

The latitudinal gradient of beta diversity in relation to climate and topography for mammals in North America

Hong Qian^{1*}, Catherine Badgley² and David L. Fox³

¹Research and Collections Center, Illinois State Museum, 1011 East Ash Street, Springfield, IL 62703, USA, ²Museum of Paleontology, University of Michigan, 1109 Geddes Road, Ann Arbor, MI 48109, USA, ³Department of Geology and Geophysics, University of Minnesota, 310 Pillsbury Drive, Minneapolis, MN 55455, USA

ABSTRACT

Aim Spatial turnover of species, or beta diversity, varies in relation to geographical distance and environmental conditions, as well as spatial scale. We evaluated the explanatory power of distance, climate and topography on beta diversity of mammalian faunas of North America in relation to latitude.

Location North America north of Mexico.

Methods The study area was divided into 313 equal-area quadrats (241×241 km). Faunal data for all continental mammals were compiled for these quadrats, which were divided among five latitudinal zones. These zones were comparable in terms of latitudinal and longitudinal span, climatic gradients and elevational gradients. We used the natural logarithm of the Jaccard index (ln*J*) to measure species turnover between pairs of quadrats within each latitudinal zone. The slope of ln*J* in relation to distance was compared among latitudinal zones. We used partial regression to partition the variance in ln*J* into the components uniquely explained by distance and by environmental differences, as well as jointly by distance and environmental differences.

Results Mammalian faunas of North America differ more from each other at lower latitudes than at higher latitudes. Regression models of ln*J* in relation to distance, climatic difference and topographic difference for each zone demonstrated that these variables have high explanatory power that diminishes with latitude. Beta diversity is higher for zones with higher mean annual temperature, lower seasonality of temperature and greater topographic complexity. For each latitudinal zone, distance and environmental differences explain a greater proportion of the variance in ln*J* than distance, climate or topography does separately.

Main conclusions The latitudinal gradient in beta diversity of North American mammals corresponds to a macroclimatic gradient of decreasing mean annual temperature and increasing seasonality of temperature from south to north. Most of the variance in spatial turnover is explained by distance and environmental differences jointly rather than distance, climate or topography separately. The high predictive power of geographical distance, climatic conditions and topography on spatial turnover could result from the direct effects of physical limiting factors or from ecological and evolutionary processes that are also influenced by the geographical template.

Keywords

Beta diversity, climate, Jaccard index, latitudinal diversity gradient, macroecology, mammalian faunas, spatial turnover.

INTRODUCTION

IL 62703, USA.

*Correspondence: Hong Qian, Research and Collections Center, Illinois State Museum,

1011 East Ash Street, Springfield,

E-mail: hqian@museum.state.il.us

The distribution of species ranges across regions and continents determines large-scale diversity gradients as well as local species richness of floras and faunas. Patterns of species richness and turnover vary across the earth at multiple spatial scales (Hutchinson, 1959; Gaston, 2000; Hillebrand, 2004). At large scales, the most familiar of these patterns is the latitudinal diversity gradient (Pianka, 1966; Rosenzweig, 1995; Willig *et al.*, 2003; Mittelbach *et al.*, 2007). At a specified scale, variation in spatial

turnover in relation to the environmental properties of the landscape provides evidence for causal mechanisms influencing recurrent patterns in spatial turnover of species, or beta diversity (Whittaker, 1977). Beta diversity can be measured with respect to geographical distance or differences in environmental variables (e.g. Qian *et al.*, 2005; Gaston *et al.*, 2007; Qian & Ricklefs, 2007).

Studies of beta diversity, while less numerous than those of species richness per se, focus on the geographical and environmental contexts in which species composition changes by addition or subtraction of species from local assemblages. Evaluation of these relationships provides fundamental insights into species richness in space and time (Rosenzweig, 1995). However, studies of beta diversity at the continental scale are few, and most of these have sought to determine whether the focal taxonomic group exhibits a latitudinal gradient in beta diversity without evaluating the contributions of the physical context - climatic and physiographic variables - to beta diversity. For many widely distributed groups of organisms, species inhabiting temperate and polar regions have larger geographical ranges than do tropically situated species - the pattern known as Rapoport's rule (Stevens, 1989). A corollary is that species turnover across the landscape decreases at higher latitudes, leading to the expectation of a latitudinal gradient in beta diversity. Studies of beta diversity thus far document patterns that vary in relation to taxonomic group, taxonomic level, spatial scale and geographical region. For example, Koleff et al. (2003) reported increased species turnover with distance for New World owls (Strigidae and Tytonidae) closer to the equator. For birds of the world, Gaston et al. (2007) evaluated turnover in relation to latitude, species richness and environmental conditions. Turnover with distance in relation to latitude showed a variable pattern. In the Northern Hemisphere, turnover was highest at low latitudes, peaking at 20° N, and declining up to 60° N, then rising at higher latitudes. Qian & Ricklefs (2007) found that beta diversity in relation to distance for North American vascular-plant assemblages decreased from low to high latitudes, representing a gradient of decreasing temperature and increasing seasonal variation in temperature, and within latitudinal zones from species to genera to families. Among studies of North American mammals, Pagel et al. (1991) documented a positive relationship between geographical range size and latitude, as well as a negative relationship between range size and longitude – implying higher turnover at lower latitudes and in western regions. Arita (1997) documented high beta diversity in relation to environmental heterogeneity (elevation and climatic variability) for Mexican mammals. Both Kaufman & Willig (1998) and Rodríguez & Arita (2004) found that beta diversity of North American mammals in relation to distance decreased with increasing latitude among non-volant mammals, but not among bats. In contrast, Stevens & Willig (2002) documented a latitudinal gradient of beta diversity with distance for New World bat faunas, with the highest values at low latitudes.

