# Evolutionary paleoecology and the biology of extinction

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#### Committee

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## Introduction and Theory

- 2 Evolutionary paleoecology is the study of the effects of ecological traits and factors on differential rate dynamics, particularly rates of faunal turnover and diversification [61].
- <sup>4</sup> Ecological traits are expressed by a taxon, at any level, and are involved in biotic-biotic or biotic-abiotic interactions while ecological factors are the environmental conditions in which
- 6 a taxon exists (the set of all biotic and abiotic interactors). Diversification is the difference between origination and extinction and is the net pattern of macroevolution. The study
- of evolutionary paleoecology is therefore the link between environmental (biotic-biotic and biotic-abiotic) interactions and macroevolution. As a corollary to Kitchell [61]'s definition,
- Allmon [3] states that in order to correctly link ecological interactions to macroevolution, one must focus on the specific traits and factors that may affect the speciation process. Tacitly included in this is the study of how ecological traits are related to extinction [60].

Geographic range size is considered one of the most obvious emergent species level traits [46] because no one property of an organism can explain this trait, is not expressed by a single organisms, and is the combination of multiple traits. Species with larger geographic ranges tend to have lower extinction rates than species with smaller geographic ranges [40, 45, 50, 75, 95]. The effects of traits expressed at the organismal level, such as those relating to environmental preference, together or in concert remain less well understood. Ecological traits have been shown to be related to differential extinction [11, 34, 65, 75], especially the relationship between adaptation to variable environments and increased species longevity. While some research has focused on the indirect effect of organismal traits on longevity [39], the additive or interactive effects of organismal traits alone and their relationship to survival remain understudied. Here I propose to study the how organismal traits are related to extinction and whether emergent traits are necessary for best explaining extinction.

- Survival can be considered the fundamental measure evolutionary success because ultimately, a successful lineage is not one that speciated greatly but one that never went extinct [25, 77].
- During periods of background extinction it is expected that extinction is non-random with respect to biology [45], so it should be possible to estimate how various ecological traits
- are correlated with survival [60, 61]. Periods of background extinction also represent the majority of geologic time, remain relatively predictable and change slowly, thus providing a
- better opportunity to study how traits are related to survival than periods of mass extinction [45, 89]. Additionally, the Law of Constant extinction [110] states that extinction rate within
- an adaptive zone is taxon-age independent. By analyzing the survival patterns within different adaptive zones during periods of background extinction, it should be possible to determine if
- extinction is best modeled as age independent or dependent.

It is under this framework that I propose to study how ecological traits associated with environmental preference and range size have affected differential survival and cosmopolitan-endemism dynamics. I will be studying two distantly related and biotically different groups:

40 Permian brachiopods and Cenozoic mammals. Both of these groups are considered to have

very good fossil records able to reflect massive long term evolutionary patterns [71]. These two time periods were chosen because they represent periods of approximately the same length (47 My and 65 My) and of climatic change, global warming and global cooling respectively. Also,

these two groups are a marine and terrestrial system respectively and the traits associated with environmental preference and range size (described below) are fundamentally very

46 different.

Importantly, this comparison allows for a series of questions to be asked with regards to emergent properties and the Law of Constant extinction. Brachiopod and mammal survival can be analyzed at the generic level for different adaptive zones (combinations of traits). The age independence or dependence of extinction for these two groups can then be tested. However because there is a known potential bias towards age-dependent extinction when analyzing generic level survival curves [87, 88, 101], the mammalian survival will be further analyzed at the specific level and the differences between the two survival functions will be examined, specifically in whether different traits best model the two curves and if either is age-independent or not. Additionally, the importance of global climatic change in modeling both brachiopod and mammalian survival will be analyzed. In the case of mammalian survival, if global climatic change is found to not be important this does not mean that regional or local climatic changes are not important to survival. By analyzing mammalian community connectedness, it should be possible to estimate how disjoint communities are and if it is reasonable to expect global, regional, or local processes to have dominated and how this may have changed over time.

## <sup>62</sup> Brachiopods, environmental preference, and extinction

Questions: In Australasian Permian brachiopods, are traits related to environmental preference correlated with differential survival? Are certain traits more explanatory of survival than others? Does changing climate, habitat or substrate availability affect survival?

Background and Predictions: Brachiopods are suspension feeders that are not actively motile, thus the environment conditions of where they occur are extremely important for estimating success. Brachiopod occurrence has been found to be strongly linked to the substrate on

which they occur [90, 91]. In contrast, there are no distinct hypotheses related to brachiopod

environmental preeference and dispersal ability or environmental energetics, both of which are related to substrate occurrence. Because brachiopods are suspension feeders, environmental

energetics are important for prey capture and individual survival. The availability of optimal environments becomes a limiting factor on the possible geographic extent of a taxon and

the expected extinction rate accompanying geographic range size. In brachiopods, three important traits involved in determining environmental preference are affixing strategy,

substrate preference, and habitat preference. While larval mode is considered important in determining range size in marine invertebrates [48, 49], this does not preserve in brachiopods

and thus cannot be used for modeling survival [49].

Substrate preference is related to the chemical and physical processes affecting an environment and may limit the range of possible environments in which an organism can optimally survive, thus limiting the possible geographic extent of a taxon. Substrate selection is mitigated via larval chemosensory abilities and thus may act as a weak proxy for larval dispersal ability [48, 49]. Affixing strategy and habitat preference are also related to range size by limiting the possible geographic extent of a taxon. Affixing strategy is the manner by with an individual interfaces with the substrate and different strategies are optimal for different environmental conditions such as flow speed or mud depth [2, 63, 64]. Habitat preference is a statement of the suitability of an environment and the accompanying energy level and avaliability might act to limit the possible geographic extent of a taxon. Additionally, habitat might capture other factors relating to the environment not captured in substrate preference. While other environmental factors such as temperature, water depth, and environmental energy have not been found to limit the distribution of modern brachiopods [90, 91] though it is unknown if these traits affect survival.

The three principle divisions of substrate affinity are carbonate, clastic, or mixed. These are
descriptions of the lithology of the sites at which a taxon is found [7, 33, 58, 74, 75]. The
Pharenozoic is characterized by an overall decline in carbonates relative to clastics [33, 74].
Because of this, it is expected that taxa with clasitic or mixed affinities will have greater
survival than taxa associated with carbonate substrates. If substrate affinity is not found
to be important for modeling survival, this may mean that depositional environment has
little to no affect on survival and that other factors relating to the environment, measured or
unmeasured, are dominant. Additionally, this may mean that depositional environments were
not a limiting factor to survival even though it is strongly related to geographic distribution.

The three ways of categorizing affixing strategy are pedunculate, reclining, and cementing. During the Permian, pedunculate taxa are associated with shallow on-shore environments while reclining taxa are associated with deep off-shore environments [20]. These associations are weak as most assemblages are composed of a heterogeneous mix of taxa [20]. Previous global level analysis of brachiopod durations showed that affixing strategy is correlated with longevity [2] and that among endemic taxa, reclining taxa had longer durations than other affixing strategies. Among cosmopolitan taxa, however, pedunculate and cementing taxa had longer durations than all other taxa, both cosmopolitan and endemic. This global analysis mixed taxa from many different geological periods and geographic regions which may produce unfair and biased comparisons. By restricting analysis to a single continuous region, I hope to alleviate these concerns and instead focus on survival of an evolving taxonomic assemblage in a continuous environmental context. If affixing strategy is not important for modeling survival this would mean that, even though affixing strategy is correlated with differential survival [2], it may only be a minor factor. For example, this may indicate that the environmental energetics of Australasia was rather uniform. 116

The primary ways of classifying habitat preference are on-shore, off-shore, or mixed. Habitat has been frequently invoked for explaining patterns of global diversity [15, 47, 51, 58, 97]. Importantly, habitat preference is related to sea-level and the availability of on-shore habitats

in particular might change dramatically and rapidly over time [76]. On-shore environments have declined in areal extent over the Pharenozoic [79], so it is expected that taxa associated with on-shore habitats would have lower durations than taxa associated with off-shore or mixed habitat preferences. If environmental preference is found to be unimportant when modeling survival, this may mean that sea-level dynamics are rather constant through out Australasia or that other factors may have dominated. The former should be apparent from the rock record itself, an observation that would help determine which processes and conditions were most likely affecting survival.

An important consideration is that taxonomic survival might not be linked to single environments per se, but the variability of environments [34, 41, 65]. Adaptation to environmental variability has been found to relate strongly with survival past origination [34]. It is then expected that taxa with mixed substrate affinities and habitat preferences would have longer durations than taxa with single preferences because this means that a taxon's potential geographic extent is not expressly limited by either of these two traits thus decreasing the expected extinction rate because of an expectedly large range size [40, 45, 50, 75, 95].

During the Permian there was a shift from an "ice house" to a "hot house" world [13, 29, 57, 80].

