Landscape Genetics of the American Badger: Understanding Challenges in Elusive Species

Elizabeth Marie Kierepka
University of Wisconsin-Milwaukee

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LANDSCAPE GENETICS OF THE AMERICAN BADGER: UNDERSTANDING CHALLENGES IN ELUSIVE SPECIES

by

Elizabeth M Kierepka

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ABSTRACT

LANDSCAPE GENETICS OF THE AMERICAN BADGER: UNDERSTANDING CHALLENGES IN ELUSIVE SPECIES

by

Elizabeth M Kierepka

The University of Wisconsin-Milwaukee, 2014

Under the Supervision of Professor Emily Latch

American badgers are one of the most poorly understood carnivores in North America due to their highly elusive, semifossorial lifestyle. To complicate understanding their biology further, badgers possess life history characteristics that predict radically different responses to habitat heterogeneity. In particular, they are considered grassland specialists, so their movement and population viability could be highly dependent on grassland habitats. Badgers are also highly mobile, which suggests they experience high gene flow. Predicting how these life history traits impact gene flow, however, is difficult based on the high diversity in responses to landscape heterogeneity among carnivores. To assess how landscape heterogeneity affects gene flow in badgers, my dissertation contains three chapters. In the first chapter, I assessed performance of individual-based landscape genetic methods to identify statistics that would be most appropriate for elusive species like badgers. Once I identified methods that would be appropriate for badgers, I combined both individual-based simulations and landscape genetic methods in my second chapter to assess how landscape heterogeneity in Wisconsin affects gene flow in a protected population of badgers. My final chapter investigated how historical and
contemporary changes in grasslands has affected genetic variation in badgers across their
North American range.
TABLE OF CONTENTS

Chapter 1 - Performance of partial statistics in individual-based landscape genetics
Introduction.........................................................................................................................1
Methods.................................................................................................................................7
   Generation of Landscapes.................................................................................................7
   Gene Flow Simulations.....................................................................................................10
   Statistical Analysis: Mantel Tests....................................................................................10
   Landscape Variable Derivation: Ordination.................................................................11
   Statistical Analysis: Ordination.......................................................................................12
   Statistical Performance.....................................................................................................14
Results..................................................................................................................................15
Discussion............................................................................................................................18
   Performance of Mantel Tests and Ordination Techniques............................................19
   Influence of Landscape Composition on Partial Statistics..........................................23
   Conclusions.......................................................................................................................25
Literature Cited....................................................................................................................35

Chapter 2 - Fine-scale landscape genetics of the American badger (Taxidea taxus):
disentangling landscape effects and sampling artifacts in an elusive species
Introduction............................................................................................................................40
Methods.................................................................................................................................44
   Study Area.........................................................................................................................45
   Laboratory Methods.........................................................................................................46
   Statistical Analysis..........................................................................................................47
Results..................................................................................................................................52
Discussion............................................................................................................................56
Literature Cited....................................................................................................................77

Chapter 3 - Range-wide gene flow in a widespread, specialist carnivore, the
American badger (Taxidea taxus)
Introduction.........................................................................................................................84
Methods.................................................................................................................................89
   Sample Collection..............................................................................................................89
   Laboratory Methods.........................................................................................................89
   Historical Gene Flow.......................................................................................................91
   Contemporary Gene Flow...............................................................................................94
Results..................................................................................................................................100
   Historical Gene Flow.......................................................................................................100
   Contemporary Gene Flow...............................................................................................102
Discussion............................................................................................................................105
   Historical Gene Flow.......................................................................................................106
   Contemporary Gene Flow...............................................................................................109
   Conclusions.......................................................................................................................113
Literature Cited....................................................................................................................134
LIST OF FIGURES

Figure 1.1. Multifractal landscapes generated for this study where 400 individuals were randomly placed on each landscape. Each landscape consisted of two landscape categories: suitable (light grey) and matrix (black) habitat. The amount of suitable habitat (P) varied from 20% to 80% with five replicates within each P category. Pixels of suitable habitat were given a resistance value of 1 whereas matrix habitat was given one of four values (5, 10, 25, and 50) across simulations for a total of 80 simulated landscapes................................................................. 28

Figure 1.2. Frequency of type I errors in partial Mantel tests, dbRDA, and RDA in simulated landscapes. IBR was incorrectly detected most often in dbRDA tests, and least often in partial Mantel tests. Error bars correspond to standard errors calculated across the 1,600 simulated populations................................................................. 29

Figure 1.3. Frequency of type II errors in partial Mantel tests (a), dbRDA (b), and RDA (c) in simulated landscapes. Five replicate landscapes were simulated for each percent suitable habitat (20%, 40%, 60%, 80%), at four matrix habitat resistance levels (R5 – light gray bars; R10 – gray bars; R25 – dark gray bars; R50 – striped bars). One hundred populations were simulated for each of the 80 total landscapes. Type II errors were calculated as the number of populations (out of 100) in which IBR was not detected in our simulated landscapes, and corresponding error bars indicate standard errors calculated across five replicate landscapes. In general, type II errors increased with amount of suitable habitat but were similar across resistance levels for all tests........ 30

Figure 2.1. Predictor variables for each georeferenced badger describe its location according to three landscape features (Wisconsin River, level III Ecoregion, and land cover). For the Wisconsin River (blue line), each individual was coded as either east or west of the river (a). All badger locations fell within one of four level III Ecoregions within Wisconsin (b): Driftless Area (DA), Northcentral Hardwood Forests (NCHF), Northern Lakes and Forest (NLF), and Southwestern Wisconsin Till Plain (SWTP). Land cover data for each badger was calculated as percent land cover within a circular buffer surrounding each badger’s location (most predominant land cover class given by point’s color; c)........................................................................................................ 63

Figure 2.2. Spatially lagged scores for the first two sPCA axes for Wisconsin badgers. Scores from Axis 1 (a) and 2 (b) were correlated with the Wisconsin River and percent agriculture respectively. Dark colors (black and dark grey) represent negative sPCA scores while positive values are light in color (white and light grey). More extreme values in sPCA axes are displayed with larger squares................................. 66

Figure 2.3. Percent of false significant tests (type I errors) for the Wisconsin River, level III Ecoregions, and Agriculture calculated in 100 simulated IBD populations. Both partial RDAs and spatially lagged regressions (sReg) for each axis separately incorrectly identified the Wisconsin River and level III Ecoregions as influences on gene flow more often than Agriculture....................... 68
Figure 3.1. Sampling locations and subspecific designations of $n = 917$ badgers. Designations are morphological subspecies as defined in Long (1972). .................................116