In this study, we analyse North American mammalian faunas across five latitudinal zones covering North America north of Mexico (Fig. 1). We evaluate the contributions of climatic and physiographic variables and geographical distance to turnover of mammal species. Variation in beta diversity with respect to



Figure 1 Division of North America into five latitudinal zones.

environmental factors provides evidence for differences in habitat specialization, whereas variation in beta diversity with respect to geographical distance, independent of environmental conditions, suggests differences in dispersal limitation (Condit *et al.*, 2002; Svenning & Skov, 2004) as well as historically different geographical sources of species (such as Nearctic and Neotropical realms). Environmental conditions and distance also interact as potential influences, since environmental conditions usually differ more among sites at greater distances along latitudinal and longitudinal gradients in North America.

The major objectives of this study are to determine whether beta diversity of mammalian assemblages in North America varies systematically with latitude and to assess the relative importance of climatic and topographic variables as well as geographical distance in predicting the beta diversity of mammalian faunas. We evaluate eight climatic and two physiographic variables variables that have demonstrated significance in earlier studies of North American mammal diversity gradients (Currie, 1991; Badgley & Fox, 2000). Faunas are compared within latitudinal zones and the zones themselves are compared in order to characterize the latitudinal gradient in beta diversity. The primary analyses are done at the species level, but we illustrate beta diversity at the genus and family levels for comparison with other studies of extant mammals (e.g. Kaufman, 1995) and with patterns in the fossil record that are often analysed at higher taxonomic levels (e.g. Janis et al., 2002; Jernvall & Fortelius, 2002).

The scale of this analysis is a grid of equal-area quadrats, 241 km (150 miles) on a side, based on the grid used by Simpson (1964) in his classic study of the species density of North American

mammals. At this scale of analysis, differences among regions are characterized well by general macroclimate and physiography rather than more localized variables. North America (north of Mexico) has a broad longitudinal span from relatively low (c. 30°) to high (> 80°) latitude, transected by physiographic features oriented generally from north to south – the Appalachian Mountains in the east and the American Cordillera in the west. North of ~30° N, variation in continental area does not confound interpretations of richness-based metrics. Thus, North America north of Mexico is an ideal testing ground for the relationship between beta diversity and latitude.

In contrast to previous analyses of spatial turnover of North American mammals, this study explicitly evaluates distance and environmental conditions as predictors of faunal differences. We include all continental mammals, in contrast to studies separating bats from non-volant mammals, because north of 30° N, most mammalian faunas have fewer than 10 species of bats (Rodríguez & Arita, 2004). Also, beta diversity is quantified in a different way from other studies, permitting us to evaluate latitudinal and longitudinal trends. Variance partitioning reveals the unique as well as joint contributions of distance and environmental factors in predicting beta diversity. Because a full understanding of the latitudinal gradient in beta diversity cannot be achieved until many groups of organisms have been examined, our study adds to the literature and offers an opportunity for comparing the latitudinal gradient in beta diversity between plants (Qian & Ricklefs, 2007) and animals.

MATERIAL AND METHODS

Mammalian species richness and some environmental variables were taken from the data set of Badgley & Fox (2000). In their study, North America was divided into 388 equal-area quadrats (241 × 241 km) on a Lambert equal-area projection, and species lists for each quadrat were compiled from species range maps of Hall (1981) and more recent sources (see Badgley & Fox, 2000, for details of data compilation). Quadrats occupied by less than 25% land area, along coastal regions, were omitted. Since the analytical method involves evaluating beta diversity across latitudinal zones of comparable magnitude, we excluded quadrats located south of 30° N because the span of longitudes is quite small. We divided North America north of 30° N into five latitudinal zones (Fig. 1). Isopleths of temperature run subparallel with latitude but tend to shift northward from east to west, and this shift is more apparent at more northern latitudes. For example, the southern boundary of the boreal zone shifts over 15° of latitude from east to west (Barbour & Billings, 1999). We constructed latitudinal zones such that the differences in temperature between eastern and western parts of each zone were minimized while keeping the latitudinal spans comparable among zones. We divided the temperate region into three zones and the boreal and arctic region into two zones (Fig. 1). We excluded a few quadrats located in the arctic region in order to make the latitudinal span of the two northernmost zones consistent. For the three southernmost zones, the latitudinal span of each zone narrows eastwards, but this trend occurs in all

the latitudinal zones. Each quadrat was assigned to a zone according to its midpoint latitude. Figure 1 illustrates the latitudinal zones and the 313 quadrats included in our analyses.