Australasian taxa are of particular interest because of their proximity to the south pole during the Permian and the repeated glacial activity in the region [13, 29, 57]. According to Olszewski and Erwin [76], however, sea-level and climate change do not wholly explain the brachiopod ecological dynamics experienced in the Permian of Texas. It is then predicted that climate will not be the best sole predictor of brachiopod survival, and that some combination of one or more the above organismal traits will be necessary to best model survival.

Proposed research: I propose a survival analysis approach to compare the patterns of survivorship in Permian brachiopods. Survival analysis is the analysis of time-till-event data.

In a paleontological context this is the time from origination (first appearance date; FAD) till extinction (last appearance date; LAD). I restrict this analysis to Australasia because it represents a relatively continually sampled and well worked area that preserves the majority of the entire Permian [8, 22, 23, 114]. Multiple models of survival with various combinations, both additive and interactive, of the traits described above will be compared. Substrate preference, affixing strategy, and habitat preference will be assumed to be constant for the duration of the taxon and modeled as time-independent covariates. If and how these traits may have evolved will remain for future study. Climate change will be modeled as either an ancillary Heaviside function or an ancillary time-dependent covariate.

Because substrate and habitat occurrences are not constant at the species level, it is necessary to determine the most probable assignments. The lithology of all occurrences will be classified into one of the three substrate affinity categories following Foote [33] while paleoenvironmental setting of all occurrences will be classified into one of the habitat preferences following Kiessling et al. [59]. Both of these traits will be assigned to all taxa following the Bayesian approach of Simpson and Harnik [102] where assignments are determined as the posterior probability of a taxon's occurrences in comparison to available options during the duration of said taxon.

Assignment probability,  $P(H_1|E)$ , will be calculated as

$$P(H_1|E) = \frac{P(E|H_1)P(H_1)}{P(E|H_1)P(H_1) + P(E|H_2)P(H_2)}$$
(1)

where  $P(H_1)$  and  $P(H_2)$  are the prior probabilities of assignment while  $P(E|H_1)$  and  $P(E|H_2)$  are conditional, binomial probabilities.

In order to asses if the best model of survival is age independent or dependent, the distribution of survival will be modeled using various probability distributions (i.e. exponential, Weibull, log-normal, etc.). An exponential distribution represents the Law of Constant extinction as the rate parameter ( $\lambda$ ) is constant with respect to time. In contrast, the Weibull distribution has two parameters, scale ( $\lambda$ ) and shape (k). The scale parameter analogous to the  $\lambda$  of the exponential distribution while the shape parameter describes whether  $\lambda$  is accelerating (k < 1), decelerating (k > 1), or constant (k = 1) with respect to taxon age.

Permian brachiopod occurrence information is available via the Paleobiology Database (PBDB; http://fossilworks.org) and is primarily based on the work of Clapham [20–24] and Waterhouse [114].

Preliminary results Preliminary analysis of brachiopod survivorship was restricted to genera that originated within the Permian. Survival was analyzed for taxa classified for substrate affinity and habitat preference. For substrate affinity, the priors of Eq. 1 were  $P(H_1) = P(H_2) = 0.5$  and if  $P(H_1|E) > \frac{2}{3}$  then the taxon was considered of carbonate affinity while if  $P(H_1|E) < \frac{1}{3}$  then the taxon was considered to have a clastic affinity. Otherwise, the taxon was considered to have mixed affinity. In the case of habitat affinity, the posterior probability for each habitat (inshore, offshore, none) was calculated using Eq. 1 with priors of  $P(H_1) = \frac{1}{3}$  and  $P(H_2) = \frac{2}{3}$  and the preference with maximum of the three posterior probabilities was the assignment.

Preliminary model fitting with both exponential and Weibull distributions and either or both traits indicated that the best model, based on comparison of AICc scores [1, 19, 44], was the model with substrate affinity as the sole predictor and a Weibull distribution of survival. This model is illustrated below (Fig. 1a). While this is the preliminarily best model of survival, the model with both substrate affinity and habitat preference as additive effects and a Weibull survival distribution may also be considered a good model of survival (ΔAICc ≈ 2.9). Additionally, as illustrated by the difference between the nonparametric Kaplan–Meier survival curves and the predictions of the parametric model of survival (Fig. 1a) and there is room for improvement in model specification.

The shape parameter (k) of the AICc best model (Fig. 1a) is estimated to be approximately 1.9. As described above, values of k greater than 1 indicate that failure (extinction) rate is accelerating with respect to taxon age, which may mean that the Law of Constant Extinction does not hold when modeling generic level extinction in brachiopods.

For brachiopod survival based on substrate affinity (Fig. 1a), survival was greater for both carbonate and clastic affinities and lowest for taxa with mixed affinity. Visual inspection

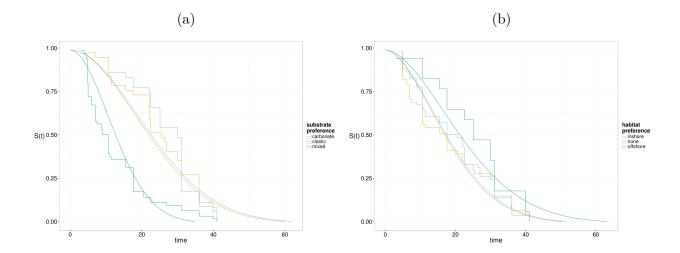


Figure 1: Survivorship curves of Australasian Permian brachiopod genera based on substrate affinity (a) and habitat preference (b). The stepwise functions are nonparametric Kaplan–Meier survival curves for each of the three substrate affinities. The three smooth lines are the predicted survivorship probabilities for taxon of the given age from parametric survival models.

of the estimated survival functions compared to the nonparametric Kaplan–Meier curves indicates that they were adequate fits to the data.

Brachiopod survival with the sole predictor being environmental and a Weibull distribution of survival was not a good model of survival, with an approximate  $\Delta \text{AICc}$  of 33 between this model and the AICc best model. There is a great degree of deviance between the nonparametric Kaplan–Meier curves and model predictions (Fig. 1b). Additionally, this model is not significantly different from the model with only an intercept ( $\chi^2 = 2.41$ , df = 2, p = 0.3). This means, preliminarily, that habitat preference alone makes no difference in generic level survival.

Further refinements to these models include modeling survival using other distributions of survival such as a log-normal distribution. Additionally the inclusion of affixing strategy and climate as predictors will increase the understanding of the biology underlying brachiopod generic survival. Additionally, the effects of sampling and uncertain durations will explored in simulations described below in the context of generic versus specific survival.

# Ecology and survival in Cenozoic mammals

Questions: How do ecological traits related to range size affect mammalian taxon duration? Is any single trait the best predictor of mammalian survivorship, or do multiple traits together best model taxon duration? Is climate an important factor in modeling mammalian taxon

#### duration?

Background and Predictions: Three potential constituent traits of range size in mammals are dietary category, locomotor category, and body size [26, 27, 55, 67, 68, 105, 106]. These traits describe different aspects of how an organism interacts with the environment based on either prey availability, dispersal ability, or energetic cost. Dietary categories are defined here as broad trophic categories that subsume further, more specific, classifications. The categories used here are carnivore, herbivore, omnivore, and insectivore. It should be noted that most prior analyses have not included insectivore as a category [55, 81]. Similarly, locomotor categories are defined as combinations of more specific classifications and are broken into arboreal, ground dwelling, and scansorial. Both of these traits are constant at the specific level.

Dietary categories limit abundance because of the availability of resources in a location [18, 26, 53, 100, 112]. Abundance is correlated with occupancy, or the number of localities at which a taxon is found [17, 35, 54]. It follows then that the limits on abundance imposed by resource availability would affect the possible range size of a taxon. Because dietary categories are fundamentally linked to primary productivity and trophic hierarchy, it is expected that the most stable categories would have the greatest survival and the least stable would have the lowest survival. Stability is used here to mean number of "steps" from primary productivity and the related relative requirement of other biotic interactors. As such, herbivores are expected to have greatest survival and carnivores the lowest survival. Omnivorous taxa are expected to have average taxon durations compared to the other two categories. Mammalian herbivores and carnivores have been found to have a greater diversification rate than omnivores [81] which may indicate that these traits are better for survival. However diversification can be caused either by an increase in origination relative to extinction or a decrease in extinction relative to origination. If the latter scenario occurred, this would agree with the predictions from Price et al. [81] that herbivorous and carnivorous taxa would be more successful and have had greater average survival than omnivores. Which scenario occurred, however, is (currently) impossible to determine from a phylogeny of only extant organisms [82] which means that analysis of the fossil record is necessary to estimate which scenario was most likely. If dietary category is not found to be important for modeling survival it may mean that trophic category is not a major factor for determining species level survival and that other factors, such as body size, may dominate.