Figure 3.2. Haplotype network for 95 d-loop haplotypes detected in $n = 785$ badgers. Each circle represents a single haplotype and the size of the circle corresponds to its frequency. Each line represents a single mutational step, and black circles indicate missing genotypes. The network was color-coded by subspecies (a) or inferred subspecies (b). The most common haplotypes (H1, H2, H3, and H4) are labeled......117

Figure 3.3. Mismatch distributions for the total dataset (a), subspecies (b-e), and SAMOVA groups (f-g). Observed distributions are represented as gray bars and expected distributions under rapid demographic expansion are black solid lines. 95% confidence intervals for the expected distribution are given as dotted black lines. All groups followed expected patterns under rapid demographic expansion (all $p < 0.008$) except for the eastern SAMOVA group ($p < 0.002$).................................................................................................118

Figure 3.4. Distribution of three genetic clusters inferred in BAPS 5. The three clusters (West: black; Wisconsin: grey, and LP: white) are largely separated by two major water barriers: the Lake Michigan and the Mississippi River. The Mississippi River is represented by the bold blue line................................................................................................. 119

Figure 3.5. Plot of first two PCA axes when grouped by large aquatic barriers (a) or subspecies (b). Each point represents a single badger’s genotype that is color-coded according to either the aquatic barriers (east of Lake Michigan, west of Lake Michigan/East of Mississippi River, and west of the Mississippi River) or subspecies (T. t. berlandieri, jacksoni, jeffersonii, or taxus). Ovals around points correspond to 95% confidence ellipses that denote where 95% of individuals within each group occur. Each square on the background grid corresponds to 0.5................................................................................................. 120

Figure 3.6. Results of the sPCA analysis. Eigenvalue plots (a) and the screeplot (b) indicate that the first two axes are most important in explaining genetic variation. The screeplot plots the relationship between variance explained by each axis and spatial autocorrelation (Moran’s I) within each axis. Spatially lagged scores from Axes 1 (c) and 2 (d) depict two distinct patterns in genetic variation. Squares represent each individual where larger squares have stronger positive (white) or negative (black) values whereas smaller, grey (light grey = positive, dark grey = negative) squares are less strongly differentiated........................................................................................................................................... 121
LIST OF FIGURES (SUPPLEMENTARY)

Figure S2.1. Results of Structure analyses from \( K = 1 \) to \( K = 10 \). Maximum likelihoods (grey squares) and \( \Delta K \) values (black diamonds) occurred at \( K = 2 \) (a). Error bars correspond to standard deviations around each likelihood across five runs per \( K \). A heat map was constructed from averaged q-values for \( K = 2 \) based on inverse distance weighting (b). The heat map revealed no conspicuous barriers, but rather exhibited a gradient from northwest to southeast............................................................................ 69

Figure S2.2. Spatial autocorrelation for all badgers within Wisconsin. Dotted lines correspond to 95% confidence intervals derived from 1000 permutations of Rousset’s \( a \) within each 5 km distance class. Each point represents average Rousset’s \( a \) at a distance class, and those below the 95% confidence intervals (< 20 km) had lower genetic distance than expected by random mating.......................................................................................... 71

Figure S2.3. Eigenvalues (a) and screeplot of sPCA (b) displaying the variance explained and spatial autocorrelation within each sPCA axis. Axis 1 and 2 were retained for further analysis because they had the highest eigenvalues and explained the most variance within the dataset.................................................................................................................. 72

Figure S3.1. Geographic distribution of the four main haplotypes and those with the 26 bp deletion region that were not included within the mtDNA analyses. Rare haplotypes had a frequency of 10 or less individuals (dark grey circles). Haplotype 1 (red squares) is found throughout North America, but its frequency is the highest in the Upper Midwest. Haplotype 2 (blue circles) primarily occurs within northern latitudes, particularly in Montana, Canada, and North Dakota. Haplotype 3 (yellow diamonds) is one base pair away from Haplotype 1, and is chiefly located in the Lower Peninsula of Michigan and Ohio. Haplotype 4 (light blue triangles) is one mutation away from Haplotype 2, and almost always occurred west of the Rocky Mountains. Like Haplotype 4, individuals with the 26 bp deletion region (green squares) are primarily found west of the Rocky Mountains. The haplotype network of the 95 haplotypes that did not include the deletion region is provided where the four main haplotypes are colored................................................................. 123

Figure S3.2. Results of the overall clustering analysis within STRUCTURE. \( \Delta K \) indicated the most likely number of clusters was 2 as shown by the large peak at \( K = 2 \) within the plot (a). The spatial arrangement of q-values (b) indicated a general cline from east to west. We interpolated q-values using inverse-weight distance to create a genetic surface across our sampling area and individuals are colored according to their q-values. Those with higher q-values (large blue circles) were more strongly assigned to the eastern cluster.......................................................................................................................... 124

Figure S3.3. Results of the iterative runs of the two main clusters detected in the overall clustering analysis within STRUCTURE. The \( \Delta K \) plot for the eastern cluster (a) indicated further substructure whereas the plot for the western cluster (b) did not strongly support any additional genetic substructure. When mapped (c), the iterative run in the eastern cluster (n = 210 individuals) suggested the presence of two additional clusters that
separated the Lower Peninsula of Michigan from the rest of the individuals. Interpolated 
$q$-values, created via inverse-weight distance interpolation, revealed a strong divergence 
between the Lower Peninsula and Wisconsin (high assignment to the Lower Peninsula is 
denoted with large blue circles). .................................................................................... 125
LIST OF TABLES

Table 1.1. Summary of variables calculated for partial Mantel tests, dbRDA, and RDA analyses. Partial Mantel tests use pair-wise distances for all variables. dbRDA uses genetic distance as the response variable, and individualistic measures of ecological and geographic data as predictors. Genetic data for RDA was derived from a spatial principal components analysis (sPCA). All three tests had three indices of performance (accuracy, type I, and type II error rates) calculated based on 100 simulated populations per landscape.................................................................32

Table 1.2. Top six models that explained variation in partial Mantel rs (a), dbRDA pseudo F-ratios (b), and RDA pseudo F-ratios. Models are ranked according the lowest AIC value, and all those with a ΔAIC < 2.0 are considered top models (bold)............ 33

Table 1.3. Model averaged parameters for the top glm models in partial Mantel tests, dbRDA, and RDA. Only parameters that were shared between models are reported (i.e., models with an ΔAIC < 2.0). When 95% confidence intervals did not include zero, parameters were considered significant (bold)..........................................................34

Table 2.1. Locus-specific summary of genetic variation for n = 233 badgers in Wisconsin. Metrics included are the number of alleles per locus (A), observed heterozygosity (H₀), expected heterozygosity (Hₑ), inbreeding coefficients (F₁S), and 95% confidence intervals for F₁S for each locus. Asterisks indicate significant deviations from HWE at a corrected α = 0.012..............................................................................73