For each latitudinal zone, we calculated a Jaccard index of faunal similarity for pairs of quadrats. We used the Jaccard index because it is commonly used in ecological analyses, including those of beta diversity (Nekola & White, 1999; Qian et al., 2005; Gaston et al., 2007; Qian & Ricklefs, 2007), and because we wanted to compare this analysis for North American mammals with that of Qian & Ricklefs (2007) for North American vascular plants, in which the Jaccard index was used. The Jaccard index (J) is defined as J = a/(a + b + c), where a is the number of taxa shared between two localities and b and c are the numbers of taxa unique to each locality (Legendre & Legendre, 1998); values of J range from 0 to 1. We calculated the Jaccard index for faunas from all pairs of quadrats within each zone in order to characterize the degree of turnover within each latitudinal zone. Calculations were done separately for species, genera and families based on the mammalian taxonomy of Wilson & Reeder (1993).

We regressed the natural logarithm of $J(\ln J)$ on spatial distance between quadrat pairs in each latitudinal zone and used the absolute value of slope of the lnJ-distance relationship as a measure of spatial beta diversity, or distance decay of similarity (Nekola & White, 1999). The distance between pairs of quadrats was measured between their centroids. Since n quadrats in a latitudinal zone result in n(n-1)/2 pairs of quadrats, taking n(n-1)/2 as the number of degrees of freedom can increase the rate of Type I errors because each quadrat is used in multiple comparisons, and thus the significance of a statistical test can be inflated. To address this problem, we used a permutation approach to adjust probabilities of statistical inference. The approach that we used to determine the significance level for each regression model was developed by Legendre et al. (1994) and was implemented in Permute! Version 3.4 (available online at http:// www.bio.umontreal.ca/casgrain/en/labo/permute/index.html). All P values were determined from 9999 permutations.

In order to evaluate the relationship between lnJ for species and differences in physiographic and climatic conditions among quadrats (environmental beta diversity), we compiled data for 10 environmental variables for each quadrat: mean elevation, topographic relief and eight climatic variables. Data for elevation (m) and relief (m) are from Badgley & Fox (2000). The eight climatic variables and their units of measurement are: (1) mean annual temperature (TEMP_{ann}, °C), (2) mean temperature of the coldest month (TEMP_{min}, °C), (3) mean temperature of the warmest month (TEMP_{max}, °C), (4) seasonality of temperature (TEMP_{max} - TEMP_{min} = TEMP_{seas}, °C), (5) mean annual precipitation (PPT_{ann}, mm), (6) seasonality of precipitation (wettest month precipitation minus driest month precipitation, PPT_{seas}, mm), (7) annual actual evapotranspiration (AET, mm), (8) annual potential evapotranspiration (PET, mm). These variables represent important aspects of climate and have high predictive power for species richness of mammals and plants over broad spatial scales (e.g. Currie, 1991; Badgley & Fox, 2000; Qian et al., 2005; Qian & Ricklefs, 2007). We obtained temperature and precipitation data from New et al. (1999) and evapotranspiration data from Ahn &

	Latitudinal zone						
	A	В	С	D	Е		
Total family richness	28	25	23	18	17		
Total genus richness	98	86	74	57	45		
Total species richness	288	252	162	108	81		
Mean species richness per quadrat	83.2	81.8	68.8	49.2	37.7		
SD of species richness per quadrat	22.9	17.5	8.0	7.0	6.8		
Total/mean species richness	3.46	3.08	2.35	2.20	2.15		
Mean lnJ for species	-1.104	-0.983	-0.629	-0.546	-0.642		
Variance of lnJ for species	0.335	0.234	0.079	0.079	0.094		
Slope of ln <i>J</i> for species versus distance	-0.681	-0.448	-0.224	-0.169	-0.165		
No. of quadrats included	50	59	58	78	68		

lnJ, natural logarithm of the Jaccard index.

Tateishi (1994). We performed a principal components analysis on the correlation matrix of the eight climatic variables in order to reduce the number of variables in further analyses and to account for collinearity among climatic variables (Tables 2, 3). In order to compare species turnover with environmental differences, we calculated the difference in elevation, relief and climate variables between pairs of quadrats. For the physiographic variables, differences were calculated from the raw data. For climatic variables, differences were calculated from the scores on each of the first four principal component (PC) axes. For each latitudinal zone, lnJ was regressed on the difference in PC scores for each pair of quadrats. For each regression model, the standardized regression coefficients indicated which PC axis made the greatest contribution to the regression model. A set of similar regression analyses was performed with the topographic variables as predictors of lnJ. A third set of regression analyses included both climatic (PC scores) and topographic variables as predictors of lnJ. These regression models were compared for each latitudinal zone (Table 4).

Finally, we conducted a series of partial regressions (Legendre & Legendre, 1998) to partition the variance in ln*J* into the components uniquely explained by spatial distance, uniquely explained by environmental difference, explained by none of these variables. In each partial regression, the spatial distance between quadrats was contrasted with a set of variables representing environmental difference, including the difference between scores of the first four PCs and differences in elevation and topographic relief for each pair of quadrats. Each latitudinal zone was analysed separately.