Locomotor category describes the motility of a taxon and the plausibility of occurrence.
For example, an obligate arboreal taxon can only occur in locations with a minimum of tree cover and can most likely only disperse to other environments with suitable tree cover.
Dispersal ability is considered important in determining the extent of a taxon's geographic range [12, 36, 48] and thus any trait that limits dispersal ability would most likely also limit the range size of that organism and increase extinction risk. During the Cenozoic, between the Paleogene–Neogene, there was a shift from a predominately closed environment to a predominately open environment [14, 52, 92]. It is expected then that arboreal taxa during the Paleogene will have a greater expected duration than Neogene taxa, and the opposite

will be true for ground dwelling taxa. In comparison, taxon duration of scansorial taxa is expected to remain relatively similar between the two time periods because it represents a mixed environmental preference that may be viable in either closed or open environments. If locomotor category is not included in the best model of survival this may mean that it is either a poor descriptor of potential dispersal ability or range size or that other factors, both measured or unmeasured, may be of greater importance. The difficulty of a Paleogene–Neogene comparison, which is potentially underminded by heterogeneous preservation potential, will be explored in simulation as described below.

Similar to dietary category, body size has an associated energetic cost in order to maintain homeostasis, which in turn necessitates a sufficient supply of previtems. Because of this, it is expected that larger organisms have higher energetic costs and thus a greater range size in order to obtain necessary resources [18, 26, 26, 68]. It is then expected that, because taxa with larger range sizes have been found to have lower extinction rates, that species with higher average body sizes will have a lower extinction rate than taxa with smaller body sizes. This expectation, however, may not be right. As body size increases, reproductive rate decreases [56], populations are smaller [116], generations are longer [73] all of which can increase extinction risk, as has been observed [28, 66]. However, the relationship between body size and extinction rate at the generic level has been found to vary between continents [66, 109]. By expanding to include a third continent, South America, and analyzing specific level data I hope to elucidate how differences in taxonomic diversity at a continental level might affect body size mediated extinction rate. If body size is found to be unimportant for modeling survival, as in the generic level analysis of Tomiya [109], this means that other biotic or abiotic factors may dominate. This may also mean that individual level home range size does not scale into increased species level range size, and there is therefore no correlated decrease in extinction rate.

The interaction of body size, locomotor category, and dietary category is also extremely important. For example, a small bodied arboreal taxon of any trophic category during the heavily forested and warm time of the Paleogene would be expected at once to have both a small body size determined range, a large potential geographic range determined by locomotion, as well as an increased availability of resources. Together this would mean that relative survival would be expected to be less than, greater than, and greater than average respectively. Determining which factors dominate during the Paleogene, as well as other parts of the Cenozoic, must be done empirically.

Proposed research: To analyze differential mammalian survival, I propose a survival analysis approach similar to that described above for Permian brachiopods. Because dietary category and locomotor category are constant at the species level, they will be modeled as time-independent covariates of survival. While a taxon's average body size may evolve over time, the resolution of the mammal fossil record does not permit observation of this for all Cenozoic taxa. Instead, following previous work, body size will be considered constant with respect to time and will also be modeled as time-independent covariate. The climate proxy  $\delta O^{18}$  oxygen curve [118] will be modeled as an ancillary time-dependent covariate. Also, as described

above for Permian brachiopods, constant versus age dependent extinction rate will be tested by comparing the fit of survival to various probability distributions. Additionally, the effect of stratification based on the Paleogene–Neogene divide will be assessed in order to determine if the time periods had distinct patterns of survival. For this Paleogene–Neogene comparison, the principle interest will be with differences in relative trait–based survival between time periods and not with exact estimates.

While many analyses of survivorship, such as the one described above for Permian brachiopods,
are done using generic data [40, 66, 109], there are potential biases in accurately modeling
specific level processes using generic level data [86, 88, 96, 101, 111]. In order to assess some of
the differences between generic and specific level survival, I will estimate specific and generic
level survival models. Using an approach similar to previous work on estimating specific level
origination and extinction rates from generic level survival curves [32], I will measure the
deviance between extinction rate directly estimated from the specific survivorship and the
specific level extinction rates estimated from the generic level survival data. In addition to
empirical comparison between generic and specific level survival, simulations of diversification
with varying levels of cryptic speciation (anagenesis). This may also act as a proxy for generic
level diversification because a lineage having a long duration because it is not correctly broken
up can be considered analogous to a genus persisting because it continues to speciate.

Homogeneous preservation across taxa is expected to have a minimal and uniform effect on estimates of duration and survival [96]. However, the above comparison between the Paleogene-Neogene may be biased due to heterogeneous preservation between environments. Differential preservation between open and closed environments might strongly bias taxon duration estimation and strongly bias comparisons of survival. I propose a series of simulations in order to explore the effects of heterogeneous preservation on survival: two groups with identical diversification and identical preservation, two groups with identical diversification but different preservation, and two groups with different diversification and different preservation. Because phylogenies are frequently simulated as a time-homogeneous birth-death process, this will be the model used to simulate diversification. The expected survival distribution for a time-homogeneous birth-death process is exponential because of the age independent rate parameter ( $\lambda$ ).

Mammalian occurrence data will be collected primarily through a combination of the PBDB, Neogene Old World Database (NOW; http://www.helsink.fi/science/now/), and museum collections. North American fossil mammal data are well represented and vetted in the PBDB because of the extensive work of Alroy [4–6]. European fossil mammal data is also well represented between the PBDB and NOW. South American fossil mammal data is available through the PBDB, but is not particularly well vetted and has poor overall coverage. Because of this, South American fossil mammal data will be gathered via various museums such as the Field Museum of Natural History and the American Museum of Natural History as well as published occurrence compilations. With the South American taxa, taxonomy and sampling may not be as well resolved as for North and South America and it may be necessary to restrict analysis to the most taxonomically resolved and sampled groups such as

## Community connectedness in Cenozoic mammals

- Questions: How does community composition change over time in terms of the ratio between endemic and cosmopolitan taxa? Is there a single global pattern, or do different continents
   have different patterns? Do patterns differ between ecological categories? Is global climate change an important predictor of these patterns?
- Background and Predictions: Community connectedness is the degree to which localities are composed of endemic versus cosmopolitan taxa, and how similar this ratio is across localities.
- How community composition changes over time and in relation to organismal traits and a changing environment is extremely important for understanding how trophic structure changes or is maintained over time. Additionally, community connectedness is important for understanding the degree to which global, regional, or local scale processes are important for shaping environmental interactions, both biotic-biotic and biotic-abiotic.
- Community connectedness is measured here using four summary statistics: average relative number of endemic taxa per locality (E), average relative locality occupancy per taxon (Occ), biogeographic connectedness (BC), and code length [99]. These summary statistics describe how unique each locality is on average compared to all others during a time period, how relatively widespread taxa are on average during a time period, how evenly distributed taxa are amongst localities during a time period, and the degree of biogeographic provincially during a time period. Analysis of summary statistics both within and between different regions across the globe allows for the expected relative importance of global versus regional versus local processes and how this might change over time to be estimated. If global climate process are important to patterns of community connectedness and environmental interactions than it is expected that these measures will be correlated with global temperature through time. Additionally, if two or more regions have similar or correlated patterns of community connectedness, it is expected that global scale processes may play a roll in shaping the environment. Regional processes are expected to be important when E is low, Occ is high, BC is high, and code length is high. In contrast, local processes are expected to be important when E is high, Occ is low, BC is low and code length is low. The different process scales are not mutually exclusive, however, and one or more scales might be involved in shaping environmental interactions. Importantly, which process scales are important may change over time.
- In addition to total community connectedness, the dynamics of taxa within various ecological categories are important for understanding whether different adaptive zones may be differently affected by global, regional, or local scale processes. As described above, two important traits for potentially determining range size in mammals which are constant at the specific level are dietary category and locomotor category.

During the Cenozoic there was a global shift from closed, forested habitat to open, savanna-like habitat. Importantly, the timing of this environmental shift was different between continents [107, 108], so patterns of community connectedness may not be globally uniform and could reflect regional differences. The global prediction is that there would have been a relative increase in E and code length accompanied by a decrease in BC and Occ in arboreal taxa over time. The opposite is expected for terrestrial taxa. These expectations are because forested environments would likely have become increasingly patchily distributed, particularly during the Neogene compared to the Paleogene.

Based on prior work, it is expected that the patterns of biogeographic community connectedness for herbivorous taxa in a region would be most similar to that for all regional taxa and potentially "drive" the regional pattern, partially because on average this category represents the majority or plurality of taxa [54]. In contrast, community connectedness for carnivorous taxa is expected to remain constant over time or be correlated with herbivore patterns. Finally, omnivorous taxa are not expected to be correlated with the patterns of either herbivorous or carnivorous taxa and have either relatively consant or random patterns of community connectedness over time. These predictions are based on the differences in resilience and relationship to primary productivity, with herbivores being more resilient than carnivores and omnivores being random in their resilience [55]. Resilience is defined here as the ability for a taxon to increase in commonness (occupancy) after a decline [55].

