDB + regression
<i>Loxolophus hyattianus</i>	1984.90	PBDB + regression
<i>Loxolophus pentacus</i>	14847.99	PBDB + regression
<i>Loxolophus priscus</i>	3899.92	PBDB + regression
<i>Loxolophus schizophrenus</i>	881.66	PBDB + regression
<i>Loxolophus spiekeri</i>	18323.92	PBDB + regression
<i>Lutravus halli</i>	454.86	[329]
<i>Lycophocyon hutchisoni</i>	4865.03	PBDB + regression
<i>Lynx proterolyncis</i>	15677.78	[329]
<i>Lynx rufus</i>	10482.00	PBDB
<i>Machaeromeryx gilchristensis</i>	4536.90	[329]
<i>Machaeromeryx tragulus</i>	4536.90	[329]
<i>Macrocranion junnei</i>	15.97	PBDB + regression
<i>Macrocranion nitens</i>	33.60	PBDB + regression
<i>Macrogenis crassigenis</i>	106737.50	PBDB
<i>Macrognathomys gemmacolis</i>	5.92	PBDB + regression
<i>Macrognathomys nanus</i>	5.16	[329]
<i>Macrotarsius montanus</i>	1152.46	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Macrotarsius roederi</i>	1640.00	[312]
<i>Macrotarsius siegerti</i>	1188.60	PBDB + regression
<i>Mahgarita stevensi</i>	358.28	[365]
<i>Malaquiferus tourteloti</i>	14816.10	PBDB + regression
<i>Mammacyon obtusidens</i>	70262.96	[329]
<i>Mammut americanum</i>	4528975.80	[305]
<i>Mammut cosoensis</i>	29076021.42	[281]
<i>Mammut furlongi</i>	29535033.05	PBDB + regression
<i>Mammuthus columbi</i>	7998342.55	[305]
<i>Mammut raki</i>	41795334.69	PBDB + regression
<i>Marmota korthi</i>	2038.56	[329]
<i>Marmota vetus</i>	658.52	[329]
<i>Martes gazini</i>	1767.00	[198]
<i>Martes parviloba</i>	5831.67	[186]
<i>Martinogale alveodens</i>	17.99	[329]
<i>Mattimys kalicola</i>	32.56	PBDB + regression
<i>Megacamelus merriami</i>	1905014.16	[329]
<i>Megadelphus lundeliusi</i>	3614.01	PBDB + regression
<i>Megahippus matthewi</i>	882046.45	[329]
<i>Megalagus abaconis</i>	1118.79	[329]
<i>Megalagus brachyodon</i>	373.57	PBDB + regression
<i>Megalagus primitivus</i>	3165.29	[329]
<i>Megalagus turgidus</i>	3071.74	[329]
<i>Megalesthonyx hopsoni</i>	15975.00	[118]
<i>Megalictis ferox</i>	1587.63	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Megalictis frazieri</i>	33591.60	PBDB
<i>Megalonyx curvidens</i>	185050.00	[194]
<i>Megalonyx leptostomus</i>	279404.72	PBDB + regression
<i>Megalonyx wheatleyi</i>	252445.15	PBDB + regression
<i>Megantereon hesperus</i>	67507.91	[329]
<i>Megapeomys bobwilsoni</i>	411.58	[329]
<i>Megasminthus tiheni</i>	55.70	[329]
<i>Megatylopus cochrani</i>	884486.95	PBDB + regression
<i>Megatylopus matthewi</i>	1383324.20	[329]
<i>Megatylopus primaevus</i>	884295.95	PBDB + regression
<i>Meniscomys hippodus</i>	69.41	[329]
<i>Meniscomys uhtoffi</i>	70.00	[198]
<i>Meniscotherium chamense</i>	17413.77	PBDB + regression
<i>Meniscotherium tapiacitum</i>	1864.20	PBDB + regression
<i>Menoceras arikarense</i>	597195.61	[329]
<i>Menoceras barbouri</i>	1251683.50	[329]
<i>Menops bakeri</i>	3993312.11	PBDB
<i>Mentoclaenodon acrogenius</i>	37446.92	PBDB + regression
<i>Mephitis mephitis</i>	2055.00	PBDB
<i>Merriamoceros coronatus</i>	15214.44	[329]
<i>Merychippus brevidontus</i>	124243.67	[329]
<i>Merychippus calamarius</i>	196811.17	[329]
<i>Merychippus californicus</i>	86681.87	[329]
<i>Merychippus goorisi</i>	222888.24	PBDB + regression
<i>Merychippus gunteri</i>	92041.97	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Merychippus insignis</i>	125492.34	[329]
<i>Merychippus primus</i>	95798.28	[329]
<i>Merychippus relictus</i>	60471.00	[198]
<i>Merychippus sejunctus</i>	75357.60	[329]
<i>Merychyus arenarum</i>	52892.75	PBDB + regression
<i>Merychyus crabilli</i>	6038.43	[154]
<i>Merychyus elegans</i>	45706.69	[329]
<i>Merychyus mediuss</i>	108012.26	[329]
<i>Merychyus minimus</i>	29620.91	PBDB + regression
<i>Merychyus relictus</i>	20923.07	PBDB
<i>Merychyus smithi</i>	135340.58	PBDB + regression
<i>Merycochoerus carrikeri</i>	313334.13	[265]
<i>Merycochoerus chelydra</i>	289450.77	PBDB
<i>Merycochoerus magnus</i>	331041.82	[329]
<i>Merycochoerus matthewi</i>	59736.09	PBDB + regression
<i>Merycochoerus proprius</i>	265667.29	[329]
<i>Merycochoerus superbus</i>	438400.00	[198]
<i>Merycodus prodromus</i>	8738.23	[157]
<i>Merycodus sabulonis</i>	10509.13	[329]
<i>Merycodus warreni</i>	12597.43	[19]
<i>Merycoides harrisonensis</i>	90219.42	[329]
<i>Merycoides longiceps</i>	56672.65	[324]
<i>Merycoides pariogonus</i>	77000.00	[198]
<i>Merycoidodon bullatus</i>	109500.00	[198]
<i>Merycoidodon culbertsoni</i>	116748.58	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Merycoidodon major</i>	62200.00	[198]
<i>Mescalerolemur horneri</i>	193.21	[185]
<i>Mesocyon brachyops</i>	7942.63	[329]
<i>Mesocyon coryphaeus</i>	10198.54	[329]
<i>Mesocyon temnodon</i>	7186.79	[329]
<i>Mesodma ambigua</i>	115.64	PBDB + regression
<i>Mesodma formosa</i>	32.59	PBDB + regression
<i>Mesodma garfieldensis</i>	37.51	PBDB + regression
<i>Mesodma pygmaea</i>	8.53	PBDB + regression
<i>Mesodma thompsoni</i>	54.13	PBDB + regression
<i>Mesogaulus paniensis</i>	926.80	PBDB
<i>Mesohippus bairdi</i>	39223.29	PBDB + regression
<i>Mesohippus exoletus</i>	47569.37	PBDB + regression
<i>Mesohippus texanus</i>	35643.43	PBDB + regression
<i>Mesohippus westoni</i>	39355.66	PBDB + regression
<i>Mesonyx obtusidens</i>	147850.73	PBDB
<i>Mesoreodon chelonyx</i>	133078.52	[24]
<i>Mesoreodon floridensis</i>	104820.01	[329]
<i>Mesoreodon minor</i>	159700.00	[198]
<i>Mesoscalops montanensis</i>	85.63	[329]
<i>Mesoscalops scopelotemos</i>	95.58	[329]
<i>Metadjidaumo hendryi</i>	10.38	[329]
<i>Metaliomys sevierensis</i>	15.90	PBDB + regression
<i>Metanoiamys agorus</i>	12.11	PBDB + regression
<i>Metanoiamys fantasma</i>	20.14	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Metanoiamys korthi</i>	13.95	PBDB + regression
<i>Metanoiamys lacus</i>	10.39	PBDB + regression
<i>Metanoiamys marinus</i>	8.78	PBDB + regression
<i>Metarhinus pater</i>	1098772.19	PBDB
<i>Metatomarctus canavus</i>	10938.02	[329]
<i>Metechinus amplior</i>	567.90	[328]
<i>Miacis deutschi</i>	968.28	PBDB + regression
<i>Miacis exiguum</i>	1775.93	PBDB + regression
<i>Miacis hookwayi</i>	2350.87	[221]
<i>Miacis latidens</i>	2236.53	PBDB + regression
<i>Miacis parvivorus</i>	2384.02	PBDB + regression
<i>Miacis petilus</i>	1591.36	PBDB + regression
<i>Michenia agatensis</i>	99707.88	[329]
<i>Michenia exilis</i>	65512.75	[329]
<i>Microcosmodon conus</i>	16.11	PBDB + regression
<i>Microcosmodon rosei</i>	27.99	PBDB + regression
<i>Microeutypomys karenae</i>	12.11	PBDB + regression
<i>Microeutypomys tilliei</i>	22.41	PBDB + regression
<i>Micromomys antelucanus</i>	5.04	PBDB
<i>Microparamys cheradius</i>	57.59	PBDB + regression
<i>Microparamys dubius</i>	20.14	PBDB + regression
<i>Microparamys hunterae</i>	14.33	PBDB
<i>Microparamys minutus</i>	24.79	PBDB + regression