RESULTS

Latitudinal gradient in taxonomic richness and turnover in relation to spatial distance

Together the five latitudinal zones have 363 species of extant mammals, belonging to 28 families and 109 genera. Family,

genus and species richness in each latitudinal zone decreases substantially with increasing latitude (Table 1). While mean species richness per quadrat also decreases with latitude (Table 1), the decline is smaller than for total species richness. The richness ratio of Zone A to Zone E is 3.6 for entire zones and 2.2 for quadrats. Also, the ratio of total to mean species richness within each zone decreases from zones A to E (Table 1). These patterns imply that faunas at higher latitudes are more similar and that spatial turnover is lower at higher latitudes.

Beta diversity of mammalian faunas decreases with latitude at the species, genus and family levels, although the differences between latitudinal zones are much lower for genus and family than for species (Fig. 2). (Pairs of quadrats with similar faunas have low negative values of lnJ and pairs with dissimilar faunas have high negative values.) Several patterns emerge. First, the mean value of lnJ within a zone increases from -1.104 in Zone A to -0.546 in Zone D and then decreases to -0.642 in Zone E (Table 1). Thus, on average, faunas are less similar at lower latitudes than at higher latitudes, but the trend changes slightly in the northernmost zone. Second, the variance of lnJ within a zone decreases with increasing latitude from 0.335 in Zone A to 0.079 in zones C and D (Table 1). Zone E has a slightly higher value (0.094). The general pattern indicates that the more northerly zones contain quadrats that are more homogeneous in taxonomic composition. Third, the slope of the relationship between lnJ and distance for all pairs of quadrats within each zone (Fig. 2) ranges from -0.681 in Zone A to -0.165 in Zone E, with monotonically declining slope values in between. The slopes are fairly shallow for zones C, D and E and substantially steeper for zones A and B (Fig. 3a), suggesting that a major decrease in spatial turnover with distance occurs between zones B and C. In addition, lnJ reaches much lower values for zones A and B than for zones C-E. Thus, faunas differ more in relation to longitude for zones at lower latitudes. Regression models using the spatial distance separating pairs of quadrats to predict lnJ for species explained from 82.8% to 49.8% of the variance in lnJ in zones A to E, respectively (Table 4).



Figure 2 Relationship between the natural logarithm of the Jaccard index of similarity ($\ln J$) and geographical distance for each of three taxonomic levels (family, genus and species) within each of five latitudinal zones (A to E). Slopes (*b*) were estimated from linear least square regressions. All the regressions were significant at *P* < 0.001.

Species turnover in relation to environmental variables

In the principal components analysis, the climatic variables are well represented by the first four PC axes, which account for 97.9% of the variance (Table 2). The first axis (PC 1) contrasts quadrats having high values of mean annual temperature, monthly minimum temperature, monthly maximum temperature and both potential and actual evapotranspiration with quadrats



Figure 3 Relationship between beta diversity at the species level, measured as the absolute value of slope of ln*J* in relation to distance from Fig. 2, and (a) mean latitude, (b) mean annual temperature, (c) mean PC 1 score (first principal component of climatic data) and (d) mean species richness per quadrat of each latitudinal zone.

having high seasonality of temperature. This climatic gradient is strongly correlated with latitude (as in Badgley & Fox, 2000). The mean value of scores on PC 1 for quadrats in each zone increases from Zone A to Zone E, and the variation among quadrats tends to decrease from Zone A to Zone E (Table 3). The second axis (PC 2) contrasts quadrats with high versus low values of annual precipitation and seasonality of precipitation (Table 2). The mean value of scores on PC 2 shows a weak latitudinal gradient, and the variation among quadrats tends to decrease from zones A to E (Table 3). PC 3 highlights climatic seasonality by contrasting quadrats having high seasonality of temperature and high annual actual evapotranspiration with quadrats having high seasonality of precipitation (Table 2). PC 4 represents additional variation in seasonality of temperature and precipitation. Neither the mean score on PC 3 and PC 4 nor variation in these scores covaries with latitude (Table 3). The variation (standard deviation) in guadrat scores on PC 1 and PC 2 within latitudinal zones decreases from Zone A to Zone E (Table 3). Since these axes represent most (83%) of the variance in the eight climatic variables (Table 2), this trend indicates that the overall climatic variation among quadrats from each latitudinal zone decreases from lower to higher latitudes.

Multiple regression of $\ln J$ on the paired differences in scores on PC 1 to PC 4 shows different relationships for different latitudinal zones. The difference in PC scores between pairs of quadrats is a measure of difference in current climatic conditions for those quadrats. The variance in $\ln J$ explained by the differences in climatic conditions decreases with increasing latitude from 81.5% to 36.2% (Table 4), and the climatic differences contributing

most to the regression model for each latitudinal zone vary with latitude (Table S1 in Supporting Information). The differences in PC scores explain less variance in ln*J* at higher latitudes.