An additional global trend during the Cenozoic was the shift from a "hot house" environment to an "ice house" environment [117, 118]. This transition was accompanied by major shifts in global climatic envelopes and the reorganization of mammalian communities [6, 14, 30, 35, 52]. For mammalian community connectedness there are two possible scenarios. First, while the environment was shifting, lineages may have adapted in place and overall trophic structure and community connectedness would have remained relatively constant through time, as observed during the Neogene of Europe [55]. Alternatively, species may have shifted ranges and changed the average set of taxa present at a locality which would be associated with non-stationary trophic structure and community connectedness.

At a regional scale, North American community connectedness is expected to follow the global predictions described above because the vast amount of prior synthesis has focused on North America [4–6, 9, 10, 14, 30, 37, 38, 103, 104]. However, the effect of global climate change on North American diversity remains unresolved and controversial [6, 10, 14, 30], thus it is necessary to determine empirically when global versus regional versus local scale processes may have dominated and how that may have changed over time.

The European mammalian fossil record is also well studied, though research has primarily focused on the Neogene [54, 55, 66, 83–85]. An important aspect about the European record is that during the Neogene there was little shift in relative dietary category abundance [55] and that the patterns within herbivores (browse–graze transition) were mostly driven by abundant, cosmopolitan taxa [54]. It is predicted then that herbivores will demonstrate the same patterns of community connectedness as Europe as a whole, while omnivores and

carnivores will be different from that of herbivores and may demonstrate random or constant patterns of community connectedness through time.

Patterns of community connectedness for South American mammalian fauna are comparatively
less synthesized than those of North American and Europe. Instead, cross—continental
dynamics between North and South America during the Neogene are much more studied
[72]. The South American mammalian faunal record reflects two distinct biotic provinces
between the North and the South [31, 69, 70, 78]. Because of this, it is expected that
South America will have a different pattern of community connectedness than either North
America or Europe. Also, there is an expected dramatic increase occupancy in land-dwelling
herbivores relative to arboreal and scansorial taxa related to the aridification of high—latitude
South America. Additionally, because of this strong biome distinction, it is predicted that
provinciality will be high but remain constant over time.

Proposed research: Using an approach proposed by Sidor et al. [99] and Vilhena et al. [113],

I will use bipartite biogeographic networks in order to understand patterns of community connectedness. Networks are between taxa, here defined as species, and localities, defined as

2x2 grid cells from an equal-area map projection. Networks will be made for every 2 million year bin of the Cenozoic. This bin width is chosen to always have a minimum of two localities present. Networks will also be constructed for taxa separated by dietary and locomotor category. Previous studies of mammalian occurrence patterns have restricted analysis to large bodied and well studied groups, such as Primates and Artiodactyls, in order to account for potential sampling and taxonomic biases. Here, analysis will be done using all available taxa and with a restricted sample of just major groups in order to observe any differences in patterns of community connectedness.

Community connectedness will be measured using the four summary statistics described above. E is defined as  $E = \frac{\sum_{i=1}^{L} \frac{u_i}{n_i}}{L}$  where L is as the number of localities, u is the number of taxa unique to a locality, and n is the number of taxa present at a locality. This is a measure of how unique localities are on average. Occ is the number of localities a taxon is, on average, found at and is defined  $Occ = \frac{\sum_{i=1}^{N} \frac{l_i}{L}}{N}$  where N is as the number of taxa present in the biogeographic network and l is the number of localities a taxon occurred in. BC is a measurement of the shared taxa between localities and is defined  $BC = \frac{O-N}{LN-N}$  where O is the total number of taxonomic occurrences. BC ranges from 0 to 1, with 0 meaning that each locality completely disconnected from all other localities and 1 indicating that all taxa are presents at all localities.

Code length is a measure of the information flow [98] of a graph as estimated from the behavior of a random surfer [16] on a graph and how often it visits each node and it's behavior
 with regards to moving into and out of different regions of the graph. Code length is the minimum length of a binary code for describing the behavior the surfer and the community
 structure of the nodes and is estimated via the map equation [93, 94]. The logic of the map equation is that a good map compresses reality into as few symbols as possible thus we
 want to compress as many nodes as possible into as few symbols as possible. Our goal is to

represent a graph better than just assigning a unique Huffman code to each node [43, 93]. By
compressing multiple nodes into a single code block, we decrease the minimum coding length
of a network and a network with a low code length has more nodes compressed into distinct
subunits/provinces without losing the underlying information flow of the graph. In the case
of measuring community connectedness, a low code length means greater site distinctness
and provinciality than a graph with a high code length [99].

Phylogenetic similarity between localities may play an important role in community structuring [115] such as closely related taxa being "repulsed" due to competitive exclusion or "clumped" because of environmental filtering. As a preliminary approach, for every pairwise combination of localities during a time period an informal phylogeny will be constructed for the pool of all taxa present in both localities. This informal phylogeny will be based solely on available taxonomic information such as order, family, and genus assignments with each of these levels being a completely unresolved polytomy. While there is formal phylogenetic information for many mammals that would provide greater resolution, this should not be necessary in most cases because only the general "phylogenetic proximity" is necessary for this study. Using an (informal) phylogeny, a number of measures of phylogenetic similarity can be measured for example the relative mean pairwise distance between all taxa at a locality [115] or the related phylogenetic species variability of a single locality Helmus et al. [42]. These values, calculated for all localities, can then be averaged to give an estimate of average phylogenetic similarity over the Cenozoic which can be used as a partial correlate or covariate for modeling changes in community connectedness.

- The next step is to compare patterns of community connectedness both within and between regions in order to understand if global, regional, or local scale processes dominate. Additionally, comparisons will be done between the different dietary and locomotor categories both within and between regions to determine which scale processes may be affecting either trait.

  The approach and methodology to accomplish these analyses is currently under development. Additionally, the possibility of integrating locality—locality distance or some other measure of topology will be explored, especially how this relates to code length and provinciality in general.
- The data necessary to complete this study is the same as for the above analysis of mammalian survival.
- Preliminary results Preliminary results of the community connectedness patterns of both North America and Europe based on PBDB data are presented here (Fig. 2). Both regions
   have qualitatively different patterns of community connectedness, primarily during the Paleogene. Almost all four of the summary statistics are extremely volatile over the Cenozoic,
   especially for Europe. However, some interesting qualitative patterns are present.
- There is a qualitative decrease in Occ in Europe until approximately the start of the Neogene (approximately 23 My), indicating that the average taxon is becoming generally less cosmopolitan over time. In contrast, North American Occ is qualitatively stationary over the entire Cenozoic and almost always lower than that observed for Europe. This means that, on

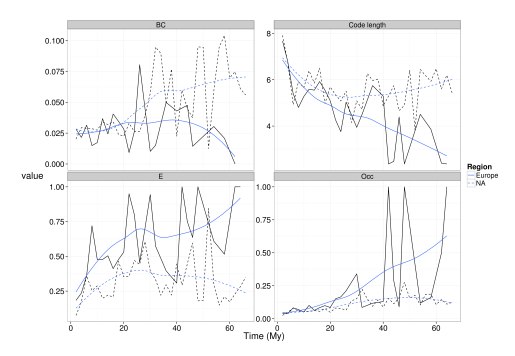


Figure 2: Biogeographic network summary statistics for mammalian communities in North America (dashed line) and Europe (solid line). The summary statistics are, clockwise from top left: biogeographic connectedness (BC), code length, average relative locality occupancy per taxon (Occ), and average relative number of endemic taxa per locality (E). Blue lines are generalized additive model smooths and are presented to illustrate the overall pattern for each region.

average, North American taxa are present in very few localities at any given point in time.

In Europe there is a qualitative rise in BC in the first few million years of the Cenozoic, but afterwards remains relatively stationary meaning that the average proportion of shared taxa remained qualitatively stationary. In comparison, North American BC remains stationary with a greater amount of shared taxa than Europe for the first half of the Cenozoic followed by a decrease and another plateau at the end of the Cenozoic.

In Europe, there is a over all qualitative decrease in E while in North America there is a qualitatively constant E over the Cenozoic with a slight decrease in the Neogene. As discussed above, E is a measure of relative uniqueness of a locality on average. Qualitatively, North America retained approximately the same amount of site uniqueness through out the Cenozoic. While the pattern of the European record shows a qualitatively nonmonotonic decrease in locality uniqueness.

The code length of European biogeographic networks increases qualitatively over the entire Cenozoic, while code length of North American networks remains relatively constant until the Neogene when there is a qualitative increase. Initial interpretation of these results indicates that North America maintains a relatively stationary degree of provinciality while Europe

has a qualitatively decreasing degree of provinciality.

## 512 Synthesis of proposed research

Underlying all of the above is a foundational question in paleobiology: why do certain taxa go extinct while others do not? In the context of evolutionary paleoecology, this question can be rephrased as "how do the set of all biotic—biotic and biotic—abiotic interactions a taxon experiences over time (i.e. adaptive zone 103) affect extinction risk?" Related to this is the Law of Constant Extinction which states that extinction risk for a given adaptive zone is taxon—age independent [110]. It is asserted that the Law of Constant Extinction only holds during periods of relatively constant environment even though this was not the context for the initial observation [65, 110] which can be interpreted as the set of dominant non-organism mediated processes do not fluctuate or fluctuate in a known manner. By understanding which non-organism mediated processes may be shaping the environment (set of all possible biotic and abiotic interactors) and how they change over time and phrasing analysis of extinction in this context, it may be possible to "test" the Law of Constant Extinction.