Table 2.2. Model selection results for spatially lagged regressions on sPCA Axes 1 and 2. Top models (ΔAIC < 2.0; in bold) for Axis 1 all included the Wisconsin River (River) whereas models for Axis 2 incorporated Agriculture (Ag). Ecoregion (Eco) was not significant for either axis following model averaging.................................74

Table 3.1. Results of SAMOVA for K = 2 where the first group contained the Lower Peninsula of Michigan, Indiana, and Ohio and the second group contained all other individuals. All variance components were significant after 1000 permutations (all p < 0.001)........................................................................................................127

Table 3.2. Diversity metrics for n = 785 badgers, for each of the four subspecies, and for the two SAMOVA groups. All diversity measures include standard errors.................128

Table 3.3. Mismatch statistics for the four subspecies and for the two SAMOVA groups. Fu’s Fs and Tajima’s D indicate demographic expansion when they are significantly negative whereas significant SSD and raggedness indices suggest deviations from patterns expected under demographic expansion. P-values for each test are given in parentheses; significant values (P<0.05) are in bold....................................................129

Table 3.4. Genetic diversity measures for three genetic clusters detected in BAPS. All three populations had significant heterozygote deficiencies (bold), and LP had lower
genetic diversity than WI and West (all \( t > 2.341, p < 0.02 \)). P-values are given in parentheses for \( F_{IS} \) values.

Table 3.5. Results from partial dbRDA analyses to evaluate the relationship between genetic distance and topographic barriers, geographic coordinates, and landscape variables. The final model after sequential selection included Latitude, Longitude, Lake Michigan, and the Mississippi River (bold), and together explained 10.11% of the total genetic variation in the dataset. Variance explained by each singular variable with all other variables partialled out are provided (% var).
LIST OF TABLES (SUPPLEMENTARY)

Table S2.1. Details of the microsatellite loci to be used in this study. Each multiplex set consisted of three loci. Locus amplification conditions used in this study are listed as footnotes..........................................................................................75

Table S3.1. Information about all the badgers, location, and source. Location corresponds to the state where badgers were sampled and for those that had small sample sizes (< 10 individuals), their SAMOVA group is listed in parentheses. All other SAMOVA groups are identical to their capture state or Canada. Sample sizes (N) correspond to the number of individuals provided by each organization.......................132
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CHAPTER 1: PERFORMANCE OF PARTIAL STATISTICS IN INDIVIDUAL-BASED LANDSCAPE GENETICS

Introduction

Classical population genetics and contemporary landscape genetics traditionally involve analyzing genetic differentiation within and among genetically discrete groups. Populations can be challenging to delineate in continuously distributed species that do not have clearly defined population boundaries. One approach for delineating cryptic population structure is to utilize Bayesian clustering algorithms (e.g., Pritchard et al. 2000; Guillot et al. 2005; Corander & Marttinen 2006) to define populations within a landscape. However, common patterns of spatial genetic structure such as isolation-by-distance (IBD; Wright 1943) or weak barriers to gene flow can lead to incorrect estimates of the number of genetically discrete populations across a landscape (Latch et al. 2006; Frantz et al. 2009; Schwartz & McKelvey 2009). In these situations, individual-based analytical approaches offer a viable alternative to population-based techniques. Because individual-based statistics do not require a priori definition of populations, they offer an appealing alternative approach to investigate a wide variety of fine-scale influences on gene flow.

Individual-based methods in landscape genetics have become particularly popular for differentiating between two models of gene flow across a heterogeneous landscape: isolation by distance, and isolation-by-resistance (IBR; McRae 2006). IBD occurs when geographic distance is positively correlated with genetic differentiation regardless of
landscape composition, and often serves as a null hypothesis in landscape genetic studies. Conversely, in IBR, landscape heterogeneity (e.g., habitat types, roads, rivers) influences gene flow. In order to evaluate the relative influence of habitat variables on gene flow, the landscape itself is parameterized using user-defined resistance values. These resistance values are assigned to each pixel on a raster map based on their hypothesized effect on gene flow. For example, Cushman et al. (2006) parameterized landscapes for black bears in Idaho such that anthropogenic structures like roads and agriculture inhibited gene flow (high resistance) and suitable forest habitats promoted gene flow (low resistance). Selection of appropriate resistance values requires *a priori* knowledge of relevant landscape features (for discussion see Zeller et al. 2012), so previous life history data such as telemetry (e.g., Cushman & Lewis 2010; Reding et al. 2013), expert opinion (e.g., O’Brien et al. 2006), or presence/absence data (e.g., Walpole et al. 2012) are often employed to derive meaningful resistance surfaces (raster maps where each pixel has a resistance value). Each parameterized resistance surface, therefore, represents a hypothesis about how landscape features affect gene flow. Hypothesis testing then involves quantifying correlations between landscape resistance and pair-wise measures of genetic differentiation to identify which landscape features, if any, explain patterns of genetic differentiation. One potential challenge to quantifying IBR is that landscape parameters of interest are often tightly correlated with geography. Autocorrelation in landscape and genetic variables can then result in spurious correlations in genetic analyses unless statistics can explicitly incorporate geographic covariates (Meirmans et al. 2012).
One of the primary remedies for spatial autocorrelation in IBR studies has been the use of partial statistics, in which the effects of geographic coordinates are separated from landscape data. Partial statistics can isolate IBR from IBD, and therefore, prevent spurious correlations observed in aspatial statistics (e.g., Cushman & Landguth 2010; Meirmans et al. 2012). Currently, the most prominent partial statistics in individual-based landscape genetics can be broadly classified as distance-based methods. Distance-based methods typically correlate pair-wise distance metrics of landscape resistance with genetic differentiation (termed genetic distance) while controlling for straight-line (Euclidean) distances between individuals. Numerous metrics exist for estimating genetic differentiation between pairs of individuals, where larger calculated values of genetic distance typically indicate higher levels of divergence. Pair-wise measures of cost to gene flow through a landscape between all pixels of the landscape (hereby called cost distance) are then calculated based on resistance surfaces. Cost distances are essentially the cumulative effect of traveling through pixels of different resistances between two individuals, irrespective of geographic distance, so like resistance values, higher cost distances represent stronger impediments to gene flow. Generally, once pair-wise distance measures are derived, distance-based partial statistics test for positive correlations between genetic and cost distances indicative of IBR. Although several distance-based methods can be adapted for genetic data, the partial Mantel test (Smouse et al. 1986) is unquestionably the most popular in landscape genetics.