Regression models using topographic relief and elevation also vary among different latitudinal zones. From low to high latitudes, the amount of variance explained in multiple regression of ln*J* on the differences in relief and elevation between quadrats decreases from 63.1% for Zone A to 5.6% for Zone E (Table 4). In most latitudinal zones, differences in topographic relief contribute more to the prediction of ln*J* than do differences in elevation (Table S1 in Supporting Information). The decline in explained variance with increasing latitude covaries with the decrease in elevation of vegetation zones at higher latitudes. The latitudinal timberline in North America lies near the boundary between zones D and E. Thus, for these latitudinal zones, differences in elevation and relief do not correspond to substantial changes in life zones.

Regression models combining climatic and topographic variables show a slight improvement in the amount of variance explained in $\ln J$ for each latitudinal zone, as well as a latitudinal gradient in the amount of variance explained (Table 4). The differences in PC scores and topographic variables together explain from 86.6% to 40.1% of the variance in $\ln J$ for zones A to E (Table 4). Over the five latitudinal zones, different climatic and physiographic conditions make substantial contributions to the prediction of beta diversity. The northward trend of decreasing variance explained in $\ln J$ by climatic and topographic variables parallels the latitudinal trend of decreasing variation in these environmental variables (Tables 5 & S2).

the correlation matrix.							
	PC 1	PC 2	PC 3	PC 4			
Eigenvalue	5.32	1.33	0.70	0.48			
Per cent of variance	66.54	16.56	8.75	6.00			
Cumulative per cent	66.54	83.10	91.85	97.85			
of variance							
Correlation with PC axis							
TEMP _{ann}	-0.97	0.19	0.01	-0.07			
TEMP _{min}	-0.96	0.18	0.18	0.09			
TEMP _{max}	-0.88	0.28	-0.19	-0.29			
TEMP _{seas}	0.82	-0.06	-0.43	-0.37			
PPT _{ann}	-0.61	-0.68	-0.23	0.30			
PPT _{seas}	-0.35	-0.75	0.41	-0.38			
AET	-0.80	-0.29	-0.48	0.01			
PET	-0.94	0.25	0.03	-0.12			

Table 2 Results of principal components (PC) analysis of climatic variables for quadrats (n = 313) in five latitudinal zones, based on the correlation matrix.

Climatic variables are: mean annual temperature (TEMP_{ann}, °C), mean temperature of the coldest month (TEMP_{min}, °C), mean temperature of the warmest month (TEMP_{max}, °C), seasonality of temperature (TEMP_{max} – TEMP_{min} = TEMP_{seas}, °C), annual precipitation (PPT_{ann}, mm), seasonality of precipitation (wettest month precipitation minus driest month precipitation, PPT_{seas}, mm), annual actual evapotranspiration (AET, mm), annual potential evapotranspiration (PET, mm).

When spatial distance, climatic conditions and topographic variables were simultaneously included in a regression model, the amount of variance explained in ln*J* improved substantially across all latitudinal zones (Table 4). These regression models explain 93.5% of the variance in ln*J* for Zone A and 57.4% for Zone E, with intermediate values for zones B, C and D (Table 4).

Variance partitioning among distance and environmental variables

The partitioning of variance explained by spatial distance alone, by environmental variables (climate and topography) alone and by distance and environmental variables jointly varies across latitudinal zones (Fig. 4). For Zone A, the greatest amount of variance (75.9%) is explained jointly by distance and environmental variables, while environmental variables alone explain





Latitudinal zone

Figure 4 The proportion of variance in ln-transformed Jaccard index of similarity explained by geographic distance and environmental difference. Environmental variables included difference between scores on the first four PC axes (principal components of eight climatic variables), difference in relief and difference in elevation between pairs of quadrats. Variance was partitioned among distance alone, difference in environmental variables alone and distance and environmental variables jointly. The total amount of variance explained in each latitudinal zone is given in Table 4 for the regression model combining spatial and environmental variables.

somewhat greater variance than distance alone. For Zone B, distance and environmental conditions jointly explain 63.2% of the variance and distance alone explains greater variance than environmental conditions alone. For Zone C, 44.0% of the explained variance is due to distance and environmental conditions jointly, and the amount explained by distance exceeds the amount explained by environmental conditions alone. For Zone D, distance and environmental conditions jointly explain 46.2% of the variance, distance alone explains 29.3% of the variance and environmental conditions jointly explain 32.5% of the variance and distance alone explains 17.3% of the variance, whereas environmental conditions alone explain yut 7.6% of the variance.

Table 3 Mean and standard deviation of principal component (PC) scores within each latitudinal zone.

Zone n		PC 1	PC 2			PC 3		PC 4	
	n	Mean	SD	Mean	SD	Mean	SD	Mean	SD
A	50	-3.202	1.280	0.881	1.460	0.157	0.974	-0.168	0.544
В	59	-1.770	1.229	0.183	1.469	0.059	1.197	0.021	0.776
С	58	-0.420	0.828	-0.277	1.228	0.037	0.935	0.118	0.833
D	78	1.283	0.827	-0.383	0.494	-0.072	0.559	-0.029	0.672
Е	68	2.777	0.832	-0.132	0.578	-0.116	0.438	0.038	0.598

n, number of quadrats included.