The first two studies proposed above investigate how organismal traits potentially related to environmental preference affect extinction rate. In effect, these traits may determine the "bounds" of a taxon's adaptive zone by limiting the total set of interactions to just those for which the taxon is adapted. The final proposed study aims to estimate what non-organism mediated processes (global, regional, and/or local) may be dominate in shaping the environment and the related set of adaptive zones. Between these studies, as well the use of two desperate groups, it should be possible to determine when, what, and if certain variables matter for survival and potentially how they matter.

## Timeline

- 534 Spring/Summer 2014
  - Evolution Meeting: preliminary brachiopod survival results
- South American fossil mammal data from Field Museum of Natural History collections Fall 2014/Winter 2015
  - GSA: survivorship simulation for anagenesis and sampling
  - Doctoral Dissertation Improvement Grant
- 540 Spring/Summer 2015

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- Evolution Meeting: mammalian survivorship analysis for North America and Europe
- South American fossil mammal data from American Museum of Natural History collections
  - write and submit survivorship simulation paper

Fall 2015/Winter 2016

- SVP or GSA: mammalian biogeographic connectedness
  - write and submit mammal connectedness paper
- 548 Spring/Summer 2016
  - Evolution Meeting: brachiopod survival analysis results
  - write and submit brachiopod survival paper

Fall 2016/Winter 2017

- SVP or GSA: mammalian survivorship analysis
- write and submit mammal survival paper
- 554 Spring/Summer 2017
  - Evolution Meeting
  - write and submit review/philosophy paper
  - Defend

## References

560

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568

582

- [1] H. Akaike. A new look at the statistical model identification. *IEEE Transactions on Automatic Control*, 19(6):716–723, 1974.
- [2] R. R. Alexander. Generic longevity of articulate brachiopods in relation to the mode of stabilization on the substrate. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 21: 209–226, 1977.
- [3] W. D. Allmon. Taxic evolutionary paleoecology and the ecological context of macroevolutionary change. *Evolutionary Ecology*, 8(2):95–112, Mar. 1994. ISSN 0269-7653. doi: 10.1007/BF01238244.
  - [4] J. Alroy. Constant extinction, constrained diversification, and uncoordinated stasis in North American mammals. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 127: 285–311, 1996.
- [5] J. Alroy. Cope's rule and the dynamics of body mass evolution in North American fossil mammals. *Science*, 280:731–734, 1998. doi: 10.1126/science.280.5364.731.
- [6] J. Alroy, P. L. Koch, and J. C. Zachos. Global climate change and North American mammalian evolution. *Paleobiology*, 26(1981):259–288, 2000.
- [7] B. M. Anderson, D. Pisani, A. I. Miller, and K. J. Peterson. The environmental affinities of marine higher taxa and possible biases in their first appearances in the fossil record. Geology, 39(10):971–974, Sept. 2011. ISSN 0091-7613. doi: 10.1130/G32413.1. URL http://geology.gsapubs.org/cgi/doi/10.1130/G32413.1.
- [8] N. W. Archbold and G. R. Shi. Permian brachiopod faunas of Western Australia: Gondwanan-Asian relationships and Permian climate. *Journal of Southeast Asian Earth Sciences*, 11(3):207–215, 1995.
  - [9] C. Badgley and J. A. Finarelli. Diversity dynamics of mammals in relation to tectonic and climatic history: comparison of three Neogene records from North America. *Paleobiology*, 39(3):373–399, Apr. 2013. ISSN 0094-8373. doi: 10.1666/12024.
- [10] A. D. Barnosky. Distinguishing the effects of the Red Queen Court Jester on Micene Mammal Evolution in the Northern Rocky Mountains. *Journal of Vertebrate Paleontology*, 21(1):172–185, 2001.
  - [11] T. K. Baumiller. Survivorship analysis of Paleozoic Crinoidea: effect of filter morphology on evolutionary rates. *Paleobiology*, 19(3):304–321, 1993.
- [12] A. Birand, A. Vose, and S. Gavrilets. Patterns of species ranges, speciation, and extinction. *The American naturalist*, 179(1):1–21, Jan. 2012. ISSN 1537-5323. doi: 10.1086/663202. URL http://www.ncbi.nlm.nih.gov/pubmed/22173457.
- [13] L. P. Birgenheier, T. D. Frank, C. R. Fielding, and M. C. Rygel. Coupled carbon isotopic and sedimentological records from the Permian system of eastern Australia

- reveal the response of atmospheric carbon dioxide to glacial growth and decay during the late Palaeozoic Ice Age. *Palaeogeography*, *Palaeoclimatology*, *Palaeoecology*, 286(3-4): 178-193, 2010. URL http://dx.doi.org/10.1016/j.palaeo.2010.01.008papers2: //publication/doi/10.1016/j.palaeo.2010.01.008.
- [14] J. L. Blois and E. A. Hadly. Mammalian Response to Cenozoic Climatic Change. Annual Review of Earth and Planetary Sciences, 37(1):181–208, May 2009. ISSN 0084-6597. doi: 10.1146/annurev.earth.031208.100055.
  - [15] D. J. Bottjer and D. Jablonski. Paleoenvironmental patterns in the evolution of Post-Palezoic benthic marine invertebrates. *Palaios*, 3(6):540–560, 1988.

- [16] S. Brin and L. Page. The anatomy of a large-scale hypertextual Web search engine. Computer Networks and ISDN Systems, 30(1-7):107-117, Apr. 1998. ISSN 01697552. doi: 10.1016/S0169-7552(98)00110-X. URL http://linkinghub.elsevier.com/retrieve/pii/S016975529800110X.
- [17] J. H. Brown. On the Relationship between Abundance and Distribution of Species.

  The American Naturalist, 124(2):255, Aug. 1984. ISSN 0003-0147. doi: 10.1086/284267.

  URL http://www.journals.uchicago.edu/doi/abs/10.1086/284267.
- [18] J. H. Brown and B. A. Maurer. Evolution of species assembalges: effects of energetic constraints and species dynamics on the diversification of the North American avidauna.

  The American Naturalist, 130(1):1–17, 1987.
  - [19] K. P. Burnham and D. R. Anderson. *Model selection and multi-model inference: a practical information-theoretic approach*. Springer, New York, 2nd edition, 2002.
- [20] M. E. Clapham and D. J. Bottjer. Permian marine paleoecology and its implications for large-scale decoupling of brachiopod and bivalve abundance and diversity during the Lopingian (Late Permian). *Palaeogeography, Palaeoclimatology, Palaeoecology*, 249 (3-4):283-301, June 2007. ISSN 00310182. doi: 10.1016/j.palaeo.2007.02.003. URL http://linkinghub.elsevier.com/retrieve/pii/S0031018207000600.
- [21] M. E. Clapham and D. J. Bottjer. Prolonged Permian Triassic ecological crisis recorded by molluscan dominance in Late Permian offshore assemblages. Proceedings of the National Academy of Sciences of the United States of America, 104(32):12971-5, Aug. 2007. ISSN 0027-8424. doi: 10.1073/pnas.0705280104. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=1941817&tool=pmcentrez&rendertype=abstract.
- [22] M. E. Clapham and N. P. James. Paleoecology Of Early-Middle Permian Marine Communities In Eastern Australia: Response To Global Climate Change In the Aftermath Of the Late Paleozoic Ice Age. Palaios, 23(11):738-750, Nov. 2008. ISSN 0883-1351. doi: 10.2110/palo.2008.p08-022r. URL http://palaios.sepmonline.org/cgi/doi/10.2110/palo.2008.p08-022r.

- [23] M. E. Clapham and N. P. James. Regional-scale marine faunal change in Eastern Australia during Permian climate fluctuations and its relationship to local community restructuring. *Palaios*, 27:627–635, Oct. 2012. ISSN 0883-1351. doi: 10.2110/palo.2012.p12-003r. URL http://palaios.sepmonline.org/cgi/doi/10. 2110/palo.2012.p12-003r.
- [24] M. E. Clapham, D. J. Bottjer, C. M. Powers, N. Bonuso, M. L. Fraiser, P. J. Marenco, S. Q. Dornbos, and S. B. Pruss. Assessing the Ecological Dominance of Phanero-zoic Marine Invertebrates. *Palaios*, 21(5):431-441, Oct. 2006. ISSN 0883-1351. doi: 10.2110/palo.2005.P05-017R. URL http://palaios.sepmonline.org/cgi/doi/10. 2110/palo.2005.P05-017R.
  - [25] W. S. Cooper. Expected time to extinction and the concept of fundamental fitness. Journal of Theoretical Biology, 107:603–629, 1984.