<i>Microparamys nimius</i>	29.87	PBDB + regression
<i>Microparamys sambucus</i>	41.22	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Microparamys tricus</i>	57.59	PBDB + regression
<i>Microparamys woodi</i>	32.56	PBDB + regression
<i>Micropternodus borealis</i>	41.19	PBDB + regression
<i>Micropternodus montrosensis</i>	103.85	PBDB + regression
<i>Micropternodus morgani</i>	72.24	[329]
<i>Microsus cuspidatus</i>	2597.76	PBDB + regression
<i>Microsyops angustidens</i>	352.30	PBDB + regression
<i>Microsyops annectens</i>	1522.82	PBDB + regression
<i>Microsyops cardiorestes</i>	186.75	PBDB + regression
<i>Microsyops elegans</i>	738.98	PBDB + regression
<i>Microsyops knightensis</i>	521.24	PBDB + regression
<i>Microsyops kratos</i>	1455.64	PBDB + regression
<i>Microsyops latidens</i>	472.21	PBDB + regression
<i>Microsyops scottianus</i>	634.92	PBDB + regression
<i>Microtomarctus conferta</i>	8866.19	[329]
<i>Mictomys vetus</i>	76.28	PBDB + regression
<i>Mimatuta morgoth</i>	512.92	PBDB + regression
<i>Mimetodon silberlingi</i>	21.68	PBDB + regression
<i>Mimomys mcknighti</i>	110.70	PBDB
<i>Mimomys panacaensis</i>	84.39	PBDB + regression
<i>Mimomys parvus</i>	99.10	PBDB
<i>Mimomys primus</i>	163.60	PBDB
<i>Mimomys taylori</i>	80.90	PBDB
<i>Mimoperadectes labrus</i>	285.50	PBDB + regression
<i>Mimotricentes fremontensis</i>	2674.19	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Mimotricentes subtrigonus</i>	1984.90	PBDB + regression
<i>Miniochoerus affinis</i>	35835.68	PBDB
<i>Miniochoerus gracilis</i>	15437.78	PBDB
<i>Minippus index</i>	4819.64	PBDB
<i>Mioclaenius turgidus</i>	4412.84	PBDB + regression
<i>Miocyon scotti</i>	41776.02	PBDB + regression
<i>Miocyon vallisrubrae</i>	14244.90	PBDB + regression
<i>Mioheteromys amplissimus</i>	41.26	[329]
<i>Miohippus anceps</i>	41650.00	[198]
<i>Miohippus assiniboiensis</i>	78519.40	PBDB + regression
<i>Miohippus equiceps</i>	41650.00	[198]
<i>Miohippus equianus</i>	35954.16	[329]
<i>Miohippus gidleyi</i>	129635.34	PBDB + regression
<i>Miohippus grandis</i>	76252.18	PBDB
<i>Miohippus intermedius</i>	82454.34	[329]
<i>Miohippus obliquidens</i>	52575.21	[329]
<i>Miolabis fricki</i>	240385.70	[329]
<i>Miomustela madisonae</i>	13.46	[329]
<i>Mionictis elegans</i>	6652.51	PBDB
<i>Mionictis incertus</i>	9269.73	PBDB
<i>Mionictis letifer</i>	134.29	[329]
<i>Mionictis pristinus</i>	32167.60	PBDB + regression
<i>Miosciurus ballovianus</i>	36.60	[329]
<i>Miosicista angulus</i>	8.15	PBDB
<i>Miospermophilus lavertyi</i>	106.70	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Miospermophilus wyomingensis</i>	83.10	[329]
<i>Miotapirus harrisonensis</i>	275463.83	PBDB + regression
<i>Miolytulus gibbi</i>	48050.12	[329]
<i>Miolytulus leonardi</i>	43044.94	[329]
<i>Miolytulus taylori</i>	73865.41	[329]
<i>Miracinonyx inexpectatus</i>	41653.42	PBDB + regression
<i>Miracinonyx studeri</i>	25529.71	PBDB + regression
<i>Mithrandir gillianus</i>	644.09	PBDB + regression
<i>Mixodectes malaris</i>	538.07	PBDB + regression
<i>Mixodectes pungens</i>	1563.78	PBDB + regression
<i>Mojavemys galushai</i>	28.22	[329]
<i>Mojavemys lophatus</i>	15.19	PBDB
<i>Mojavemys magnumarcus</i>	56.26	[329]
<i>Montanatylopus matthewi</i>	176770.65	PBDB + regression
<i>Mookomys altifluminis</i>	13.07	[329]
<i>Mookomys formicarum</i>	11.86	PBDB
<i>Mookomys thrinax</i>	11.94	[329]
<i>Moropus elatus</i>	707858.86	[329]
<i>Moropus merriami</i>	173767.00	[198]
<i>Moropus oregonensis</i>	189094.09	[329]
<i>Mustela frenata</i>	146.89	[305]
<i>Mustela meltoni</i>	895.71	PBDB + regression
<i>Mustela rexroadensis</i>	9.58	[329]
<i>Mylanodon rosei</i>	27.99	PBDB + regression
<i>Mylohyus elmorei</i>	168990.43	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Myrmecoboides montanensis</i>	45.21	PBDB + regression
<i>Myrmecophaga tridactyla</i>	32544.00	PBDB
<i>Mysops parvus</i>	54.13	PBDB + regression
<i>Mystipterus martini</i>	29.37	[329]
<i>Mystipterus pacificus</i>	14.88	[329]
<i>Mytonolagus petersoni</i>	213.83	PBDB + regression
<i>Mytonolagus wyomingensis</i>	84.09	PBDB + regression
<i>Mytonomeryx scotti</i>	5465.38	PBDB + regression
<i>Mytonomys burkei</i>	2757.96	PBDB + regression
<i>Mytonomys mytonensis</i>	1177.52	PBDB + regression
<i>Mytonomys robustus</i>	2795.71	PBDB + regression
<i>Namatomys lloydii</i>	24.79	PBDB + regression
<i>Nannippus aztecus</i>	106272.86	PBDB + regression
<i>Nannippus beckensis</i>	137564.87	PBDB + regression
<i>Nannippus peninsulatus</i>	68871.66	[329]
<i>Nannippus westoni</i>	62317.65	[329]
<i>Nannodectes gidleyi</i>	521.24	PBDB + regression
<i>Nannodectes intermedius</i>	133.09	PBDB + regression
<i>Nannodectes simpsoni</i>	402.81	PBDB + regression
<i>Nanodelphys huntii</i>	7.27	PBDB + regression
<i>Nanotragulus loomisi</i>	2724.39	[329]
<i>Nanotragulus ordinatus</i>	5541.39	[329]
<i>Nanotragulus planiceps</i>	3678.00	[198]
<i>Navajovius kohlhaasae</i>	12.82	PBDB + regression
<i>Nebraskomys mcgrewi</i>	46.06	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Nebraskomys rexroadensis</i>	47.48	PBDB + regression
<i>Nekrolagus progressus</i>	1510.20	[329]
<i>Neohipparion affine</i>	167711.41	[329]
<i>Neohipparion eurystyle</i>	133252.35	[329]
<i>Neohipparion leptode</i>	233281.23	[329]
<i>Neohipparion trampasense</i>	120571.71	[329]
<i>Neoliotomus conventus</i>	1894.05	PBDB + regression
<i>Neoliotomus ultimus</i>	2252.70	PBDB + regression
<i>Neoplagiaulax donaldorum</i>	52.00	[362]
<i>Neoplagiaulax grangeri</i>	93.00	[362]
<i>Neoplagiaulax hazeni</i>	95.39	PBDB + regression
<i>Neoplagiaulax hunteri</i>	42.74	PBDB + regression
<i>Neoplagiaulax macrotomeus</i>	14.42	PBDB + regression
<i>Neoplagiaulax mckennai</i>	60.27	PBDB + regression
<i>Neotoma cinerea</i>	299.23	[305]
<i>Neotoma fossilis</i>	161.61	PBDB + regression
<i>Neotoma leucopetrica</i>	389.66	[110]
<i>Neotoma quadriplicata</i>	477.49	PBDB + regression
<i>Neotoma sawrockensis</i>	99.48	[329]
<i>Neotoma taylori</i>	190.07	PBDB + regression
<i>Neotoma vaughani</i>	190.07	PBDB + regression
<i>Nerterogeomys anzensis</i>	80.55	PBDB
<i>Nerterogeomys garbanii</i>	157.02	PBDB
<i>Nerterogeomys minor</i>	38.23	PBDB + regression
<i>Nerterogeomys persimilis</i>	49.62	[110]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Nexuotapirus marslandensis</i>	144350.55	[329]
<i>Nexuotapirus robustus</i>	302549.45	[329]
<i>Niglarodon koernerri</i>	68.03	[329]
<i>Nimravides galiani</i>	159532.03	[329]
<i>Nimravides thinobates</i>	9582364.97	PBDB
<i>Nimravus brachyops</i>	3789.54	[329]
<i>Nimravus sectator</i>	112444.51	PBDB + regression
<i>Niptomomys doreenae</i>	17.88	PBDB + regression
<i>Niptomomys thelmae</i>	19.74	PBDB + regression