Table 4 Amount of variation (coefficient of determination × 100) in natural logarithm of the Jaccard index (ln*J*) explained by different sets of explanatory variables in multiple regression models. All coefficients of determination are significant at P < 0.001. Climate is represented by the first four principal components from Table 2; topography includes relief and elevation.

	Latituc	linal zone	2		
Explanatory variable	A	В	С	D	Е
Spatial distance	82.8	79.6	62.1	75.5	49.8
Climate	81.5	55.4	44.5	46.7	36.2
Topography	63.1	53.1	41.1	13.2	5.6
Climate and topography	86.6	74.1	57.2	48.9	40.1
Spatial distance, climate and topography	93.5	90.5	75.3	78.2	57.4

General latitudinal trends emerge from Fig. 4 but the trends are not consistently monotonic with latitude. The role of spatial distance alone increases in relation to the distances traversed by each latitudinal zone; Zone D traverses the greatest distance (about 6000 km, see Fig. 2), and distance alone explains more of the variance (29.3%) in Zone D than in any other zone. Distance and differences in environmental variables jointly explain more of the variance in lnJ at lower latitudes, rising from less than 50% in zones C-E to more than 60% in zones A and B. Differences in the environmental variables alone explain less than 10% of the variance in lnJ in zones D and E and more than 10% of the variance in zones A-C. From Table 4, it is evident that regression models based on distance alone or on differences in environmental conditions alone predict relatively large amounts of the variance in lnJ. But distance and differences in environmental conditions are strongly correlated; when both are included in a regression model their joint contribution dominates the explained variance.

DISCUSSION

The beta-diversity gradient

For the area of North America analysed, mammalian faunas show a decrease in zonal beta diversity with increasing latitude.

This trend is supported by the systematic shallowing of slope with latitude in the distance-decay plots in Figs 2 and 3(a) and the increase in mean ln*J* of zones with latitude (Table 1). At the levels of family and genus, the distance-decay plots reach an asymptote, reflecting the broad geographical distribution of most families and many genera (Fig. 2). The decrease in slope of ln*J* with latitude (Fig. 3a) has the same pattern as the decrease in slope of ln*J* with temperature and PC 1 for each latitudinal zone (Fig. 3b,c). This similarity indicates that 'latitude' is merely a proxy for underlying environmental gradients, including several measures of annual and seasonal temperature (Badgley & Fox, 2000; Hawkins & Diniz-Filho, 2004). Beta diversity is also strongly correlated with mean species richness per quadrat of each latitudinal zone (Fig. 3d).

Over most of North America, mammalian faunas at the species level become more dissimilar with greater distance apart. Zones A and B both show a steep decay of similarity with increasing distance. These temperate zones traverse considerable climatic and topographic heterogeneity from east to west. Mammalian faunas in these zones show higher species richness and ecological diversity and smaller geographical ranges west of the Great Plains (Pagel et al., 1991; Badgley & Fox, 2000). Zone C is the only zone to show an asymptote in lnJ. The lowest values of lnJ are about -1.2 in Zone C, whereas the lowest values of lnJ reach more negative values in the other four zones. Zone C spans the transition from temperate to boreal regions. The geographical ranges of North American mammals become substantially larger from Zone C northward. South of c. 48° N, all species ranges comprise less than 20% of the area of North America, but north of c. 48° N, many species ranges occupy more than 20% of the land area of North America (Fig. 5 in Pagel et al., 1991). This transition implies an increase in similarity among faunas of Zone C compared with those in zones A and B, as documented.

Longitudinally, zones D and E span greater maximum distances than do the more southerly zones. Both zones have faunas with greater dissimilarity than the most dissimilar faunas of Zone C. Zones D and E both traverse areas of Alaska that were unglaciated during the last ice age and supported a Beringian mammal fauna. After deglaciation, many Beringian species dispersed across the formerly glaciated regions, as immigrants also moved northward. A few Beringian species show little southern expansion after deglaciation, while the distribution of permafrost has limited the

 Table 5
 Mean and standard deviation of relief (m), mean elevation (m), mean annual temperature ($^{\circ}$ C) and annual precipitation (mm) of the quadrats in each latitudinal zone. Table S2 lists the mean and standard deviation of additional climatic variables.

Zone	n	Relief		Elevation		Temperature		Precipitation	
		Mean	SD	Mean	SD	Mean	SD	Mean	SD
A	50	1469.9	1316.5	871.3	784.1	16.3	3.8	722.1	489.0
В	59	1504.4	1177.4	1058.6	831.7	9.9	3.6	759.4	383.2
С	58	1232.8	1153.2	770.3	513.5	4.9	2.5	755.6	336.5
D	78	1238.1	1511.5	515.7	337.1	-1.2	2.8	622.4	269.3
Е	68	777.8	748.8	384.2	249.4	-7.4	3.2	424.8	272.4

n, number of quadrats included.

expansion of a number of northward immigrants, especially of rodents (Youngman, 1975). This combination of environmental history and modern environmental conditions may explain why more dissimilar faunas occur over the largest distances in zones D and E than in Zone C. Over shorter distances, Zone C shows greater dissimilarity among faunas than zones D and E.