- [26] J. Damuth. Home range, home range overlap, and species energy use among herbivorous mammals. *Biological Journal of the Linnean Society*, 15:185–193, 1979.
- [27] J. Damuth. Population density and body size in mammals. Nature, 290:699-700, 1981. URL http://www.jstor.org/stable/10.2307/2461771http://www.nature.com/nature/journal/v290/n5808/abs/290699a0.html.
- [28] A. D. Davidson, A. G. Boyer, H. Kim, S. Pompa-Mansilla, M. J. Hamilton, D. P. Costa, G. Ceballos, and J. H. Brown. Drivers and hotspots of extinction risk in marine mammals.

  Proceedings of the National Academy of Sciences, 109(9):3395-400, Feb. 2012. ISSN 1091-6490. doi: 10.1073/pnas.1121469109. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=3295301&tool=pmcentrez&rendertype=abstract.
- [29] C. R. Fielding, K. L. Bann, J. A. Maceachern, S. C. Tye, and B. G. Jones. Cyclicity in the nearshore marine to coastal, Lower Permian, Pebbley Beach Formation, southern Sydney Basin, Australia: a record of relative sea-level fluctuations at the close of the Late Palaeozoic Gondwanan ice age. Sedimentology, 53(2):435–463, 2006. URL http://doi.wiley.com/10.1111/j.1365-3091.2006.00770.xpapers2: //publication/doi/10.1111/j.1365-3091.2006.00770.x.
- [30] B. Figueirido, C. M. Janis, J. A. Pérez-Claros, M. De Renzi, and P. Palmqvist. Cenozoic climate change influences mammalian evolutionary dynamics. *Proceedings of the National Academy of Sciences*, 109(3):722–727, Jan. 2012. ISSN 1091-6490. doi: 10.1073/pnas.1110246108.
- [31] J. J. Flynn and A. R. Wyss. Recent advances in South American mammalian paleontology. *Trends in ecology & evolution*, 13(11):449–54, Nov. 1998. ISSN 0169-5347. URL http://www.ncbi.nlm.nih.gov/pubmed/21238387.
- [32] M. Foote. Survivorship analysis of Cambrian and Ordovician Trilobites. *Paleobiology*, 14(3):258–271, 1988.

- [33] M. Foote. Substrate affinity and diversity dynamics of Paleozoic marine animals. Paleobiology, 32(3):345–366, Sept. 2006. ISSN 0094-8373. doi: 10.1666/05062.1. URL http://www.bioone.org/doi/abs/10.1666/05062.1.
- [34] M. Foote and A. I. Miller. Determinants of early survival in marine animal genera.

  \*Paleobiology\*, 39(2):171–192, Mar. 2013. ISSN 0094-8373. doi: 10.1666/12028. URL http://www.bioone.org/doi/abs/10.1666/12028.
- [35] M. Fortelius, J. Eronen, J. Jernvall, L. Liu, D. Pushkina, J. Rinne, A. Tesakov, I. Vislobokova, Z. Zhang, and L. Zhou. Fossil mammals resolve regional patterns of Eurasian climate change over 20 million years. *Evolutionary Ecology Research*, 4: 1005–1016, 2002.
- [36] K. J. Gaston. Geographic range limits: achieving synthesis. *Proceedings. Biological sciences / The Royal Society*, 276(1661):1395-406, Apr. 2009. ISSN 0962-8452. doi: 10.1098/rspb.2008.1480. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=2677218&tool=pmcentrez&rendertype=abstract.
- [37] G. Gunnell, M. Morgan, M. C. Mass, and P. D. Gingerich. Comparative paleoecology of Paleogene and Neogene mammalian faunas: trophic structure and composition.

  Palaeogeography, Palaeoclimatology, Palaeoecology, 115:265–286, 1995. URL http://www.sciencedirect.com/science/article/pii/0031018294001150.
- [38] E. A. Hadly and B. A. Maurer. Spatial and temporal patterns of species diversity in montane mammal communities of western North America. *Evolutionary Ecology Research*, 3:477–486, 2001.
- [39] P. G. Harnik. Direct and indirect effects of biological factors on extinction risk in fossil bivalves. Proceedings of the National Academy of Sciences of the United States of America, 108(33):13594-9, Aug. 2011. ISSN 1091-6490. doi: 10.1073/pnas.1100572108. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=3158225&tool=pmcentrez&rendertype=abstract.
- [40] P. G. Harnik, C. Simpson, and J. L. Payne. Long-term differences in extinction risk among the seven forms of rarity. *Proceedings of the Royal Society B: Biological Sciences*, (October), Oct. 2013. ISSN 0962-8452. doi: 10.1098/rspb.2012.1902. URL http://rspb.royalsocietypublishing.org/cgi/doi/10.1098/rspb.2012.1902.
- [41] N. A. Heim and S. E. Peters. Regional environmental breadth predicts geographic range and longevity in fossil marine genera. *PloS one*, 6(5):e18946, Jan. 2011. ISSN 1932-6203. doi: 10.1371/journal.pone.0018946. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=3087726&tool=pmcentrez&rendertype=abstract.
- [42] M. R. Helmus, T. J. Bland, C. K. Williams, and A. R. Ives. Phylogenetic Measures of Biodiversity. The American naturalist, 169(3), Jan. 2007. ISSN 1537-5323. doi: 10.1086/511334. URL http://www.ncbi.nlm.nih.gov/pubmed/17230400.

- [43] D. Huffman. A Method for the Construction of Minimum-Redundancy Codes.

  \*\*Proceedings of the IRE, 40(9):1098-1101, Sept. 1952. ISSN 0096-8390. doi: 10.

  1109/JRPROC.1952.273898. URL http://ieeexplore.ieee.org/lpdocs/epic03/

  \*\*wrapper.htm?arnumber=4051119.\*\*
- [44] C. M. Hurvich and C.-L. Tsai. Regression and time series model selection in small samples. *Biometrika*, 76(2):297-307, 1989. URL http://biomet.oxfordjournals.org/content/76/2/297.shortpapers2: //publication/uuid/FC795396-A722-468D-AF8E-4E1F7C97998C.
  - [45] D. Jablonski. Background and mass extincitons: the alternation of macroevolutionary regimes. *Science*, 231(4734):129–133, 1986.

716

718

- [46] D. Jablonski. Species Selection: Theory and Data. Annual Review of Ecology, Evolution, and Systematics, 39(1):501-524, Dec. 2008. ISSN 1543-592X. doi: 10.1146/annurev. ecolsys.39.110707.173510. URL http://www.annualreviews.org/doi/abs/10.1146/annurev.ecolsys.39.110707.173510.
- [47] D. Jablonski and D. J. Bottjer. Environmental patterns in the origins of higher taxa: the post-paleozoic fossil record. Science (New York, N.Y.), 252(5014):1831—3, June 1991. ISSN 0036-8075. doi: 10.1126/science.252.5014.1831. URL http://www.ncbi.nlm.nih.gov/pubmed/17753259.
- [48] D. Jablonski and G. Hunt. Larval ecology, geographic range, and species survivorship in Cretaceous mollusks: organismic versus species-level explanations. *The American naturalist*, 168(4):556–64, Oct. 2006. ISSN 1537-5323. doi: 10.1086/507994. URL http://www.ncbi.nlm.nih.gov/pubmed/17004227.
- [49] D. Jablonski and R. a. Lutz. Larval Ecology of Marine Benthic Invertebrates: Paleobiological Implications. *Biological Reviews*, 58(1):21–89, Feb. 1983. ISSN 1464-7931. doi: 10.1111/j.1469-185X.1983.tb00380.x. URL http://doi.wiley.com/10.1111/j. 1469-185X.1983.tb00380.x.
- [50] D. Jablonski and K. Roy. Geographical range and speciation in fossil and living molluscs.