<i>Nonomys gutzleri</i>	20.14	PBDB + regression
<i>Nonomys simplicidens</i>	15.90	PBDB + regression
<i>Notharctus pugnax</i>	3034.76	PBDB + regression
<i>Notharctus robinsoni</i>	3318.67	PBDB + regression
<i>Notharctus robustior</i>	6900.00	[312]
<i>Notharctus tenebrosus</i>	4325.36	PBDB + regression
<i>Notharctus ventalinus</i>	2691.73	PBDB + regression
<i>Nothokemas floridanus</i>	100709.96	[329]
<i>Nothokemas waldropi</i>	22925.38	[329]
<i>Nothotylopus campognathus</i>	334368.85	[329]
<i>Nothrotheriops shastensis</i>	613762.01	[30]
<i>Nothrotheriops texanus</i>	124768.59	PBDB + regression
<i>Notiosorex jacksoni</i>	11.14	[189]
<i>Notiosorex repenningi</i>	17.08	[167]
<i>Notiotitanops mississippiensis</i>	4455686.16	PBDB + regression
<i>Notolagus lepusculus</i>	372.41	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Nototamias hulberti</i>	17.12	[329]
<i>Nototamias quadratus</i>	35.87	[329]
<i>Nyctitherium serotinum</i>	12.08	PBDB + regression
<i>Nyctitherium velox</i>	24.97	PBDB + regression
<i>Ochotona spanglei</i>	188.67	[329]
<i>Odocoileus virginianus</i>	52607.00	PBDB
<i>Ogmodontomys poaphagus</i>	131.63	[329]
<i>Ogmodontomys sawrockensis</i>	140.40	PBDB
<i>Oklahomalagus whisenhunti</i>	387.61	[329]
<i>Oligobunis floridanus</i>	24237.19	PBDB + regression
<i>Oligoryctes cameronensis</i>	4.73	PBDB + regression
<i>Oligoscalops galbreathi</i>	15.04	PBDB + regression
<i>Oligospermophilus douglassi</i>	115.51	PBDB + regression
<i>Omomys carteri</i>	171.31	PBDB + regression
<i>Omomys lloydii</i>	66.71	PBDB + regression
<i>Onychomys holisteri</i>	20.14	PBDB + regression
<i>Onychomys martini</i>	18.36	[329]
<i>Onychomys pedroensis</i>	36.23	[329]
<i>Oregonomys magnus</i>	20.14	PBDB + regression
<i>Oregonomys pebblespringsensis</i>	26.58	[329]
<i>Oregonomys sargentii</i>	26.05	[329]
<i>Oreodontoides oregonensis</i>	25336.47	[329]
<i>Oreolagus colteri</i>	48.28	PBDB + regression
<i>Oreolagus nebrascensis</i>	336.97	[329]
<i>Oreolagus nevadensis</i>	78.00	[198]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Oreolagus wallacei</i>	354.25	[329]
<i>Oreotalpa florissantensis</i>	10.87	PBDB + regression
<i>Orohippus pumilus</i>	11144.64	PBDB + regression
<i>Orohippus sylvaticus</i>	10210.29	PBDB + regression
<i>Oromeryx plicatus</i>	8545.03	PBDB + regression
<i>Oropyctis pediasius</i>	161.61	PBDB + regression
<i>Osbornoceros osborni</i>	17326.63	[329]
<i>Osbornodon brachypus</i>	12282.74	[43]
<i>Osbornodon fricki</i>	25848.30	[329]
<i>Osbornodon iamonensis</i>	13904.95	[329]
<i>Osbornodon renjiei</i>	9337.04	PBDB + regression
<i>Osbornodon scitulus</i>	11849.01	[329]
<i>Osbornodon sesnoni</i>	8349.86	[329]
<i>Otarocyon cooki</i>	1826.21	[329]
<i>Otarocyon macdonaldi</i>	1270.84	PBDB + regression
<i>Ottoceros peacevalleyensis</i>	13493.99	[329]
<i>Ourayia hopsoni</i>	590.00	PBDB + regression
<i>Ourayia uintensis</i>	758.66	PBDB + regression
<i>Oxetocyon cuspidatus</i>	2440.60	[329]
<i>Oxyacodon agapetillus</i>	352.30	PBDB + regression
<i>Oxyacodon apiculatus</i>	1116.82	PBDB + regression
<i>Oxyacodon ferronensis</i>	616.78	PBDB + regression
<i>Oxyacodon priscilla</i>	808.90	PBDB + regression
<i>Oxyaena forcipata</i>	49289.23	PBDB + regression
<i>Oxyaena gulo</i>	42202.38	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Oxyaena intermedia</i>	44676.22	PBDB + regression
<i>Oxycleaenus cuspidatus</i>	760.59	PBDB + regression
<i>Oxycleaenus pugnax</i>	1812.53	PBDB + regression
<i>Oxycleaenus simplex</i>	566.68	PBDB + regression
<i>Oxydactylus longipes</i>	112420.32	[329]
<i>Oxydactylus lulli</i>	140198.81	PBDB
<i>Oxyprimus erikseni</i>	133.09	PBDB + regression
<i>Pachyaena gigantea</i>	71758.33	PBDB + regression
<i>Pachyaena gracilis</i>	38119.28	PBDB + regression
<i>Pachyaena ossifraga</i>	460896.74	PBDB
<i>Pachyarmatherium leiseyi</i>	15420.00	[194]
<i>Paciculus montanus</i>	68.53	PBDB + regression
<i>Paciculus nebraskensis</i>	80.64	[329]
<i>Paciculus woodi</i>	32.63	[162]
<i>Paenemarmota barbouri</i>	10301.04	[329]
<i>Paenemarmota mexicana</i>	10977.77	PBDB + regression
<i>Paenemarmota nevadensis</i>	7644.00	[198]
<i>Paenemarmota sawrockensis</i>	5943.18	[329]
<i>Palaechthon alticuspis</i>	51.17	PBDB + regression
<i>Palaechthon woodi</i>	27.99	PBDB + regression
<i>Palaeictops bicuspis</i>	143.19	[293]
<i>Palaeictops bridgeri</i>	139.74	PBDB + regression
<i>Palaeictops multicuspis</i>	149.35	PBDB + regression
<i>Palaeogale dorothiae</i>	607.89	[329]
<i>Palaeogale minuta</i>	400.00	NOW

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Palaeogale sectoria</i>	716.63	PBDB
<i>Palaeolagus burkei</i>	307.97	[329]
<i>Palaeolagus hemirhizis</i>	128.62	PBDB + regression
<i>Palaeolagus hypsodus</i>	678.58	[329]
<i>Palaeolagus philoi</i>	1211.97	[329]
<i>Palaeolagus primus</i>	119.90	PBDB + regression
<i>Palaeolagus temnodon</i>	171.31	PBDB + regression
<i>Palaeonictis occidentalis</i>	36870.62	PBDB + regression
<i>Palaeonictis peloria</i>	60916.03	PBDB + regression
<i>Palaeoryctes cruoris</i>	62.68	PBDB + regression
<i>Palaeoryctes puercensis</i>	24.97	PBDB + regression
<i>Palaeosyops laevidens</i>	1052429.44	PBDB + regression
<i>Palaeosyops paludosus</i>	1388030.44	PBDB + regression
<i>Palaeosyops robustus</i>	1619966.85	PBDB + regression
<i>Palenochtha minor</i>	9.87	PBDB + regression
<i>Palenochtha weissae</i>	9.87	PBDB + regression
<i>Paleotomus junior</i>	171.31	PBDB + regression
<i>Paleotomus radagasti</i>	1351.50	PBDB
<i>Panthera onca</i>	100000.00	PBDB
<i>Pantolambda bathmodon</i>	63252.15	PBDB
<i>Pantolambda cavirictus</i>	35087.78	PBDB + regression
<i>Pantolambda intermedius</i>	19134.73	PBDB + regression
<i>Parablastomeryx galushi</i>	19345.66	PBDB + regression
<i>Paracosoryx furlongi</i>	13493.99	[329]
<i>Paracosoryx wilsoni</i>	15473.78	PBDB

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Paracryptotis</i> <i>gidleyi</i>	28.22	[329]
<i>Paracryptotis</i> <i>rex</i>	42.10	[329]
<i>Paracynarctus</i> <i>kelloggi</i>	8349.86	[329]
<i>Paracynarctus</i> <i>sinclairi</i>	8022.46	[329]
<i>Paradaphoenus</i> <i>cuspigerus</i>	4023.87	[329]
<i>Paradaphoenus</i> <i>minimus</i>	3314.52	PBDB + regression
<i>Paradaphoenus</i> <i>tooheyi</i>	3498.19	[329]
<i>Paradjidaumo</i> <i>alberti</i>	15.90	PBDB + regression
<i>Paradjidaumo</i> <i>hypodus</i>	29.87	PBDB + regression
<i>Paradjidaumo</i> <i>reynoldsi</i>	20.14	PBDB + regression
<i>Paradjidaumo</i> <i>spokanensis</i>	54.13	PBDB + regression
<i>Paradjidaumo</i> <i>trilophus</i>	32.56	PBDB + regression
<i>Paradjidaumo</i> <i>validus</i>	61.14	PBDB + regression
<i>Paradomnina</i> <i>relictus</i>	23.81	[329]
<i>Paraenhydrocyon</i> <i>josephi</i>	7942.63	[329]
<i>Paraenhydrocyon</i> <i>wallowianus</i>	14185.85	[329]