Spatial distance and physical aspects of the environment, separately or together, predict a greater fraction of the variance in turnover for zones at lower latitude than higher latitude. For Zone A, differences in climatic and topographic variables predict more variance in ln*I* than does spatial distance alone (Table 4). For zones B-D, spatial distance predicts more of the variance in lnJ than do differences in climate and topography. However, when distance and environmental variables are evaluated simultaneously by variance partitioning (Fig. 4), the variance in lnJ explained jointly by distance and environmental conditions exceeds that explained by distance alone or environmental conditions alone. Unmeasured factors have an increasingly large influence on beta diversity at higher latitudes. The drop in explained variance (Table 4, Fig. 4) is especially large between zones A and B and zones C-E, with a large drop also between Zone D and Zone E. Zones C-E cover the formerly glaciated region of North America. Time since deglaciation may influence beta diversity of zones C-E in ways that are not directly correlated with the variables included here. For example, Zone E contains much of the area of continuous permafrost (Brown, 1960), which limits species distributions, especially those of burrowers, in the high Arctic.

Beta diversity decreases little from Zone D to Zone E (the absolute slope being 0.169 for Zone D and 0.165 for Zone E) and the absolute mean of lnJ for Zone E is slightly higher than those for zones C and D (Table 1). In Fig. 2, a cluster of quadrats separated by relatively short distances has lower values of lnJ for both species and genera. This pattern does not occur in the other four zones (Fig. 2). Several factors may contribute to the patterns of Zone E. First, much of the westernmost area of Zone E was ice-free during the last glacial advance, as discussed above, and was a refugium for plants and animals. In contrast, the rest of Zone E was covered by an ice sheet for thousands of years. Some species from Beringia may have encountered barriers to dispersal into the eastern part of Zone E, resulting in quite dissimilar faunas at the two ends of the zone. Second, Hudson Bay separates Zone E into two parts, and this separation may have accentuated the faunal differences in pairs of quadrats from either side. Third, some species of Zone E are boreal species. These species are usually widely distributed in the boreal zone, but their ranges may have penetrated into only limited areas of Zone E, with different species ranges penetrating into different longitudinal regions of Zone E, resulting in little or no overlap of their ranges in Zone E. As a result, mammalian faunas separated by a short distance may be quite dissimilar in species composition. For areas with low species richness, differences of a few species in different quadrats may lead to a large dissimilarity (high negative lnJ) among the faunas, resulting in high species turnover in some species-poor areas (e.g. Gaston et al., 2007).

Comparison with other studies of beta diversity

Our results are consistent with those of earlier studies of North American mammal faunas, using different analytical approaches. Kaufman & Willig (1998) and Rodríguez & Arita (2004) documented higher turnover versus distance of non-volant mammals at lower latitudes of North America. In the present study, bats contribute to the beta-diversity gradient across zones A–E. Although Pagel *et al.* (1991) did not document turnover directly, their analysis of mammalian geographical range size in relation to latitude and longitude implies greater turnover at lower latitudes and western longitudes in North America. While previous studies, including the aforementioned ones, have documented species turnover of North American mammals, the present study has for the time demonstrated the contributions of distance and environmental conditions separately and jointly to spatial turnover of North American mammals.

In a similarly designed analysis of vascular plants, Qian & Ricklefs (2007) found that beta diversity for vascular-plant assemblages decreases strongly with latitude. Regression models that included distance and climatic variables also explained a high proportion of the variance in lnJ; the amount of explained variance showed a different relationship with latitude than in this study. For their Zone D (approximately similar to Zone D in this study), spatial distance explained most of the variance in lnJ, whereas for mammals, distance and environmental factors jointly explain more variance than distance and environmental variables do separately (Fig. 4). Both studies demonstrate a decline in explained variance at higher latitudes. These similarities and differences suggest that mammal faunas and vascular-plant floras show broadly similar patterns of beta diversity over North America but that the spatial and environmental determinants of the pattern differ somewhat between these two groups.

Storch *et al.* (2005) demonstrated that the slope of the species– area relationship (SAR) for birds is shallower in areas of higher levels of available energy. A higher slope of the SAR reflects a faster accumulation of species with increasing sample area, and thus a greater rate of species turnover. Hence, their study implies that beta diversity is lower in areas with higher levels of available energy. In contrast, our study demonstrates that species turnover of mammals is greater in areas with higher ambient energy and decreases monotonically with decreasing ambient energy (measured as mean annual temperature; Fig. 3b).