Mantel tests quantify correlations between two or more matrices, and in landscape genetics, these matrices are typically composed of pair-wise genetic, Euclidean, and cost distances. Based on their widespread use in landscape genetics, simulation studies have
largely focused on identifying factors that affect performance of Mantel tests like resistance strength (i.e., values of pixels within raster maps; Cushman et al. 2012) and landscape configuration (Landguth et al. 2012). Resistance strength and landscape configuration have received attention in simulation studies because both can dramatically affect conclusions about gene flow (Cushman et al. 2012; Oyler-McCance et al. 2013). In general, Mantel tests perform best in highly fragmented landscapes with high amounts of resistant matrix habitat (Cushman et al. 2012). When resistance values of matrix habitat are much higher than suitable habitat (generally 5 to 10 times higher), partial Mantel tests detect inhibitory effects of habitat variables on gene flow (Cushman et al. 2012). Even when landscape configuration and resistance values are set to maximize statistical performance within simulated datasets, multiple authors have raised concerns about potential problems with Mantel tests including low power (Legendre & Fortín 2010; Graves et al. 2013) and high type I error rates (Balkenhol et al. 2009; Guillot & Rousset 2013). These studies have provided important caveats for empirical studies that rely on Mantel tests, and elucidate the need for alternative statistical approaches to complement results from Mantel tests and improve interpretation of empirical datasets.

Few other partial statistics have received the mass utilization as Mantel tests, but ordination techniques such as redundancy analysis (RDA) have recently been suggested as a viable alternative (e.g., Balkenhol et al. 2009; Legendre & Fortín 2010). Ordination techniques offer considerable flexibility as compared to Mantel tests because they do not require distance-based metrics and can overcome core assumptions of Mantel tests (e.g., linear relationships between variables). RDA is an unconstrained ordination technique where genetic data can be either genetic distances (distance-based redundancy analysis
[dbRDA]; Legendre and Anderson 1999) or individualistic measures of genetic variation like allele frequencies. Individualistic metrics for landscape and geographic variables (e.g., spatial coordinates, habitat type identity, precipitation measures, side of putative barrier) increase the diversity of explanatory variables that can be tested while negating the requirement for pairwise distance calculations that can result in losses of statistical power (Legendre & Fortín 2010). Also, ordination techniques like RDA and dbRDA can utilize transformations such as principal coordinates analysis (PCoA) to linearize genetic variables, thus removing any potential violations of linearity observed in Mantel tests (Graves et al. 2013).

Another potential advantage of ordination methods is that they provide measures of variance around parameter estimates, which allows for improved interpretation of results over Mantel tests that only provide a correlation coefficient and p-value (Legendre & Fortín 2010). Correlation coefficients from Mantel tests can be highly variable (Guillot & Rousset 2013; Graves et al. 2013), and without measures of variance, researchers may have trouble distinguishing between a type I error and the correct conclusion. In contrast, RDA provides statistics typical of ANOVAs including sum of squares, variance explained by each component, and measures of variance around F-ratios. These additional results generated by RDA provide more information upon which to base interpretations about whether and how particular landscape variables influence gene flow (i.e., partition variance among landscape variables) and the statistical validity of the entire model (Legendre & Fortín 2010). Given these characteristics – the ability to overcome linearity assumptions, flexibility in type and number of explanatory variables,
and statistical outputs – ordination methods offer potentially useful alternatives to Mantel tests.

Within individual-based genetics, simulation studies have evaluated the performance of Mantel tests under different scenarios that occur in empirical datasets (e.g., Cushman et al. 2012, 2013). Our goal was to use simulations to test performance of ordination techniques, a potentially viable alternative to Mantel tests for individual-based landscape genetic analyses. We quantified the performance of partial Mantel tests and ordination methods by assessing each statistic’s accuracy and explanatory power for detecting IBR (alternative hypothesis where landscape parameter was significant) while controlling for underlying patterns of IBD (null hypothesis of no effect of landscape heterogeneity). Simulations were designed such that spatial genetic structure was driven by IBD in all landscapes, and additionally included varying levels of resistance across the landscape (IBR). Given that RDA does not require quantification of pair-wise distances and tend to be more robust in population-based analyses (Balkenhol et al. 2009; Legendre & Fortin 2010), RDA was predicted to be more effective at detecting the effect of landscape resistance (i.e., costs to gene flow) than Mantel tests. In addition to comparing statistical approaches, we also designed simulations to assess the role of underlying landscape structure (habitat amount and configuration) and resistance strength of matrix habitat on statistical accuracy and explanatory power of IBR models. Both landscape structure and resistance strength have been identified as important predictors of accuracy in Mantel tests (Cushman et al. 2012), and we expected that these variables would likewise influence performance in ordination methods, though perhaps to a different
degree. Our findings are relevant to the larger field of work investigating optimal methods for detecting IBR in individual-based landscape genetics.

**Methods**

*Generation of Landscapes*

Landscapes in nature are composed of a mosaic of features and resources that each species can perceive as suitable or unsuitable based on their life history requirements. Many species utilize areas within a heterogeneous landscape in a non-random way, and preference for or avoidance of certain landscape variables is often assumed to promote or inhibit gene flow respectively (e.g., Shafer *et al.* 2012). Our simulated landscapes were composed of two habitat types: suitable and unsuitable (matrix) habitats. Heterogeneity of a landscape can be expressed using many different metrics, but we controlled two aspects of landscape configuration, level of fragmentation and proportion of suitable habitat, because these aspects have been shown to impact the strength of correlations between genetic differentiation and landscape resistance (Cushman *et al.* 2012). The level of fragmentation, in particular, seems to have direct impacts on both demographic (Revilla & Wiegand 2008) and genetic processes (Keyghobadi 2007; Bruggeman *et al.* 2010) in empirical and simulated datasets, so controlling fragmentation and proportion of habitat types should allow us to isolate the effects of different cost measures and resistance strength on partial statistics.

We manipulated habitat configuration of the simulated landscapes by creating artificial landscapes in QRULE (Gardner 1999) using the “Multifactual Random Map”
function. Multifactual random maps were created by defining the amount of aggregation of pixels or level of fragmentation (H) as well as the proportion of suitable habitat (P). We chose an H value of 0.50 with suitable habitat comprising either 20%, 40%, 60%, or 80% of the total landscape because extreme values in H or P in either direction are probably less realistic in nature or prevent meaningful levels of IBR or IBD within simulations. Only variation in P was tested within fractal landscapes because Cushman et al. (2012) found that configuration metrics (e.g., patch cohesion; Schumaker 1996; clumpy index of habitat aggregation; McGarigal et al. 2002, and correlation length of habitat; McGarigal et al. 2002) that explained genetic differentiation were highly correlated with P, but not with H. Therefore, varying P is likely a more effective way to create landscapes that result in detectable levels of genetic differentiation. Five replicate landscapes were generated within each P category (20%, 40%, 60%, or 80% suitable habitat) to evaluate variation in statistics, resulting in 20 independent landscape configurations (Figure 1.1).