<i>Parahippus</i> <i>leonensis</i>	94845.07	[329]
<i>Parahippus</i> <i>pawniensis</i>	99707.88	[329]
<i>Parahippus</i> <i>tyleri</i>	105994.41	PBDB + regression
<i>Parahippus</i> <i>wyomingensis</i>	98400.00	[188]
<i>Parahyus</i> <i>vagus</i>	185903.42	PBDB + regression
<i>Paralabis</i> <i>cedrensis</i>	75771.66	PBDB + regression
<i>Parallomys</i> <i>americanus</i>	162.39	[329]
<i>Paramiolabis</i> <i>taylori</i>	131926.47	[329]
<i>Paramylodon</i> <i>harlani</i>	1185306.75	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
Paramys adamus	50.76	PBDB + regression
Paramys atavus	47.48	PBDB + regression
Paramys compressidens	1746.66	PBDB + regression
Paramys copei	540.72	PBDB + regression
Paramys delicatior	708.27	PBDB + regression
Paramys delicatus	1793.22	PBDB + regression
Paramys excavatus	259.62	PBDB + regression
Paramys taurus	316.00	PBDB + regression
Paranamatomys storeri	7.59	PBDB
Parapliohippus carizzoensis	80821.64	[329]
Parapliosaccomys oregonensis	24.29	[329]
Parapliosaccomys transversus	45.58	PBDB
Parapotos tedfordi	3533.34	[329]
Pararyctes pattersoni	17.36	PBDB + regression
Paratomarctus euthos	14472.42	[329]
Paratomarctus temerarius	11498.82	[329]
Paratylopus labiatus	47357.63	PBDB + regression
Paratylopus primaevus	29814.41	PBDB + regression
Parectypodus clemensi	23.70	PBDB + regression
Parectypodus corystes	40.09	PBDB + regression
Parectypodus laytoni	11.30	PBDB + regression
Parectypodus lunatus	21.68	PBDB + regression
Parectypodus simpsoni	37.51	PBDB + regression
Parectypodus sinclairi	19.00	[362]
Parectypodus sylviae	12.82	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Parectypodus trovessartianus</i>	87.78	PBDB + regression
<i>Pareumys boskeyi</i>	88.57	PBDB + regression
<i>Pareumys grangeri</i>	57.59	PBDB + regression
<i>Pareumys guensburgi</i>	140.29	PBDB + regression
<i>Pareumys milleri</i>	76.28	PBDB + regression
<i>Parictis parvus</i>	1594.41	PBDB + regression
<i>Parictis personi</i>	2218.53	PBDB + regression
<i>Paromomys depressidens</i>	57.16	PBDB + regression
<i>Paromomys maturus</i>	197.38	PBDB + regression
<i>Paronychomys alticuspis</i>	17.12	[329]
<i>Paronychomys lemredfieldi</i>	20.09	[329]
<i>Paronychomys tuttlei</i>	38.86	[329]
<i>Paroreodon parvus</i>	51150.00	[198]
<i>Parvericius montanus</i>	41.26	[329]
<i>Parvericius voorhiesi</i>	28.79	[329]
<i>Patriofelis ferox</i>	22418.00	[198]
<i>Patriofelis ulta</i>	48311.29	PBDB + regression
<i>Patriolestes novaceki</i>	246.37	PBDB + regression
<i>Pauromys exallos</i>	17.97	PBDB + regression
<i>Pauromys lillegraveni</i>	12.11	PBDB + regression
<i>Pauromys simplex</i>	13.95	PBDB + regression
<i>Pauromys texensis</i>	17.97	PBDB + regression
<i>Pediomeryx hemphillensis</i>	167711.41	[329]
<i>Pelycodus jarrovii</i>	3222.75	PBDB + regression
<i>Pelycomys brulanus</i>	214.27	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Pelycomys rugosus</i>	330.84	PBDB + regression
<i>Penetrigonias hudsoni</i>	601751.47	PBDB
<i>Pentacemylus leotensis</i>	5601.22	PBDB + regression
<i>Pentacemylus progressus</i>	7131.24	PBDB + regression
<i>Pentacodon inversus</i>	604.36	PBDB + regression
<i>Pentacodon occultus</i>	1560.69	PBDB + regression
<i>Peraceras hessei</i>	936589.16	[329]
<i>Peraceras profectum</i>	2326789.55	[329]
<i>Peraceras superciliosum</i>	1639660.88	[329]
<i>Peradectes californicus</i>	6.11	PBDB + regression
<i>Peradectes chesteri</i>	5.04	PBDB + regression
<i>Peradectes elegans</i>	11.30	PBDB + regression
<i>Peradectes minor</i>	11.23	[327]
<i>Peradectes protinnominatus</i>	12.82	PBDB + regression
<i>Peratherium comstocki</i>	87.78	PBDB + regression
<i>Peratherium marsupium</i>	91.55	PBDB + regression
<i>Perchoerus probus</i>	24602.41	PBDB + regression
<i>Peridiomys halis</i>	63.43	[329]
<i>Peridiomys oregonensis</i>	35.35	PBDB + regression
<i>Peridiomys rusticus</i>	69.41	[329]
<i>Peritychus carinidens</i>	30641.27	PBDB + regression
<i>Peritychus coarctatus</i>	22723.44	PBDB + regression
<i>Perognathus ancenensis</i>	11.47	[329]
<i>Perognathus coquorum</i>	35.35	PBDB + regression
<i>Perognathus dunklei</i>	7.92	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Perognathus furlongi</i>	9.68	[329]
<i>Perognathus gidleyi</i>	11.82	[329]
<i>Perognathus maldei</i>	10.39	PBDB + regression
<i>Perognathus mclaughlini</i>	8.33	[329]
<i>Perognathus minutus</i>	5.93	[329]
<i>Perognathus pearlettensis</i>	6.89	[329]
<i>Perognathus rexroadensis</i>	17.97	PBDB + regression
<i>Perognathus projectioansrum</i>	3.63	[329]
<i>Peromyscus antiquus</i>	26.84	[329]
<i>Peromyscus brachygnathus</i>	10.39	PBDB + regression
<i>Peromyscus complexus</i>	25.46	PBDB
<i>Peromyscus cragini</i>	15.49	[329]
<i>Peromyscus dentalis</i>	12.11	PBDB + regression
<i>Peromyscus hagermanensis</i>	19.89	[329]
<i>Peromyscus minimus</i>	4.67	PBDB + regression
<i>Peromyscus nosher</i>	15.90	PBDB + regression
<i>Peromyscus polionotus</i>	14.30	PBDB
<i>Peromyscus sarmocophinus</i>	17.97	PBDB + regression
<i>Petauristodon jamesi</i>	307.97	[329]
<i>Petauristodon mathewsi</i>	214.86	[329]
<i>Petauristodon pattersoni</i>	336.97	[329]
<i>Pewelagus dawsonae</i>	447.84	[150]
<i>Pewelagus mexicanus</i>	186.75	PBDB + regression
<i>Phelosaccomys annae</i>	26.31	[329]
<i>Phelosaccomys hibbardi</i>	46.53	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Phelosaccomys neomexicanus</i>	19.89	[329]
<i>Phelosaccomys shotwelli</i>	20.49	[329]
<i>Phenacocoelus typus</i>	52052.08	[329]
<i>Phenacodaptes sabulosus</i>	109.55	PBDB + regression
<i>Phenacodus bisonensis</i>	17047.49	PBDB + regression
<i>Phenacodus grangeri</i>	28510.66	PBDB + regression
<i>Phenacodus intermedius</i>	45430.80	PBDB + regression
<i>Phenacodus magnus</i>	72471.82	PBDB + regression
<i>Phenacodus matthewi</i>	5446.34	[50]
<i>Phenacodus trilobatus</i>	64494.29	PBDB + regression
<i>Phenacodus vortmani</i>	10636.06	PBDB + regression
<i>Phenacolemur fortior</i>	186.75	[364]
<i>Phenacolemur mcgrewi</i>	63.45	PBDB + regression
<i>Phenacolemur praecox</i>	151.65	PBDB + regression
<i>Phenacolemur simonsi</i>	40.09	PBDB + regression
<i>Phenacomys gryci</i>	228.00	[198]
<i>Philotrox condoni</i>	11968.10	[329]
<i>Phlaocyon achoros</i>	2951.30	[329]
<i>Phlaocyon annectens</i>	3498.19	[329]
<i>Phlaocyon latidens</i>	2779.43	[329]
<i>Phlaocyon leucosteus</i>	3827.63	[329]
<i>Phlaocyon minor</i>	3498.19	[329]
<i>Phlaocyon taylori</i>	1939.14	[329]
<i>Phlaocyon yatkolai</i>	9604.62	[329]
<i>Phoberocyon johnhenryi</i>	179871.86	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Picroidus calgariensis</i>	12.82	PBDB + regression
<i>Picroidus canpaci</i> s	48.28	PBDB + regression
<i>Picroidus silberlingi</i>	48.28	PBDB + regression