  \*\*Proceedings. Biological sciences / The Royal Society, 270(1513):401-6, Feb. 2003. ISSN 0962-8452. doi: 10.1098/rspb.2002.2243. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=1691247&tool=pmcentrez&rendertype=abstract.
  - [51] D. Jablonski, J. J. Sepkoski, D. J. Bottjer, and P. M. Sheehan. Onshore-offshore patterns in the evolution of Phanerozoic shelf communities. *Science*, 222(4628):1123–1125, 1983.
- [52] C. M. Janis. Tertiary mammal evolution in the context of changing climates, vegetation, and tectonic events. *Annual Review of Ecology and Systematics*, 24:467–500, 1993.
- [53] C. M. Janis, J. Damuth, and J. M. Theodor. Miocene ungulates and terrestrial primary productivity: where have all the browsers gone? *Proceedings* of the National Academy of Sciences, 97(14):7899–904, July 2000. ISSN 0027-

- National Nat
- [54] J. Jernvall and M. Fortelius. Common mammals drive the evolutionary increase of hypsodonty in the Neogene. *Nature*, 417(6888):538–40, May 2002. ISSN 0028-0836. doi: 10.1038/417538a.
- [55] J. Jernvall and M. Fortelius. Maintenance of trophic structure in fossil mammal communities: site occupancy and taxon resilience. *American Naturalist*, 164(5):614–624, Nov. 2004. ISSN 1537-5323. doi: 10.1086/424967.
- <sup>750</sup> [56] C. N. Johnson. Determinate of loss of mammal species during the Late Quaternary 'megafauna' extinctions: life history and ecology, but not body size. *Proceedings of the Royal Society B: Biological Sciences*, 269:2221–2227, 2002. doi: 10.1098/rspb.2002.2130.
- [57] A. T. Jones, T. D. Frank, and C. R. Fielding. Cold climate in the eastern

  Australian mid to late Permian may reflect cold upwelling waters. Palaeogeography, Palaeoclimatology, Palaeoecology, 237(2-4):370-377, 2006. URL

  http://linkinghub.elsevier.com/retrieve/pii/S0031018205007327papers2:
  //publication/doi/10.1016/j.palaeo.2005.12.009.
- <sup>758</sup> [58] W. Kiessling and M. Aberhan. Environmental determinants of marine benthic biodiversity dynamics through Triassic Jurassic time. *Paleobiology*, 33(3):414–434, 2007.
- [59] W. Kiessling, M. Aberhan, B. Brenneis, and P. J. Wagner. Extinction trajectories of benthic organisms across the TriassicJurassic boundary. *Palaeogeography*,
   Palaeoclimatology, Palaeoecology, 244(1-4):201-222, Feb. 2007. ISSN 00310182. doi: 10.1016/j.palaeo.2006.06.029. URL http://linkinghub.elsevier.com/retrieve/pii/S0031018206004494.
- [60] J. Kitchell. Biological selectivity of extinction. In E. G. Kaufman and O. H. Walliser, editors, *Extinction Events in Earth History*, pages 31–43. Springer-Verlang, Berlin, 1990. URL http://link.springer.com/chapter/10.1007/BFb0011132.
- <sup>768</sup> [61] J. A. Kitchell. Evolutionary paleoecology: recent contributions to evolutionary theory. Paleobiology, 11(1):91–104, 1985.
- [62] D. G. Kleinbaum and M. Klein. Survival analysis: a self-learning text. Springer, New York, NY, 2 edition, 2005.
- [63] M. LaBarbera. Brachiopod orientation to water movement: functional morphology. Lethaia, 11(1):67-79, Jan. 1978. ISSN 0024-1164. doi: 10.1111/j.1502-3931.1978. tb01219.x. URL http://doi.wiley.com/10.1111/j.1502-3931.1978.tb01219.x.
- [64] M. LaBarbera. Water flow patterns in and around three species of articulate brachiopods.

  Journal of Experimental Marine Biology and Ecology, 55:185–206, 1981.
  - [65] L. H. Liow. Does versatility as measured by geographic range, bathymetric range and

- morphological variability contribute to taxon longevity? Global Ecology and Biogeography, 16(1):117-128, 2007. URL http://doi.wiley.com/10.1111/j.1466-8238.2006. 00269.xpapers2://publication/doi/10.1111/j.1466-8238.2006.00269.x.
- [66] L. H. Liow, M. Fortelius, E. Bingham, K. Lintulaakso, H. Mannila, L. Flynn, and N. C. Stenseth. Higher origination and extinction rates in larger mammals. *Proceedings of the National Academy of Sciences*, 105(16):6097, 2008. URL papers2://publication/uuid/5A93DDA3-204F-4D9C-AFAC-A15722A18C61.
- [67] S. K. Lyons. A quantitative model for assessing community dynamics of pleistocene mammals. *The American naturalist*, 165(6):E168–85, June 2005. ISSN 1537-5323. doi: 10.1086/429699. URL http://www.ncbi.nlm.nih.gov/pubmed/15937741.
- [68] S. K. Lyons, P. J. Wagner, and K. Dzikiewicz. Ecological correlates of range shifts of Late Pleistocene mammals. *Philosophical transactions of the Royal Society of London.*Series B, Biological sciences, 365(1558):3681-93, Nov. 2010. ISSN 1471-2970. doi: 10.1098/rstb.2010.0263. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=2981999&tool=pmcentrez&rendertype=abstract.
- [69] B. J. Macfadden. Origin and evolution of the grazing guild in New World terrestrial mammals. *Trends in ecology & evolution*, 12(5):182–187, 1997.
- [70] B. J. Macfadden. Extinct mammalian biodiversity of the ancient New World tropics.

  Trends in ecology & evolution, 21(3):157-65, Mar. 2006. ISSN 0169-5347. doi: 10.1016/j.tree.2005.12.003. URL http://www.ncbi.nlm.nih.gov/pubmed/16701492.
- <sup>798</sup> [71] G. A. Mark and K. W. Flessa. A test for evolutionary equilibria: Phanerozoic brachiopods and Cenozoic mammals. *Paleobiology*, 3(1):17–22, 1977.
- [72] L. G. Marshall, S. D. Webb, J. J. Sepkoski, and D. M. Raup. Mammalian evolution and the Great American interchange. *Science*, 215(4538):1351–1357, 1982.
- [73] A. P. Martin and S. R. Palumbi. Body size, metabolic rate, generation time, and the molecular clock. *Proceedings of the National Academy of Sciences*, 90(9):4087-91, May 1993. ISSN 0027-8424. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=46451&tool=pmcentrez&rendertype=abstract.
- [74] A. I. Miller and S. R. Connolly. Substrate affinities of higher taxa and the Ordovician Radiation. *Paleobiology*, 27(4):768-778, Dec. 2001. ISSN 0094-8373. doi: 10.1666/0094-8373(2001)027\(0768:SAOHTA\)\(0)20.CO;\(0.2001)20.C
- [75] S. Nürnberg and M. Aberhan. Habitat breadth and geographic range predict diversity dynamics in marine Mesozoic bivalves. *Paleobiology*, 39(3):360–372, Apr. 2013. ISSN 0094-8373. doi: 10.1666/12047. URL http://www.bioone.org/doi/abs/10.1666/12047.

- [76] T. Olszewski and D. Erwin. Dynamic response of Permian brachiopod communities to long-term environmental change. *Nature*, 428(April):2-5, 2004. doi: 10. 1038/nature02471.1. URL http://www.nature.com/nature/journal/v428/n6984/abs/nature02464.html.
- [77] M. E. Palmer and M. W. Feldman. Survivability is more fundamental than evolvability. *PloS one*, 7(6):e38025, Jan. 2012. ISSN 1932-6203. doi: 10.1371/journal.pone. 0038025. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid= 3377627&tool=pmcentrez&rendertype=abstract.
- [78] B. Patterson and R. Pascual. The fossil mammal fauna of South America. *The Quarterly review of biology*, 43(4):409–451, 1968.
- [79] S. E. Peters. Environmental determinants of extinction selectivity in the fossil record. Nature, 454(7204):626-9, July 2008. ISSN 1476-4687. doi: 10.1038/nature07032. URL http://www.ncbi.nlm.nih.gov/pubmed/18552839.
- [80] M. G. Powell. Latitudinal diversity gradients for brachiopod genera during late Palaeozoic time: links between climate, biogeography and evolutionary rates. *Global Ecology* and Biogeography, 16(4):519–528, July 2007. ISSN 1466-822X. doi: 10.1111/j.1466-8238. 2007.00300.x. URL http://doi.wiley.com/10.1111/j.1466-8238.2007.00300.x.
- [81] S. A. Price, S. S. B. Hopkins, K. K. Smith, and V. L. Roth. Tempo of trophic evolution and its impact on mammalian diversification. *Proceedings of the National Academy of Sciences of the United States of America*, 109(18):7008–12, May 2012. ISSN 1091-6490. doi: 10.1073/pnas.1117133109. URL http://www.ncbi.nlm.nih.gov/pubmed/22509033.
- [82] D. L. Rabosky. Extinction rates should not be estimated from molecular phylogenies. *Evolution*, 64(6):1816–24, June 2010. ISSN 1558-5646. doi: 10.1111/j.1558-5646.2009. 00926.x. URL http://www.ncbi.nlm.nih.gov/pubmed/20030708.
- [83] P. Raia, P. Piras, and T. Kotsakis. Turnover pulse or Red Queen? Evidence from the large mammal communities during the Plio-Pleistocene of Italy. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 221(3-4):293-312, June 2005. ISSN 00310182. doi: 10.1016/j.palaeo.2005.02.014. URL http://linkinghub.elsevier.com/retrieve/pii/S0031018205001033.
- [84] P. Raia, C. Meloro, A. Loy, and C. Barbera. Species occupancy and its course in the past: macroecological patterns in extinct communities. Evolutionary Ecology Research, 8:181–194, 2006.
- [85] P. Raia, F. Carotenuto, J. T. Eronen, and M. Fortelius. Longer in the tooth, shorter in the record? The evolutionary correlates of hypsodonty in Neogene ruminants. *Proceedings. Biological sciences / The Royal Society*, 278(1724):3474-81, Dec. 2011. ISSN 1471-2954. doi: 10.1098/rspb.2011.0273. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=3189366&tool=pmcentrez&rendertype=abstract.