Meta-analyses of beta diversity (e.g. Koleff *et al.*, 2003; Soininen *et al.*, 2007) have shown that beta-diversity patterns vary in relation to the spatial scale of the study, the metrics used to characterize turnover, the taxonomic group, major environment (marine versus terrestrial), as well as the specific region or continent. Comparisons within major taxonomic groups (e.g. montane and boreal floras, Nekola & White, 1999; North American mammals, Rodríguez & Arita, 2004) have also shown that beta-diversity patterns vary among different functional groups (e.g. volant versus non-volant mammals). In the present study, climatic and topographic features in combination with distance explain much of the variance in beta diversity for mammalian faunas in latitudinal zones. Turnover is also correlated with species richness (Fig. 3).

Insights about mechanisms

Several covarying properties of mammalian faunas or environments provide insights into possible mechanisms underlying these patterns of spatial turnover. (1) Geographical range size. The increase in geographical range size with latitude, or Rapoport's rule, for a group is correlated with the latitudinal gradient in beta diversity for North American mammals. Pagel et al. (1991) documented a decrease in geographical range size at lower latitudes and higher longitudes for North American mammals. Together these trends locate the highest beta diversity in the south-western part of the area analysed - a region that includes the Colorado Plateau, the Great Basin and California. (2) Ecological structure of mammalian faunas. If geographical range size varies in relation to species body size or trophic habit, then geographical gradients in the ecological structure of mammalian faunas may also contribute to the continental pattern of beta diversity. The ecological structure of North American mammalian faunas shows considerable geographical variation (Badgley & Fox, 2000): frugivores, aerial insectivores and terrestrial invertivores of small body size $(\leq 100 \text{ g})$ dominate faunas at lower latitudes, whereas herbivorous and carnivorous species of larger body size are more prevalent at high latitudes. Granivores and herbivores of intermediate size are prevalent in desert and montane habitats of the west. (3) Environmental history. More than half of the area analysed was covered by massive ice sheets only 14,000 years ago. Areas with a much shorter duration of occupancy may exhibit lower species density and different patterns of spatial turnover. For North American mammals, the time since deglaciation has a measurable influence on extant species density in the formerly glaciated region (Hawkins & Porter, 2003), implying a lag in recolonization. Furthermore, north-western Canada and eastern Alaska were recolonized by species from both the south and the west (Beringia), resulting in distinctive modern faunas unique to this region.

These covarying patterns suggest mechanistic hypotheses to elaborate and test. Ecological mechanisms include greater accommodation of species per unit area in regions with high topographic and resource heterogeneity; finer partitioning of resources among species at lower latitudes, as documented for African mammals (Hernández Fernández & Vrba, 2005) and herbivorous insects (Dyer et al., 2007); narrower environmental tolerances of species in habitats with lower climatic seasonality, resulting in lower dispersal ability (e.g. Janzen, 1967); and the longevity of viable ecosystems for species to invade and inhabit. Evolutionary mechanisms include higher speciation rates either in more topographically complex landscapes or at lower latitude more generally (e.g. Rohde, 1992); higher extinction rates in areas of high environmental stress and massive ecosystem disturbance (from glacial cycles); and the long-term evolutionary interactions of mammalian faunas from different, historically separate regions (Simpson, 1964). Our results indicate that the combination of spatial distance and environmental factors (climate, topography) has more explanatory power than either physical dimension alone in predicting beta diversity. This combination is consistent with most of the mechanisms proposed above, and further studies are needed to evaluate the contributions of these different mechanisms.

CONCLUSION

The latitudinal gradient in beta diversity of North American mammals corresponds to a macroclimatic gradient of decreasing mean annual temperature and increasing seasonality of temperature from south to north. Spatial distance, differences in climatic conditions and differences in topography predict a substantial amount of the variation in beta diversity within latitudinal zones, and this predictive power declines towards higher latitudes. Most of the variance in spatial turnover is explained by distance and environmental differences jointly rather than distance, climate or topography separately. Faunas at the highest latitudes (Zone E) show slightly greater spatial turnover than do faunas from the adjacent zone to the south. This deviation from the primary latitudinal gradient in beta diversity probably reflects different post-glacial sources for recolonizing the formerly glaciated region at high latitudes and environmental barriers that limit the ranges of a few species at both the eastern and western ends of this broad latitudinal zone. The high predictive power of geographical distance, climatic conditions and topography on spatial turnover could result from the direct effects of physical limiting factors or from ecological and evolutionary processes that are also influenced by the geographical template.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Table S1 Coefficients and standardized coefficients of variablesused in five regression models for each latitudinal zone.

Table S2Average values and standard deviations of climate datafor the quadrats in each latitudinal zone.

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BIOSKETCHES

Hong Qian is interested in searching for and interpreting large-scale patterns and processes of biodiversity and biogeography. In particular, he is interested in understanding the relative roles of historical and modern factors in determining the patterns in biodiversity.

Catherine Badgley is interested in palaeoecology of mammals, taphonomy, Miocene mammals of Indo-Pakistan and biogeography.

David L. Fox uses stable isotopes of biogenic and sedimentary materials to answer questions in palaeobiology and palaeoclimatology, analyses the role of stratigraphic data in phylogenetic analysis through the use of computer simulated evolutionary histories, and studies the ecological biogeography of modern mammals in relation to the spatial variation in climate.

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