<i>Pipestoneomys bisulcatus</i>	47.48	PBDB + regression
<i>Plagioctenodon krausae</i>	4.73	PBDB + regression
<i>Plagioctenodon rosei</i>	10.87	PBDB + regression
<i>Plagiomene accola</i>	359.33	PBDB + regression
<i>Plagiomene multicuspis</i>	850.13	PBDB + regression
<i>Planisorex dixonensis</i>	12.08	PBDB + regression
<i>Platygonus bicalcaratus</i>	114741.07	PBDB + regression
<i>Platygonus oregonensis</i>	40134.84	[329]
<i>Platygonus pearcei</i>	55826.28	[329]
<i>Platygonus vetus</i>	65463.62	[30]
<i>Plesiadapis anceps</i>	291.90	PBDB + regression
<i>Plesiadapis churchilli</i>	871.09	PBDB + regression
<i>Plesiadapis cookei</i>	4434.83	PBDB + regression
<i>Plesiadapis dubius</i>	417.80	PBDB + regression
<i>Plesiadapis fordinatus</i>	709.89	PBDB + regression
<i>Plesiadapis gingerichi</i>	2726.96	PBDB + regression
<i>Plesiadapis praecursor</i>	192.03	PBDB + regression
<i>Plesiadapis rex</i>	690.80	PBDB + regression
<i>Plesiocolopirus hancocki</i>	42441.52	PBDB + regression
<i>Plesiogulo lindsayi</i>	4628.55	[329]
<i>Plesiogulo marshalli</i>	3133.79	[329]
<i>Plesiolestes nacimenti</i>	233.00	[312]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Plesiolestes problematicus</i>	115.64	PBDB + regression
<i>Plesiolestes wilsoni</i>	366.42	PBDB + regression
<i>Plesiosminthus clivosus</i>	9.30	[329]
<i>Plesiosorex coloradensis</i>	192.48	[329]
<i>Plesiosorex donroosai</i>	685.40	[329]
<i>Pleurolicus dakotensis</i>	60.95	[329]
<i>Pleurolicus exiguum</i>	24.79	PBDB + regression
<i>Pleurolicus sellardsi</i>	53.28	[370]
<i>Pleurolicus sulcifrons</i>	83.93	[329]
<i>Pliocyon medius</i>	172818.99	[329]
<i>Pliocyon robustus</i>	176310.16	[329]
<i>Pliogale furlongi</i>	456.36	[364]
<i>Pliogale manka</i>	2986.47	PBDB + regression
<i>Pliogeomys parvus</i>	10.28	[329]
<i>Pliogeomys russelli</i>	15.18	[329]
<i>Pliohippus fossilatus</i>	257815.63	[329]
<i>Pliohippus pernix</i>	198789.15	[329]
<i>Pliohippus tehonensis</i>	164875.58	PBDB + regression
<i>Pliometanastes galushai</i>	8636.25	PBDB
<i>Pliometanastes protistus</i>	88139.12	PBDB + regression
<i>Plionarctos edensis</i>	56954.05	[329]
<i>Plionarctos harroldorum</i>	47351.79	PBDB + regression
<i>Plionictis ogygia</i>	36.97	[329]
<i>Pliophenacomys dixonensis</i>	50.76	PBDB + regression
<i>Pliophenacomys finneyi</i>	64.79	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Pliophenacomys meadensis</i>	54.13	PBDB + regression
<i>Pliophenacomys osborni</i>	86.49	[329]
<i>Pliophenacomys primaevus</i>	61.56	[329]
<i>Pliosaccommys dubius</i>	27.66	[329]
<i>Pliosaccommys higginsensis</i>	18.17	[329]
<i>Pliotaxidea garberi</i>	4440.12	PBDB + regression
<i>Pliotaxidea nevadensis</i>	130.32	[329]
<i>Pliotomodon primitivus</i>	107.77	[329]
<i>Pliozapus solus</i>	18.73	[329]
<i>Plithocyon ursinus</i>	189094.09	[329]
<i>Poabromylus golzi</i>	11885.48	PBDB + regression
<i>Poabromylus kayi</i>	23487.74	PBDB + regression
<i>Poebrotherium eximium</i>	50423.02	PBDB + regression
<i>Poebrotherium wilsoni</i>	60615.67	PBDB + regression
<i>Pogonodon eileenae</i>	138453.42	[92]
<i>Pratifelis martini</i>	215345.72	[329]
<i>Pratilepus kansasensis</i>	972.63	[329]
<i>Premnoides douglassi</i>	40.09	PBDB + regression
<i>Presbytymys lophatus</i>	72.36	PBDB + regression
<i>Presbytherium rhodorugatus</i>	2901.91	PBDB + regression
<i>Princetonia yalensis</i>	1703.86	PBDB + regression
<i>Probassariscus matthewi</i>	2396.46	PBDB + regression
<i>Probathyopsis harrisorum</i>	96406.85	PBDB + regression
<i>Probathyopsis praecursor</i>	119830.36	PBDB + regression
<i>Problastomyx primus</i>	14913.17	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Procamelus grandis</i>	400312.19	[329]
<i>Procamelus occidentalis</i>	189094.09	[329]
<i>Procerberus formicarum</i>	67.38	PBDB + regression
<i>Prochetodon cavus</i>	213.83	PBDB + regression
<i>Prochetodon foxi</i>	202.80	PBDB + regression
<i>Prochetodon speirsae</i>	417.88	PBDB
<i>Prochetodon taxus</i>	480.23	PBDB + regression
<i>Procranioceras skinneri</i>	169396.94	[329]
<i>Procynodictis progressus</i>	3665.64	PBDB + regression
<i>Procyon lotor</i>	5814.00	PBDB
<i>Procyon rexroadensis</i>	9632.47	PBDB + regression
<i>Prodiacodon concordiarcensis</i>	43.19	PBDB + regression
<i>Prodiacodon crustulum</i>	125.81	[168]
<i>Prodiacodon furor</i>	66.91	[280]
<i>Prodiacodon puercensis</i>	225.49	[200]
<i>Prodiacodon tauricinerei</i>	82.27	PBDB + regression
<i>Prodipodomys centralis</i>	21.26	[23]
<i>Prodipodomys idahoensis</i>	22.42	[329]
<i>Prodipodomys kansensis</i>	12.81	[329]
<i>Prodipodomys timoteoensis</i>	20.14	PBDB + regression
<i>Prohesperocyon wilsoni</i>	3787.93	PBDB + regression
<i>Proheteromys fedti</i>	17.97	PBDB + regression
<i>Proheteromys floridanus</i>	5.37	[329]
<i>Proheteromys gremmelsi</i>	27.97	[23]
<i>Proheteromys ironcloudi</i>	10.59	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Proheteromys maximus</i>	87.36	[329]
<i>Proheteromys nebraskensis</i>	21.54	PBDB
<i>Proheteromys sulculus</i>	15.90	PBDB + regression
<i>Proheteromys toledoensis</i>	59.15	[329]
<i>Prolapsus junctionis</i>	72.36	PBDB + regression
<i>Prolapsus sibilatoris</i>	178.45	PBDB + regression
<i>Prolimnocyon antiquus</i>	1442.37	PBDB + regression
<i>Prolimnocyon atavus</i>	2467.80	PBDB + regression
<i>Prolimnocyon haematus</i>	1058.55	PBDB + regression
<i>Promartes darbyi</i>	3029.14	[60]
<i>Promartes gemmarosae</i>	1945.16	[60]
<i>Promartes lepidus</i>	6776.78	[207]
<i>Promioclaenus acolytus</i>	410.27	PBDB + regression
<i>Promioclaenus pipiringosi</i>	748.79	PBDB + regression
<i>Promioclaenus thnetus</i>	260.57	[98]
<i>Promylagaulus riggsi</i>	85.63	[329]
<i>Pronodens silberlingi</i>	22247.84	[329]
<i>Pronothodectes gaoi</i>	254.50	PBDB + regression
<i>Pronothodectes jepi</i>	242.55	PBDB + regression
<i>Pronothodectes matthewi</i>	142.24	PBDB + regression
<i>Pronotolagus apachensis</i>	445.86	[329]
<i>Pronotolagus nevadensis</i>	60.34	[329]
<i>Pronotolagus whitei</i>	1436.55	[329]
<i>Proscalops miocaenus</i>	28.30	PBDB + regression
<i>Proscalops secundus</i>	72.97	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Proscalops tertius</i>	96.54	[329]
<i>Prosciurus magnus</i>	106.19	PBDB + regression
<i>Prosciurus parvus</i>	68.53	PBDB + regression
<i>Prosciurus relictus</i>	64.79	PBDB + regression
<i>Prosigmodon chihuahuensis</i>	72.36	PBDB + regression
<i>Prosigmodon ferrusquiai</i>	48.33	PBDB
<i>Prosigmodon holocuspis</i>	113.30	[329]
<i>Prosigmodon oroscoi</i>	27.28	PBDB + regression
<i>Prosomys mimus</i>	32.56	PBDB + regression
<i>Prosthennops niobrarensis</i>	43044.94	[329]
<i>Prosthennops serus</i>	53637.30	[329]
<i>Prosthennops xiphodonticus</i>	23860.99	[329]
<i>Prosynthetoceras francisi</i>	134591.56	[329]