We calculated three configuration metrics (patch cohesion, clumpy index of habitat aggregation, and correlation length of habitat) for all simulated landscapes. Configuration metrics have higher predictive power to explain how landscapes impact IBR than just suitable habitat alone (Cushman et al. 2012). Patch cohesion, a measure of habitat class aggregation, measures the physical connectedness of a focal habitat class as a percentage (McGarigal et al. 2002). Clumpy index of habitat aggregation (hereby called clumpy) also measures habitat class aggregation, but is not influenced by area of a focal habitat class (Neel et al. 2004). Clumpy ranges from -1 to 1 where -1 is completely disaggregated, 0 is random, and 1 is maximally clumped (McGarigal et al. 2002).
Finally, correlation length of habitat defines the average distance an individual could travel in a random direction without leaving suitable habitat when randomly placed in a habitat patch, making correlation length a measure of the extensiveness of suitable habitat (McGarigal et al. 2002).

By utilizing multiple landscapes from QRULE, we were able to isolate the impact of different resistance strengths in addition to suitable habitat amount and configuration on detection of IBR. Each simulated raster was 512 X 512 pixels (each pixel = 10 m x 10 m) where each pixel contained one of two cost values: suitable habitat was always set to 1 and the matrix was given a value of 1 (indicating IBD only), 5, 10, 25, or 50. Essentially, matrix habitat was parameterized such that high valued pixels impeded gene flow, and the higher the value, the stronger the inhibitory effect. In total, five landscapes (R1, R5, R10, R25, and R50) per habitat area category (total = 100 landscapes) were utilized for gene flow simulations.

The spatial arrangement (and geographic coordinates) of individuals was identical for each population to facilitate comparison among the five resistance surfaces. Four hundred individuals were randomly generated onto the landscape using the “Create Random Points” function in ArcMap. In addition to spatial coordinates for each individual, pair-wise cost matrices were calculated based on raster values using the “cost matrix” function found in the landscape genetics toolbox for ArcGIS (Etherington 2011). This tool sums the resistances of pixels along a Euclidean line connecting two individuals, and then produces a pair-wise cost distance matrix. Euclidean distances between all pairs of individuals were also calculated for Mantel tests using the “Euclidean Distance” function in the toolbox.
**Gene Flow Simulations**

Individual-based simulations were performed in the program CDPOP v. 1.3 (Landguth & Cushman 2010) using the cost matrices of the simulated landscapes. In total, 10,000 populations were simulated, 500 populations (5 replicates of 100 populations) for each of the 5 resistances (R1, R5, R10, R25, R50) across four habitat area categories (20%, 40%, 60%, 80% suitable habitat). Each simulation started with 400 individuals (200 male, 200 female) characterized at 10 polymorphic microsatellite loci (15 alleles per locus). Dispersal and mating movements occurred according to an inverse square distribution. An inverse square distribution describes a dispersal pattern in which it is far more common for individuals to remain near their birthplace than to disperse long distances. This dispersal distribution typifies many species in nature due to the risks associated with long-distance dispersal (Mayr 1963; Endler 1977). Furthermore, an inverse square dispersal distribution ensures an underlying IBD pattern in all simulated populations. Pair-wise cost distance matrices constrained both mating and dispersal where the maximum movement through suitable habitat was the length of the landscape (6120 m) and decreased according to resistance strength in matrix (R5 = 1224 m, R10 = 612 m, R25 = 244.8 m, R50 = 122.4 m). Mating was sexual with replacement, and each female could have a maximum of five offspring with an average of two. Generations were non-overlapping and results were saved after 500 generations. Statistics were then run on 400 sampled individuals.

**Statistical Analysis: Mantel Tests**
Mantel tests require pair-wise genetic distances, so we used Spagedi v. 1.3 (Hardy & Vekemans 2002) to calculate matrices of inter-individual genetic distances (Rousset’s $a$; Rousset 2000), for each simulated population. Matrices of Euclidean, cost, and genetic distances (Table 1.1) were then utilized for full and partial Mantel tests in the R package vegan (functions mantel and mantel.partial; Oksanen et al. 2013). Simple Mantel tests compared genetic distance to Euclidean distances, and partial Mantel tests compared genetic and cost distances while controlling for Euclidean distance. This procedure was repeated for each population across resistance value categories (R1, R5, R10, R25, R50) to quantify the ability of partial Mantel tests to separate the effects of cost and Euclidean distances as the relative effect of cost decreased. Statistical significance in all tests was assessed via Pearson’s correlation coefficients ($r$) after 9999 permutations.

Landscape Variable Derivation: Ordination

Ordination techniques can incorporate multiple explanatory variables into the analysis, but require individualistic measures of genetic diversity, geography, and costs (Table 1.1). One difficulty in adapting ordination techniques to individual-based landscape genetics is quantification of landscape resistance to gene flow typically relies on inter-individual distances. Multiple techniques exist to quantify cost distances (e.g., least cost path analysis; circuit theory; McRae and Beier 2007), but cost distance calculations often require substantial life history information to properly parameterize a landscape (Spear et al. 2010). Information on dispersal itself instead of habitat use is very difficult to obtain, so even with habitat use data, cost distances may not reflect the
actors that affect dispersal (Zeller et al. 2012). Therefore, we quantified habitat area around each individual to derive a landscape variable that does not depend on cost distances.

Ordination tests incorporated point estimates of habitat area surrounding each individual derived using following methodology. Circular buffers of 100, 200, and 300 m were initially drawn around each individual point using the “Buffer” function in ArcMap v. 10. Then, the area of matrix habitat was calculated within each buffer and expressed as a proportion (area of matrix/total area in buffer) for use the landscape predictor variable. Point estimates of landscape resistance do not require \textit{a priori} parameterization, which makes them an important complement to analyses that utilize cost distances derived from natural history data or expert opinion.

\textit{Statistical Analysis: Ordination}

Calculations in dbRDA involve transforming genetic distances (Rousset’s \textit{a}) using a principal coordinates analysis (PCoA; Legendre & Anderson 1999) and then applying a normal RDA on the derived PCoA axes. All PCoA axes with positive eigenvalues are retained for analysis. When distances are completely linear, no axes will have negative eigenvalues. Genetic distances, however, are often not linear (Graves et al. 2013), so dbRDA effectively removes errors created by non-linear distances to make them appropriate for RDA. As an alternative to PCoA, we also utilized spatial principal components analysis (sPCA; Jombart et al. 2008) to derive linear, spatially lagged scores as explanatory variables for a normal RDA.
In population-based analyses, population averages of genetic diversity are often employed as response variables in ordination techniques, but this method is not feasible for individual-based genetics. sPCA incorporates both genetic data and spatial autocorrelation to summarize overall genetic and spatial patterns into independent ordination axes. We chose to use sPCA because this technique removes much of the extraneous variation in genetic data to isolate important patterns and has been successful in isolating landscape effects in empirical individual-based studies (e.g., Robinson et al. 2012; Chapter 2). Unlike dbRDA that retains all positive PCoA axes, authors can choose the number of retained ordination axes from sPCA (Jombart et al. 2008). Typically, axes with the highest eigenvalues are retained for further analysis, but the exact number often depends on a study’s focus and underlying processes affecting gene flow. For simplicity, we retained the first two sPCA axes in all calculations, and derived sPCA lagged scores for genetic explanatory variables. Lagged scores correspond to each individual’s position in ordination space and incorporate both genetic diversity and spatial autocorrelation. All sPCA calculations were performed in the R package ADEGENET (Jombart et al. 2008).