- [86] D. M. Raup. Taxonomic survivorship curves and Van Valen's Law. *Paleobiology*, 1 (1):82–96, Jan. 1975. ISSN 0036-8075. doi: 10.1126/science.49.1254.50. URL http://www.ncbi.nlm.nih.gov/pubmed/17777225.
  - [87] D. M. Raup. Cohort Analysis of generic survivorship. *Paleobiology*, 4(1):1–15, 1978.
- <sup>856</sup> [88] D. M. Raup. A kill curve for Phanerozoic marine species. *Paleobiology*, 17(1):37–48, 1991.
- [89] D. M. Raup and G. E. Boyajian. Patterns of generic extinction in the fossil record. Paleobiology, 14(2):109–125, 1988.
- [90] J. R. Richardson. Ecology of articulated brachiopods. In A. Williams, C. H. C. Brunton, and S. J. Carlson, editors, *Treatise on Invertebrate Paleontology, Part H, Brachiopoda* 1, pages 441–462. The Geological Society of America, Boulder, Colorado, 1997.
- [91] J. R. Richardson. Biogeography of articulated brachiopods. In A. Williams, C. H. C.
   Brunton, and S. J. Carlson, editors, Treatise on Invertebrate Paleontology, Part H,
   Brachiopoda 1, pages 463–472. The Geological Society of America, Boulder, Colorado,
   1997.
  - [92] K. D. Rose. The beginning of the age of mammals. Johns Hopkins University Press, Baltimore, Md, 2006.

- [93] M. Rosvall and C. T. Bergstrom. Maps of random walks on complex networks reveal community structure. Proceedings of the National Academy of Sciences, 105(4):1118-23, Jan. 2008. ISSN 1091-6490. doi: 10.1073/pnas. 0706851105. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=2234100&tool=pmcentrez&rendertype=abstract.
- [94] M. Rosvall, D. Axelsson, and C. Bergstrom. The map equation. *The European Physical Journal Special Topics*, 178(14):13-24, 2009. URL http://www.springerlink.com/index/H8193132U6432363.pdf.
- [95] K. Roy, G. Hunt, D. Jablonski, A. Z. Krug, and J. W. Valentine. A macroevolutionary perspective on species range limits. *Proceedings. Biological sciences / The Royal Society*, 276(1661):1485-93, Apr. 2009. ISSN 0962-8452. doi: 10.1098/rspb.2008.1232. URL http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=2677224&tool=pmcentrez&rendertype=abstract.
- [96] J. J. Sepkoski. Stratigraphic biases in the analysis of taxonomic survivorship. *Paleobiology*, 1(4):343–355, 1975.
- [97] J. J. Sepkoski. A model of onshore-offshore change in faunal diversity. *Paleobiology*, 17 (1):58–77, 1991.
- [98] C. E. Shannon. A mathematical theory of communication. The Bell System Tehenical

- $Journal, \ 27:379-423,623-656, \ 1948. \ \ URL \ \ http://dl.acm.org/citation.cfm?id=584093papers2://publication/uuid/D2D2682E-1332-4611-96E0-A0B51712373C.$
- [99] C. A. Sidor, D. A. Vilhena, K. D. Angielczyk, A. K. Huttenlocker, S. J. Nesbitt,
   B. R. Peecook, J. S. Steyer, R. M. H. Smith, and L. A. Tsuji. Provincialization of terrestrial faunas following the end-Permian mass extinction. *Proceedings of the National Academy of Sciences*, 110(20):8129–33, May 2013. ISSN 1091-6490. doi: 10.1073/pnas.1302323110.
- [100] M. Silva, J. H. Brown, and J. a. Downing. Differences in Population Density and Energy Use between Birds and Mammals: A Macroecological Perspective. *The Journal of Animal Ecology*, 66(3):327, May 1997. ISSN 00218790. doi: 10.2307/5979. URL http://www.jstor.org/stable/5979?origin=crossref.
- <sup>898</sup> [101] C. Simpson. Levels of selection and large-scale morphological trends. PhD thesis, University of Chicago, 2006.
- [102] C. Simpson and P. G. Harnik. Assessing the role of abundance in marine bivalve extinction over the post-Paleozoic. *Paleobiology*, 35(4):631-647, Dec. 2009. ISSN 0094-8373. doi: 10.1666/0094-8373-35.4.631. URL http://www.bioone.org/doi/abs/10.1666/0094-8373-35.4.631.
- 904 [103] G. G. Simpson. Tempo and Mode in Evolution. Columbia University Press, New York, 1944.
- 906 [104] G. G. Simpson. *The Major Features of Evolution*. Columbia University Press, New York, 1953.
- [105] F. A. Smith, J. Brown, J. Haskell, and S. Lyons. Similarity of mammalian body size across the taxonomic hierarchy and across space and .... The American Naturalist, 2004. URL http://www.journals.uchicago.edu/doi/abs/10.1086/382898papers2://publication/uuid/D5606802-FD91-49EB-BE2F-E2D314A5E71D.
- [106] F. A. Smith, S. K. Lyons, S. Morgan Ernest, and J. H. Brown. Macroecology: more than the division of food and space among species on continents. *Progress in Physical Geography*, 32(2):115–138, Apr. 2008. ISSN 0309-1333. doi: 10.1177/0309133308094425.
   URL http://ppg.sagepub.com/cgi/doi/10.1177/0309133308094425.
- [107] C. A. E. Strömberg. Decoupled taxonomic radiation and ecological expansion of openhabitat grasses in the Cenozoic of North America. *Proceedings of the National Academy* of Sciences of the United States of America, 102(34):11980-4, Aug. 2005. ISSN 0027-8424. doi: 10.1073/pnas.0505700102. URL http://www.pubmedcentral.nih.gov/ articlerender.fcgi?artid=1189350&tool=pmcentrez&rendertype=abstract.
- [108] C. A. E. Strömberg, R. E. Dunn, R. H. Madden, M. J. Kohn, and A. A. Carlini.
  Decoupling the spread of grasslands from the evolution of grazer-type herbivores in

- South America. Nature communications, 4:1478, Jan. 2013. ISSN 2041-1723. doi: 10.1038/ncomms2508. URL http://www.ncbi.nlm.nih.gov/pubmed/23403579.
- [109] S. Tomiya. Body Size and Extinction Risk in Terrestrial Mammals Above the Species Level. *The American Naturalist*, pages E000–E000, Sept. 2013. ISSN 00030147. doi: 10.1086/673489. URL http://www.jstor.org/stable/info/10.1086/673489.
- 928 [110] L. Van Valen. A new evolutionary law. Evolutionary Theory, 1:1-30, 1973. URL http://ci.nii.ac.jp/naid/10011264287/.
- 930 [111] L. Van Valen. Taxonomic survivorship curves. Evolutionary Theory, 4:129–142, 1979.
  - [112] L. Van Valen. Three paradigms of evoltuion. Evolutionary Theory, 9:1–17, 1989.
- [113] D. A. Vilhena, E. B. Harris, C. T. Bergstrom, M. E. Maliska, P. D. Ward, C. A. Sidor, C. A. E. Strömberg, and G. P. Wilson. Bivalve network reveals latitudinal selectivity
   gradient at the end-Cretaceous mass extinction. Scientific reports, 3:1790, May 2013. ISSN 2045-2322. doi: 10.1038/srep01790.
- 936 [114] J. B. Waterhouse. Late Palaeozoic Mollusca and correlations from the south-east Bowen Basin, east Australia. *Palaeontographica Abteilung A*, 198:129–233, 1987.
- [115] C. O. Webb, D. D. Ackerly, M. a. McPeek, and M. J. Donoghue. Phylogenies and Community Ecology. *Annual Review of Ecology and Systematics*, 33(1):475-505, Nov. 2002. ISSN 0066-4162. doi: 10.1146/annurev.ecolsys.33.010802.150448. URL http://arjournals.annualreviews.org/doi/abs/10.1146%2Fannurev.ecolsys.33.010802.150448.
- [116] E. P. White, S. K. M. Ernest, A. J. Kerkhoff, and B. J. Enquist. Relationships between body size and abundance in ecology. TRENDS in Ecology and Evolutioncology & evolution, 22(6):323–30, June 2007. ISSN 0169-5347. doi: 10.1016/j.tree.2007.03.007. URL http://www.ncbi.nlm.nih.gov/pubmed/17399851.
- [117] J. C. Zachos, M. Pagani, L. Sloan, E. Thomas, and K. Billups. Trends, rhythms, and aberrations in global climate 65 Ma to present. Science, 292:686-693, 2001. doi: 10.1126/science.1059412. URL http://www.sciencemag.org/content/292/5517/686.short.
- [118] J. C. Zachos, G. R. Dickens, and R. E. Zeebe. An early Cenozoic perspective on greenhouse warming and carbon-cycle dynamics. *Nature*, 451(7176):279–283, Jan. 2008. ISSN 1476-4687. doi: 10.1038/nature06588.