<i>Prosynthetoceras orthrionanus</i>	40134.84	[329]
<i>Protadjidaumo pauli</i>	17.97	PBDB + regression
<i>Protadjidaumo typus</i>	12.11	PBDB + regression
<i>Protapirus obliquidens</i>	168000.80	[264]
<i>Protapirus simplex</i>	189283.62	PBDB + regression
<i>Protepycyon raki</i>	23623.56	[329]
<i>Proterix bicuspis</i>	194.51	[48]
<i>Proterix loomisi</i>	523.22	PBDB + regression
<i>Proterixoides davisi</i>	242.45	PBDB + regression
<i>Prothyptacodon albertensis</i>	464.26	PBDB + regression
<i>Prothyptacodon furens</i>	1351.03	PBDB + regression
<i>Prothyptacodon hilli</i>	1954.40	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Protictis haydenianus</i>	2074.37	PBDB + regression
<i>Protictis microlestes</i>	722.34	PBDB
<i>Protictis minor</i>	1270.84	PBDB + regression
<i>Protictis paralus</i>	354.83	PBDB + regression
<i>Protictis paulus</i>	298.35	PBDB
<i>Protictis simpsoni</i>	3202.65	PBDB + regression
<i>Protitanops curryi</i>	142477.00	[198]
<i>Protitanotherium superbum</i>	2858807.95	PBDB + regression
<i>Protoceras celer</i>	105068.04	PBDB + regression
<i>Protoceras skinneri</i>	81023.96	PBDB
<i>Protohippus gidleyi</i>	164390.50	[329]
<i>Protohippus perditus</i>	135944.23	[329]
<i>Protohippus supremus</i>	167711.41	[329]
<i>Protohippus vetus</i>	123329.89	PBDB
<i>Protolabis coartatus</i>	110194.25	[329]
<i>Protolabis heterodontus</i>	219695.99	[329]
<i>Protomarcus optatus</i>	11271.13	[329]
<i>Protoreodon pacificus</i>	34181.30	PBDB + regression
<i>Protoreodon parvus</i>	33051.65	PBDB + regression
<i>Protoreodon pearcei</i>	38169.88	PBDB + regression
<i>Protoreodon petersoni</i>	18828.04	PBDB + regression
<i>Protoreodon pumilus</i>	51870.94	PBDB + regression
<i>Protoreodon walshi</i>	59926.66	PBDB + regression
<i>Protorohippus venticulus</i>	14768.64	PBDB + regression
<i>Protosciurus mengi</i>	401.25	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Protosciurus tecuyensis</i>	478.19	[329]
<i>Protoselene griphus</i>	1619.16	PBDB + regression
<i>Protoselene opisthacus</i>	1703.86	PBDB + regression
<i>Protospermophilus kelloggi</i>	202.35	[329]
<i>Protospermophilus malheurensis</i>	103.54	[329]
<i>Protospermophilus oregonensis</i>	450.34	[329]
<i>Protospermophilus quatalensis</i>	273.14	[329]
<i>Protospermophilus vortmani</i>	230.44	[329]
<i>Prototomus deimos</i>	504.65	PBDB + regression
<i>Prototomus martis</i>	3573.98	PBDB + regression
<i>Prototomus phobos</i>	2691.73	PBDB + regression
<i>Prototomus robustus</i>	2434.13	PBDB + regression
<i>Prototomus secundarius</i>	1522.82	PBDB + regression
<i>Protungulatum donnae</i>	546.57	PBDB + regression
<i>Protylopus annexens</i>	14816.10	PBDB + regression
<i>Protylopus pearsonensis</i>	38366.65	PBDB + regression
<i>Protylopus petersoni</i>	18468.57	PBDB + regression
<i>Protylopus robustus</i>	17655.81	PBDB + regression
<i>Protylopus stocki</i>	15437.78	PBDB + regression
<i>Proviverroides piercei</i>	7343.84	PBDB + regression
<i>Psalidocyon marianae</i>	8777.97	[329]
<i>Pseudaelurus aeluroides</i>	31571.18	[329]
<i>Pseudaelurus intrepidus</i>	40945.61	[329]
<i>Pseudaelurus marshi</i>	33189.87	[329]
<i>Pseudaelurus stouti</i>	5767.53	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Pseudhipparion curtivallum</i>	58104.59	[329]
<i>Pseudhipparion gratum</i>	108012.26	[329]
<i>Pseudhipparion hessei</i>	75357.60	[329]
<i>Pseudhipparion retrusum</i>	86681.87	[329]
<i>Pseudhipparion simpsoni</i>	48050.12	[329]
<i>Pseudhipparion skinneri</i>	54176.36	[329]
<i>Pseudoblastomeryx advena</i>	10097.06	[329]
<i>Pseudoceras skinneri</i>	10938.02	[329]
<i>Pseudocylindrodon lateriviae</i>	101.65	PBDB + regression
<i>Pseudocylindrodon medius</i>	44.30	PBDB + regression
<i>Pseudocylindrodon neglectus</i>	88.57	PBDB + regression
<i>Pseudocylindrodon pintoensis</i>	194.89	MIOMAP
<i>Pseudodiplacodon progressum</i>	1263102.01	PBDB
<i>Pseudolabis dakotensis</i>	59874.14	[329]
<i>Pseudoparablastomeryx francescita</i>	4900.87	PBDB
<i>Pseudoparablastomeryx scotti</i>	5884.05	[329]
<i>Pseudoprotoceras longinaris</i>	35005.16	PBDB + regression
<i>Pseudoprotoceras minor</i>	19973.07	PBDB + regression
<i>Pseudotheridomys cuyamensis</i>	12.43	[329]
<i>Pseudotheridomys hesperus</i>	14.15	[329]
<i>Pseudotheridomys pagei</i>	8.17	[329]
<i>Pseudotomus californicus</i>	3660.37	PBDB + regression
<i>Pseudotomus eugenei</i>	14363.43	PBDB + regression
<i>Pseudotomus hians</i>	2299.39	PBDB
<i>Pseudotomus horribilis</i>	2971.99	[39]

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Pseudotomus johanniculi</i>	12241.50	PBDB + regression
<i>Pseudotomus littoralis</i>	2500.02	PBDB + regression
<i>Pseudotomus petersoni</i>	4209.29	[39]
<i>Pseudotomus robustus</i>	4052.00	PBDB + regression
<i>Pseudotrimylus mawbyi</i>	142.92	PBDB + regression
<i>Ptilodus fractus</i>	105.00	[362]
<i>Ptilodus gnomus</i>	87.78	PBDB + regression
<i>Ptilodus kummae</i>	208.28	PBDB + regression
<i>Ptilodus mediaevus</i>	311.46	PBDB + regression
<i>Ptilodus montanus</i>	373.57	PBDB + regression
<i>Ptilodus wyomingensis</i>	147.00	[362]
<i>Puercolestes simpsoni</i>	138.73	PBDB + regression
<i>Puma concolor</i>	48009.00	PBDB
<i>Puma lacustris</i>	17724.85	PBDB + regression
<i>Pyrocyon dioctetus</i>	1834.56	PBDB + regression
<i>Quadratomus grandis</i>	1595.31	PBDB + regression
<i>Quadratomus grossus</i>	4624.34	PBDB + regression
<i>Quadrodens wilsoni</i>	46.06	[329]
<i>Rakomeryx sinclairi</i>	111301.72	[329]
<i>Rapamys fricki</i>	1050.61	PBDB + regression
<i>Raphictis gausion</i>	261.58	PBDB + regression
<i>Reithrodontomys galushai</i>	10.39	PBDB + regression
<i>Reithrodontomys rexroadensis</i>	7.29	PBDB + regression
<i>Reithrodontomys wetmorei</i>	9.03	[329]
<i>Reithroparamys debequensis</i>	190.07	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Reithroparamys delicatissimus</i>	460.05	PBDB + regression
<i>Reithroparamys huerfanensis</i>	338.37	PBDB + regression
<i>Reithroparamys sciuroides</i>	482.77	[353]
<i>Repomys arizonensis</i>	54.13	PBDB + regression
<i>Repomys gustelyi</i>	81.45	[329]
<i>Repomys maxumi</i>	122.73	[329]
<i>Repomys panacaensis</i>	38.47	[329]
<i>Rhizocyon oregonensis</i>	3361.02	[329]
<i>Rhynchotherium falconeri</i>	54956347.86	PBDB + regression
<i>Russellagus vonhofi</i>	214.86	[329]
<i>Sanctimus falkenbachi</i>	151.41	[329]
<i>Sanctimus stouti</i>	120.30	[329]
<i>Sanctimus stuartae</i>	100.48	[329]
<i>Satherium piscinarium</i>	934.49	[329]
<i>Saxonella naylori</i>	45.48	PBDB + regression
<i>Scalopoides isodens</i>	32.14	[329]
<i>Scalopoides ripafodiator</i>	27.39	[329]
<i>Scalopus aquaticus</i>	39.60	PBDB
<i>Scapanoscapter simplicidens</i>	58.56	[329]
<i>Scapanus hagermanensis</i>	7.00	[198]