Full and partial dbRDAs (function capscale) and RDAs (function rda) were performed in the R package VEGAN (Oksanen et al. 2013). Statistical significance for all ordination techniques (dbRDA and RDA) was assessed using the anova.cca function in VEGAN; this function is specifically designed to calculate pseudo F-ratios, variance components, and p-values from ordination methods (Oksanen et al. 2013). Since the total variance explained by ecological and geographic variables is typically very small (1.20 – 12.35% in our simulated datasets), we extracted pseudo F-ratios because like Pearson
correlation coefficients, higher $F$-ratios indicate a stronger relationship between landscape or geographic variables and genetic variation.

**Statistical Performance**

In this study, statistical performance was first defined by counting the number of times each test correctly identified the underlying pattern of IBD or IBR (hereby termed accuracy; Table 1.1). Because all landscapes were characterized by an underlying IBD pattern, a correct test occurred when a full test was significant in the R1-R50 landscapes. In the IBR case (R5-R50 landscapes), accuracy of a partial test (in which geographic distances were controlled) was defined by how many significant partial tests occurred out of 100 simulated populations. Along with accuracy, we also counted how often each test did not correctly identify the appropriate mechanism driving gene flow (type I or II errors; Table 1.1). Type I errors occur only in the R1 (IBD only) scenario when partial tests are significant for IBR despite no underlying IBR. In contrast, type II errors were defined as the inability to detect the effect of matrix habitat in the IBR landscapes (R5-R50).

We also assessed the effect of landscape heterogeneity (percent suitable habitat, landscape configuration metrics, and resistance strength) on the strength of detected IBR relationship (i.e., explanatory power of the model). General linear models (glm) for each partial test incorporated either $F$-ratios (ordination techniques) or Pearson correlation coefficients (Mantel tests) calculated from the IBR scenarios (R5-R50) as explanatory variables. Predictor variables included six variables: percent suitable habitat, clumpy, and resistance as well as the residuals of regressions between percent suitable habitat and
correlation length or patch cohesion. Separating configuration (i.e., correlation length and patch cohesion) from habitat amount (percent suitable habitat) effects can be difficult because these landscape variables are highly correlated (all Mantel $r > 0.866; p < 0.001$), but Cushman et al. (2012) showed that configuration metrics explained additional variance in Pearson correlation coefficients. Therefore, percent suitable habitat was regressed against patch cohesion and correlation length to derive residuals that represent the variance in patch cohesion and correlation length not explained by percent suitable habitat. As our total dataset contained 8,000 populations (excluding IBD cases), we randomly selected 20 populations out of the total 100 within each landscape/resistance combination to minimize errors created by large sample size (final $n = 1,600$).

Model selection used an information criterion approach (Burnham & Anderson 2002) to identify the best model among 31 candidate models. Model ranking occurred according to Burnham & Anderson (2002) where models with the lowest Akaike Information Criterion (AIC) and a $\Delta$AIC value $< 2.0$ were the best models. We calculated AIC weights to examine how likely the top model is the best model among all candidate models. Final parameter estimates for resistance and landscape variables were calculated through model averaging of all top models, and parameters that do not include zero explained significant variation within Mantel $r$ or $F$-ratios. All glm and AIC procedures were performed in R.

Results

Both partial Mantel tests and dbRDA identified IBD in all populations (100%) whereas RDA could not explicitly test for IBD because sPCA axes explicitly incorporate
both spatial autocorrelation and genetic variation, making it unable to isolate IBD in a simple test. Unlike IBD detection, where both partial Mantel tests and dbRDA consistently yielded high accuracy rates, IBR detection was highly variable across landscapes and resistance strengths. In the 20% suitable habitat landscapes, partial Mantel tests performed better (53-100% accuracy) than ordination methods (4-94% accuracy), but then were less accurate than ordination as the amount of suitable habitat increased. Type I error rates (i.e., detecting IBR when the resistance of matrix habitat was 1), were lower (5-26%) than type II error rates (17% - 96%) in Mantel tests. Ordination approaches also had higher type I errors, particularly in dbRDA (25-99%; details below; Figure 1.2). Type II errors (i.e., failure to detect IBR when resistance of matrix habitat was greater than 1) occurred in all landscapes and resistances with relatively high frequencies for all tests. High type II error rates were not unexpected given that intrapopulation levels of genetic structure are often quite small. Resistance strength, percent suitable habitat, and landscape configuration influenced all statistical tests’ abilities to detect the effect of matrix habitat.

The effects of landscape configuration and resistance strength on IBR detection accuracy were different for each test. Maximum performance in partial Mantel tests occurred within the 20% suitable habitat at resistance 50 where significant IBR was detected in 82-99% of populations. Correlation coefficients describing relationships between cost distances and genetic differentiation were low overall, ranging from -0.0487-0.125. The two top ranked glm models (summed AIC weight = 0.9991; Table 1.2) included percent suitable habitat, resistance strength, correlation length, and clumpy (all p < 0.005). Pearson correlation coefficients were negatively correlated with percent
suitable habitat, correlation length, and clumpy and positively correlated with resistance strength (Table 1.3). Taken together, these results show that partial Mantel tests detected IBR more often in landscapes with low percent suitable habitat, very high resistance of matrix habitat, and high fragmentation of suitable habitat (low clumpy and correlation length).

Overall, dbRDA had very high type I error rates where IBR was falsely detected in 62-99% of R1 populations (Figure 1.2). In IBR landscapes (R5-R50), dbRDA rarely detected IBR as frequently as the R1 landscapes (Figure 1.3). Accuracy within IBR landscapes was highly variable among resistances and amount of suitable habitat, but highest performance generally occurred within the 20% and 40% suitable habitat landscapes. The three top glm models (summed AIC weight = 0.9304) indicated that F-ratios in R5-50 populations were positively associated with clumpy and negatively related to percent suitable habitat and correlation length (Table 1.3). All three variables remained significant after model averaging. Resistance and patch cohesion were not found to be significant factors in explaining F-ratios (Table 1.2).