<i>Scapanus latimanus</i>	55.00	PBDB
<i>Scapanus proceridens</i>	65.37	[329]
<i>Scapanus shultzii</i>	97.00	[198]
<i>Scapanus townsendii</i>	141.58	[305]
<i>Scaphohippus sumani</i>	103960.05	PBDB

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Scenopagus curtidens</i>	26.62	PBDB + regression
<i>Scenopagus edenensis</i>	72.22	PBDB + regression
<i>Scenopagus priscus</i>	17.36	PBDB + regression
<i>Schaubeumys galbreathi</i>	19.54	[179]
<i>Schaubeumys grangeri</i>	20.14	PBDB + regression
<i>Schaubeumys sabrae</i>	17.54	[193]
<i>Schizodontomys amnicolus</i>	111.05	[329]
<i>Schizodontomys greeni</i>	94.00	[198]
<i>Schizodontomys harkseni</i>	105.64	[329]
<i>Sciuravus bridgeri</i>	41.22	PBDB + regression
<i>Sciuravus nitidus</i>	167.14	PBDB + regression
<i>Sciuravus popi</i>	338.37	PBDB + regression
<i>Sciuravus powayensis</i>	88.57	PBDB + regression
<i>Sciuravus wilsoni</i>	84.39	PBDB + regression
<i>Sciurion campestre</i>	27.28	PBDB + regression
<i>Sciurus carolinensis</i>	518.00	PBDB
<i>Sciurus olsoni</i>	22.73	PBDB
<i>Scottimus exiguus</i>	68.99	[215]
<i>Scottimus longiquus</i>	111.05	[329]
<i>Scottimus lophatus</i>	132.82	PBDB
<i>Scottimus vidiuus</i>	44.30	PBDB + regression
<i>Selenaletes scopaeus</i>	8232.01	PBDB + regression
<i>Serbelodon barboureensis</i>	11462314.59	[281]
<i>Sespedectes singularis</i>	28.30	PBDB + regression
<i>Sespedectes stocki</i>	23.37	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Sespemys thurstoni</i>	290.03	[329]
<i>Sesphia californica</i>	3604.72	[329]
<i>Sesphia nitida</i>	2394.82	[339]
<i>Shoshonius bowni</i>	142.24	PBDB + regression
<i>Shoshonius cooperi</i>	57.16	PBDB + regression
<i>Sifrhippus aemulor</i>	6498.52	[219]
<i>Sifrhippus grangeri</i>	11578.47	PBDB + regression
<i>Sifrhippus sandrae</i>	10631.54	PBDB + regression
<i>Sigmodon curtisi</i>	92.85	PBDB + regression
<i>Sigmodon hudspethensis</i>	68.53	PBDB + regression
<i>Sigmodon minor</i>	52.98	[329]
<i>Simidectes magnus</i>	3787.93	PBDB + regression
<i>Simidectes medius</i>	2882.11	PBDB + regression
<i>Simidectes merriami</i>	6833.74	PBDB + regression
<i>Similisciurus maxwelli</i>	259.82	[329]
<i>Simimeryx hudsoni</i>	4171.22	PBDB + regression
<i>Simimeryx minutus</i>	1736.45	PBDB + regression
<i>Simimys landeri</i>	38.23	PBDB + regression
<i>Simimys simplex</i>	13.95	PBDB + regression
<i>Simocyon primigenius</i>	70000.00	NOW
<i>Simojovelhyus pocitosense</i>	19763.18	[355]
<i>Simpsonictis pegus</i>	245.14	PBDB + regression
<i>Simpsonictis tenuis</i>	126.87	PBDB + regression
<i>Simpsonlemur citatus</i>	171.00	[312]
<i>Simpsonlemur jepseni</i>	121.00	[312]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Simpsonodus chacensis</i>	1262.40	PBDB + regression
<i>Sinclairella dakotensis</i>	139.74	PBDB + regression
<i>Sinopa major</i>	7566.66	PBDB + regression
<i>Sinopa rapax</i>	3299.38	PBDB + regression
<i>Smilodectes gracilis</i>	1035.64	PBDB + regression
<i>Smilodectes mcgrewi</i>	1212.97	PBDB + regression
<i>Smilodectes sororius</i>	935.37	PBDB + regression
<i>Smilodon gracilis</i>	57206.56	PBDB + regression
<i>Sminthosinus bowleri</i>	100.48	[329]
<i>Sorex cinereus</i>	3.80	PBDB
<i>Sorex edwardsi</i>	6.84	PBDB
<i>Sorex hagermanensis</i>	7.56	PBDB + regression
<i>Sorex meltoni</i>	3.46	[329]
<i>Sorex palustris</i>	13.46	[305]
<i>Sorex powersi</i>	7.77	[329]
<i>Sorex rexroadensis</i>	8.00	[198]
<i>Sorex yatkolai</i>	4.91	PBDB
<i>Spermophilus argonautus</i>	131.63	[329]
<i>Spermophilus bensoni</i>	184.93	[329]
<i>Spermophilus boothii</i>	498.11	[281]
<i>Spermophilus cragini</i>	578.25	[329]
<i>Spermophilus dotti</i>	270.43	[329]
<i>Spermophilus fricki</i>	231.96	[293]
<i>Spermophilus gidleyi</i>	700.00	[198]
<i>Spermophilus howelli</i>	139.77	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Spermophilus jerae</i>	87.36	[329]
<i>Spermophilus matalchicensis</i>	220.52	PBDB + regression
<i>Spermophilus matthewi</i>	399.62	[22]
<i>Spermophilus meadensis</i>	100.48	[329]
<i>Spermophilus rexroadensis</i>	345.98	PBDB + regression
<i>Spermophilus russelli</i>	166.00	[198]
<i>Spermophilus shotwelli</i>	204.00	[198]
<i>Spermophilus tephrus</i>	89.12	[329]
<i>Spermophilus wellingtonensis</i>	295.89	[329]
<i>Spermophilus wilsoni</i>	247.15	[329]
<i>Sphacorhysis burntforkensis</i>	40.09	PBDB + regression
<i>Sphenophalos nevadanus</i>	8789.20	PBDB
<i>Spilogale microdens</i>	1270.84	[43]
<i>Spilogale putorius</i>	341.19	[305]
<i>Spilogale rexroadi</i>	895.71	PBDB + regression
<i>Stegomastodon mirificus</i>	35283150.07	PBDB + regression
<i>Steinius annectens</i>	397.75	[325]
<i>Steinius vespertinus</i>	119.90	PBDB + regression
<i>Stelocyon arctylus</i>	2132.95	PBDB + regression
<i>Stenoechinus tantalus</i>	48.42	[329]
<i>Stenomylus gracilis</i>	44801.64	[329]
<i>Stenomylus hitchcocki</i>	38948.67	[329]
<i>Stenomylus taylori</i>	47980.97	PBDB
<i>Sthenictis dolichops</i>	665.14	[329]
<i>Sthenictis junturensis</i>	330.30	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Stibarus montanus</i>	1152.46	PBDB + regression
<i>Stibarus obtusilobus</i>	1416.00	PBDB + regression
<i>Stibarus quadricuspis</i>	2093.31	PBDB + regression
<i>Stockia powayensis</i>	161.34	PBDB + regression
<i>Stratimus strobeli</i>	20.09	[329]
<i>Strigorhysis bridgerensis</i>	91.55	PBDB + regression
<i>Strigorhysis huerfanensis</i>	176.39	PBDB + regression
<i>Stygimys gratus</i>	128.62	PBDB + regression
<i>Stygimys jepseni</i>	84.00	[362]
<i>Stygimys kuszmauli</i>	291.90	PBDB + regression
<i>Stylinodon mirus</i>	47987.28	PBDB + regression
<i>Subdromomeryx antilopinus</i>	59874.14	[329]
<i>Subhyracodon mitis</i>	458341.99	[279]
<i>Subhyracodon occidentalis</i>	662391.44	PBDB + regression
<i>Sunkahetanka geringensis</i>	11158.98	[329]
<i>Swaindelphys cifellii</i>	24.57	PBDB
<i>Symmetrodontomys simplicidens</i>	26.84	[329]
<i>Syndyoceras cooki</i>	73865.41	[329]
<i>Tachylagus gawneae</i>	146.91	PBDB + regression
<i>Taeniolabis taoensis</i>	75621.27	PBDB + regression
<i>Talpavoides dartoni</i>	8.53	PBDB + regression
<i>Talpavus conjunctus</i>	45.48	PBDB + regression
<i>Talpavus duplus</i>	17.88	PBDB + regression
<i>Talpavus nitidus</i>	11.30	PBDB + regression
<i>Tamias ateles</i>	32.56	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Tanymycter brachydontus</i>	102744.44	[329]
<i>Tapiravus validus</i>	64860.88	[329]
<i>Tapirus simpsoni</i>	369534.73	[329]
<i>Tapochoerus egressus</i>	9671.40	PBDB + regression
<i>Tapochoerus mcmillini</i>	4357.69	PBDB + regression
<i>Tapocyon dawsonae</i>	17454.68	PBDB + regression
<i>Tapocyon robustus</i>	21673.71	PBDB + regression
<i>Tardontia nevadans</i>	157.59	[329]
<i>Tardontia occidentale</i>	184.22	PBDB + regression