Like dbRDA, RDA with sPCA spatially lagged scores had higher type I errors than partial Mantel tests (Figure 1.2). Type I errors in RDA ranged from 25-60%, which was similar to accuracy rates in IBR scenarios (20-68%; Figure 1.3). However, IBR landscapes always had higher numbers of significant tests than the IBD case. The three top glm models included percent suitable habitat, clumpy, and correlation length (summed AIC weight = 0.6259; Table 1.2). Only clumpy and the intercept remained significant after model averaging (Table 1.3), which is likely due to over half of competing models having ΔAICs from 3.0 to 9.0. Clumpy was positively associated with
F-ratios whereas percent suitable habitat and correlation length were negatively correlated with F-ratios in RDA. Taken together, the results for dbRDA and RDA are concordant, and show that IBR detection is highest in fragmented landscapes with a few clumped patches (low correlation length and high clumpy) and low-intermediate amounts of suitable habitat.

Discussion

Ordination techniques possess numerous characteristics that were predicted to enhance differentiation between IBD and IBR models of gene flow in individual-based studies over more common distance-based methods like Mantel tests. Ordination methods are able to overcome the linearity assumptions that characterize distance-based methods, offer flexibility in the type and number of explanatory variables, and provide a suite of statistical outputs that permit thorough interpretation of results. Despite predictions that ordination methods would outperform Mantel tests, our data show that no one test was uniformly superior to any other. Across all landscapes, both partial Mantel tests and dbRDA were effective at detecting IBD regardless of the strength of IBR. The need to transform genetic data for RDA prevented complete isolation of IBD and IBR, making it impossible to explicitly test for IBD. In landscapes that were characterized by both IBD and IBR, IBR detection was highly variable across tests. Mantel tests use pairwise distances that have considerable noise, likely resulting in decreased power to detect IBR. In contrast, ordination techniques are more powerful tests (i.e., can detect IBR more often), but suffer from false detections of IBR likely created by variable
transformations. Landscape composition also had a powerful impact on all test statistics. Maximum performance occurred in landscapes where IBR was strongest (highly fragmented with low suitable habitat). Ordination methods were more robust to changes in the underlying landscape than Mantel tests, suggesting that ordination methods might provide more meaningful conclusions about gene flow when little is known about the resistance of the landscape.

Performance of Mantel Tests and Ordination Techniques

We found that both Mantel tests and dbRDA were highly effective at detecting IBD because they always detected IBD when it was present regardless of the strength of IBR. These high accuracy rates indicate that investigators can be confident that significant Mantel tests or dbRDA are strong evidence for IBD in a natural population. RDA, in contrast, could not test for IBD because sPCA does not explicitly control for IBD (Jombart et al. 2008), resulting in spatially lagged scores that reflect both IBR and IBD. IBD and IBR typically occur simultaneously in nature, so verification of IBD is a critical step before isolating the effects of IBR. Unlike the high accuracy and power in IBD tests, IBR detection was highly variable across tests and landscape configurations.

Mantel tests, the most common test in individual-based landscape genetics, generally had low accuracy rates for IBR tests, failing to detect IBR when it was present. These high type II error rates suggest that Mantel tests’ dependence on pair-wise genetic, cost, and Euclidean distances may reduce power to detect IBR. Indeed, Legendre and Fortín (2010) found that Mantel tests often suffer from reduced power as compared to ordination techniques due to data transformations into pair-wise distances (Legendre &
Fortin 2010). If Mantel tests inherently have reduced power to detect IBR, then the subtle genetic differentiation present in individual-based studies may be too low to consistently detect IBR. The inability of Mantel tests to detect intrapopulation levels of genetic differentiation is concerning because Mantel tests may not detect IBR in empirical studies with levels of genetic differentiation similar to this study. For example, failure to identify genetic differentiation caused by anthropogenic land use could be a critical problem when trying to inform management agencies about how fragmentation impacts populations of conservation concern. Mantel tests can detect barriers if there is adequate lag time (Landguth et al. 2010), but recent anthropogenic barriers may not result in sufficient genetic differentiation to detect IBR. In cases where Mantel tests do not detect a barrier, we recommend use of more powerful tests and field data to better understand anthropogenic barriers to gene flow. Alternatively, if Mantel tests do detect a barrier, researchers can be confident that IBR is present because Mantel tests rarely detected IBR in IBD only landscapes within our simulations.

Despite their low accuracy rates in IBR detection, Mantel tests rarely detect IBR when only IBD is present (i.e., type I errors). This finding illustrates the utility of Mantel tests for individual-based landscape genetics. The low type I error rates found in this study are in contrast to findings for population-based simulation studies, which have found high rates of false IBR detection using Mantel tests (e.g., Balkenhol et al. 2009; Guillot & Rousset 2013). High type I errors for Mantel tests in population-based studies are likely due to greater genetic differentiation between population pairs and substantially fewer pair-wise comparisons than individual-based studies, both of which increase power to detect IBR (and the potential for type I errors). Individual-based simulation studies
have also recorded greater type I errors than this study, especially in complex landscapes with 3 or more habitat types (Graves et al. 2013). In such complex landscapes, a type I error can stem from false significance of an IBR test in an IBD landscape or from detection of an incorrect landscape factor driving gene flow. More ways to generate type I errors in complex landscapes leads to higher overall type I error rates (Graves et al. 2013).

Our results suggest that ordination techniques (dbRDA and RDA) detect IBR more often than partial Mantel tests based on the higher number of significant partial tests in most IBR landscapes. The lower type II error rates in ordination techniques suggest they have higher power to detect IBR patterns than partial Mantel tests, which can be highly useful in situations where genetic differentiation is low, as is common in individual-based studies. For example, anthropogenic changes to once continuous landscapes (e.g., roads or land conversion) require multiple generations of reduced gene flow to produce genetic differentiation within a population (Landguth et al. 2010). Ordination techniques could potentially detect such contemporary barriers faster than Mantel tests because they can detect weaker IBR. However, one critical caveat of ordination techniques is their propensity to detect IBR even in populations that only exhibit IBD (i.e., type I errors). Both dbRDA and RDA exhibited high type I errors, which indicates they have difficulty removing the effects of IBD from IBR.

Type I errors were highest in dbRDA, a likely result of the transformation of genetic distances. Pair-wise genetic distances are transformed using a PCoA, and then all resultant PCoA axes with positive eigenvectors are retained for analysis. In essence, PCoA removes the often high variability in raw genetic distances between individuals, so
the reduced noise within genetic data may inflate type I error rates. PCoA is based on linear distances, so as genetic distances become less linear due to IBR, negative PCoA axes will occur. Thus, the amount of variation explained by PCoA axes is maximized within the R1 landscape because genetic distances are only correlated with Euclidean distance, making the amount of variation contained within PCoA axes very high. When IBR occurs, genetic distances are no longer solely correlated with Euclidean distance, so the PCoA attempts to force the non-Euclidean genetic distances into linear axes. Any remaining variation goes into negative PCoA axes and is not used in the dbRDA, so the amount of variation explained in genetic distances is less than in the R1 landscape.