<i>Tarka stylifera</i>	3858.47	PBDB + regression
<i>Tatmanius szalayi</i>	46.59	[71]
<i>Taxidea mexicana</i>	10434.28	[27]
<i>Taxidea taxus</i>	7112.14	[305]
<i>Tayassu protervus</i>	45119.24	PBDB
<i>Teilhardina americana</i>	66.71	PBDB + regression
<i>Teilhardina crassidens</i>	57.16	PBDB + regression
<i>Teleoceras meridianum</i>	2022813.66	[329]
<i>Teletaceras mortivallis</i>	89575.08	PBDB + regression
<i>Telmatherium altidens</i>	2087486.17	PBDB + regression
<i>Telmatherium cultridens</i>	627910.03	PBDB + regression
<i>Telmatherium manteoceras</i>	478340.42	PBDB + regression
<i>Temnocyon altigenis</i>	32532.67	[329]
<i>Temnocyon percussor</i>	68871.66	[329]
<i>Tenudomys bodei</i>	43.38	[329]
<i>Tenudomys macdonaldi</i>	79.84	[329]

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Tephrocyon rurestris</i>	13095.19	[329]
<i>Tetonius ambiguus</i>	92.52	PBDB
<i>Tetonius matthewi</i>	107.32	PBDB + regression
<i>Tetonius mckennai</i>	57.16	PBDB + regression
<i>Tetraclaenodon puerensis</i>	9425.94	PBDB + regression
<i>Tetrapassalus mckennai</i>	5.04	PBDB
<i>Texomys ritchiei</i>	97.51	[329]
<i>Thinobadistes segnus</i>	645890.00	[194]
<i>Thinocyon velox</i>	1924.12	PBDB + regression
<i>Thinohyus latus</i>	101400.00	[198]
<i>Thisbemys corrugatus</i>	965.53	PBDB + regression
<i>Thisbemys elachistos</i>	140.29	PBDB + regression
<i>Thisbemys perditus</i>	587.96	PBDB + regression
<i>Thisbemys uintensis</i>	1777.64	PBDB + regression
<i>Thomomys bottae</i>	114.82	[305]
<i>Thomomys carsonensis</i>	29.87	PBDB + regression
<i>Thomomys gidleyi</i>	37.34	[329]
<i>Thryptacodon antiquus</i>	4110.15	PBDB + regression
<i>Thryptacodon australis</i>	2744.65	PBDB + regression
<i>Thryptacodon orthogonius</i>	1536.41	PBDB + regression
<i>Thryptacodon pseudarctos</i>	6070.61	PBDB + regression
<i>Thylacaelurus campester</i>	35.01	PBDB + regression
<i>Thylacaelurus montanus</i>	63.45	PBDB + regression
<i>Thylacodon pusillus</i>	48.28	PBDB + regression
<i>Ticholeptus zygomaticus</i>	106937.52	[329]

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Tillodon fodiens</i>	49838.54	PBDB + regression
<i>Tillomys senex</i>	84.39	PBDB + regression
<i>Tinimomys graybulliensis</i>	11.30	PBDB + regression
<i>Tinimomys tribos</i>	5.32	[301]
<i>Titanoides gidleyi</i>	48022.06	PBDB + regression
<i>Titanoides nanus</i>	37406.72	PBDB + regression
<i>Titanoides primaevus</i>	72645.54	PBDB + regression
<i>Tomarctus brevirostris</i>	17500.77	[329]
<i>Tomarctus hippophaga</i>	13766.59	[329]
<i>Torrejonia sirokii</i>	440.75	PBDB + regression
<i>Toxotherium hunteri</i>	91951.16	PBDB + regression
<i>Tregosorex holmani</i>	26.84	[329]
<i>Tremarctos floridanus</i>	149968.48	[305]
<i>Trigenicus profectus</i>	28723.88	PBDB + regression
<i>Trigonias osborni</i>	599072.86	PBDB + regression
<i>Trigonias yoderensis</i>	264875.92	PBDB + regression
<i>Trigonictis cookii</i>	7397.01	PBDB + regression
<i>Trigonictis macrodon</i>	419.89	[329]
<i>Triisodon quivirensis</i>	22180.76	PBDB + regression
<i>Trilaccogaulus ovatus</i>	103.54	[329]
<i>Triplopus cubitalis</i>	34118.20	PBDB + regression
<i>Triplopus implicatus</i>	64570.33	PBDB + regression
<i>Triplopus obliquidens</i>	95152.03	PBDB + regression
<i>Triplopus rhinocerinus</i>	88700.44	PBDB
<i>Triplopus woodi</i>	69515.35	PBDB

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Tritemnodon agilis</i>	10780.00	[63]
<i>Tritemnodon strenuus</i>	4702.60	PBDB + regression
<i>Trogolemur amplior</i>	60.27	PBDB + regression
<i>Trogolemur myodes</i>	32.59	PBDB + regression
<i>Trogomys rupinimenthae</i>	12.55	[329]
<i>Trogosus castoridens</i>	25945.98	PBDB + regression
<i>Trogosus grangeri</i>	42730.29	PBDB + regression
<i>Trogosus latidens</i>	86402.00	PBDB + regression
<i>Tubulodon atopum</i>	186.75	PBDB + regression
<i>Tubulodon taylori</i>	113.11	PBDB
<i>Tuscahomys mediuss</i>	84.22	PBDB
<i>Tuscahomys minor</i>	44.37	PBDB
<i>Tylocephalonyx skinneri</i>	1696671.69	PBDB
<i>Uintaceras radinskyi</i>	516493.43	PBDB + regression
<i>Uintacyon asodes</i>	6096.02	PBDB + regression
<i>Uintacyon massetericus</i>	3203.72	PBDB + regression
<i>Uintacyon rudis</i>	1864.20	PBDB + regression
<i>Uintanius ameghini</i>	48.28	PBDB + regression
<i>Uintanius rutherfordi</i>	51.17	PBDB + regression
<i>Uintasorex montezumicus</i>	2.40	PBDB + regression
<i>Uintasorex parvulus</i>	6.11	PBDB + regression
<i>Uintatherium anceps</i>	523092.59	PBDB + regression
<i>Untermannerix copiosus</i>	121.51	[329]
<i>Unuchinia dysmathes</i>	114.44	PBDB
<i>Uriscus californicus</i>	130.13	PBDB + regression

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Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Urocyon cinereoargenteus</i>	3829.00	PBDB
<i>Ursavus brevirhinus</i>	80000.00	NOW
<i>Ursavus pawniensis</i>	61697.58	[329]
<i>Ursavus primaevus</i>	90000.00	NOW
<i>Ursus abstrusus</i>	43499.17	PBDB + regression
<i>Ursus americanus</i>	93431.00	PBDB
<i>Utahia carina</i>	17.88	PBDB + regression
<i>Utahia kayi</i>	37.51	PBDB + regression
<i>Valenia wilsoni</i>	279.17	PBDB + regression
<i>Vassacyon promicrodon</i>	5427.03	PBDB + regression
<i>Viverravus acutus</i>	483.44	PBDB + regression
<i>Viverravus gracilis</i>	849.04	PBDB + regression
<i>Viverravus laytoni</i>	241.14	PBDB + regression
<i>Viverravus lutosus</i>	602.46	PBDB + regression
<i>Viverravus minutus</i>	602.46	PBDB + regression
<i>Viverravus politus</i>	1009.45	PBDB + regression
<i>Viverravus rosei</i>	148.81	PBDB + regression
<i>Viverravus sicarius</i>	1594.41	PBDB + regression
<i>Vulpavus australis</i>	1746.94	PBDB + regression
<i>Vulpavus palustris</i>	2833.89	PBDB + regression
<i>Vulpavus profectus</i>	3388.44	[360]
<i>Vulpes stenognathus</i>	7331.97	[329]
<i>Vulpes velox</i>	2197.86	[305]
<i>Washakius insignis</i>	111.45	PBDB + regression
<i>Washakius izetti</i>	73.44	PBDB + regression

Continued on next page

Table A.1 – continued from previous page

Species	Mass (g)	Source
<i>Washakius woodringi</i>	48.28	PBDB + regression
<i>Wilsoneumys planidens</i>	51.70	PBDB
<i>Worlandia inusitata</i>	73.44	PBDB + regression
<i>Wyolestes apheles</i>	1594.41	PBDB + regression
<i>Wyolestes iglesius</i>	1288.90	PBDB + regression
<i>Wyonycteris chalix</i>	8.61	PBDB + regression
<i>Xenicohippus craspedotum</i>	10895.00	[198]
<i>Xenicohippus grangeri</i>	12284.22	PBDB + regression
<i>Ysengrinia americana</i>	110194.25	[329]
<i>Yumaceras figginsi</i>	293607.76	[329]
<i>Yumaceras hamiltoni</i>	247706.54	[329]
<i>Yumaceras ruminalis</i>	314896.72	[329]
<i>Zapus burti</i>	21.33	[329]
<i>Zapus rinker</i>	27.28	PBDB + regression
<i>Zapus sandersi</i>	18.92	[329]
<i>Zemiodontomys burkei</i>	120.30	PBDB + regression
<i>Zetamys nebraskensis</i>	60.34	[329]