Though PCoA transformations are designed to linearize and reduce noise in genetic data, dbRDA may not produce valid results in individual-based studies because it cannot effectively distinguish between IBD and IBR due to high type I errors. Like dbRDA, RDA with sPCA lagged scores exhibited high type I errors, which also is likely related to transformation of genetic data into ordination axes. Further investigation is warranted to understand if including more or fewer ordination axes impacts performance in sPCA (and other similar techniques that would collapse genetic variation into ordination axes). In this study, we retained the first two sPCA axes because they explained the most variance and their eigenvalues were usually well differentiated from other axes. Even though sPCA summarizes patterns in spatial autocorrelation and genetic diversity, it does not explicitly control for it, so IBD can contribute to multiple sPCA axes (Jombart et al. 2008). As a result, ordination axes may just reflect IBD instead of patterns in IBR, so use of sPCA and RDA requires individual inspection of each axis to identify patterns that may be associated with IBD and IBR.
Screeplots from sPCA aid in this process because they provide a visual representation of the amount of spatial autocorrelation within each axis (Jombart et al. 2008). Essentially, there is no rule for the number of sPCA axes to retain for genetic analysis, and each axis should be inspected individually to help understand how IBD and IBR impact genetic variation.

Although ordination techniques are more powerful tests (i.e., can detect IBR more often), removing IBD from IBR is difficult when using ordination-based genetic variables. Population-based studies often do not suffer from this problem because they can utilize population-specific variables such as allele frequencies and connectivity indices (see Balkenhol et al. 2009). Population-specific allele frequencies and connectivity indices cannot be calculated for individuals leaving few alternatives to partial Mantel tests in individual-based studies. To date, focus in landscape genetic methods has generally been placed on comparing or developing methods that are alternatives to partial Mantel tests in population-based studies (e.g., Robinson et al. 2012; Wang 2013). Many of these population-based techniques offer considerable promise in individual-based landscape genetics, but development of proper genetic variables for individual-based techniques remains problematic. Therefore, we recommend exploration into alternative individual-based variables that could better disentangle IBR from IBD to maximize the utility of ordination and regression statistics in landscape genetics.

*Influence of Landscape Composition on Partial Statistics*

While intrinsic factors (i.e., variable type) of each test certainly contributed to accuracy and power, the underlying landscape also had a powerful impact on each tests’
ability to detect IBR. Landscapes with high fragmentation and low suitable habitat had the highest performance indices (highest accuracy and test statistics and lowest error rates) across all tests. The high performance in the most extreme cases of fragmentation and habitat amount supported our prediction that highly fragmented landscapes with low suitable habitat would produce the strongest amount of genetic differentiation, making IBR easier to detect. Despite that maximum performance occurred in the same fragmented landscapes with low suitable habitat, partial Mantel and ordination test statistics were sensitive to different landscape metrics (i.e., percent suitable habitat, correlation length, patch cohesion, clumpy, and resistance strength).

Mantel correlation coefficients were significantly associated with all landscape metrics except patch cohesion, indicating Mantel tests are highly sensitive to changes in the underlying landscape. Mantel tests detected IBR more often than ordination techniques only in the landscapes with the most extreme levels of fragmentation (i.e., low correlation length), suitable habitat, and matrix resistance. Correlation length was particularly important for explaining Mantel correlation coefficients because correlation length is an indicator of the extensiveness of suitable habitat. Mantel tests rely on pairwise distance calculations, so measures of landscape resistance and genetic differentiation are highly dependent on how much suitable habitat occurs between two individuals. As correlation length decreases, the amount of suitable habitat between two individuals also decreases, which in turn elevates genetic distances. Therefore, a strong Mantel correlation coefficient will occur in landscapes with small correlation lengths (see Cushman et al. 2012). Another important factor in explaining Mantel correlation coefficients was resistance strength of matrix habitat where a positive relationship was
recorded. Higher resistance strength increases genetic differentiation, so this relationship was not surprising, but it does raise an important issue for use of Mantel tests in empirical studies. Estimating resistances can be difficult without proper life history (Spear et al. 2010; Zeller et al. 2012), so the degree of error in Mantel tests will be unknown. Based on the fairly limited situations where Mantel tests have high accuracy rates, partial Mantel tests should be paired with other analyses like ordination techniques to help distinguish between a type II error and IBD.

Maximum performance (i.e., highest accuracy rates and F-ratios) of ordination methods occurred within the same fragmented, low suitable habitat landscapes as Mantel tests, but clumpy was the most important landscape variable. Clumpy had the biggest impact on strength of IBR in both dbRDA and RDA, which likely reflects the buffer calculation utilized as landscape variables. In more clumped landscapes, percent suitable habitat within a buffer was similar for many individuals. One critical advantage in ordination techniques was that resistance strength of the matrix had no impact on test statistics in either dbRDA or RDA. Ordination techniques do not require accurate parameterization of landscape resistance and are robust to changes in landscape resistance, so ordination techniques could be particularly useful for species lacking relevant life history data (e.g., presence-absence, mark-recapture, path analysis; Zeller et al. 2012) to parameterize resistance landscapes. Ordination techniques offer considerable promise for empirical studies, and we encourage further study with more complex landscapes and natural populations to understand performance of ordination techniques in individual-based landscape genetics.

Conclusions
Investigation of IBR using ordination techniques will probably be most informative when paired with simulated populations that exhibit IBD to quantify type I errors. In particular, simulations of populations under IBD (i.e., null hypothesis) with geographic locations, landscape configuration, and genetic diversity that mimic the study population can aid in characterizing how often statistics produce erroneous conclusions about gene flow (Chapter 2). Simulations should closely mirror the empirical dataset because deviating from observed landscape configuration, genetic data parameters, or sampling schemes can have drastic impacts on statistical results (e.g., Cushman et al. 2012; Cushman et al. 2013). Interpretation of simulated populations in an individual-based framework could follow several paths, but we recommend quantifying type I errors for each proposed landscape hypothesis and only accepting factors that were not detected in IBD landscapes at high frequencies (e.g., 5%). Simulating populations under IBR and comparing them to empirical datasets is another possibility, but meaningful simulations require considerable life history data (i.e., population-specific demographics) that is largely unavailable for many species.

Despite the risks of type I errors with ordination tests, they are potentially more versatile than Mantel tests based on their ability to reduce noise within datasets and utility in a wide variety of landscape genetic scenarios. Partial Mantel tests are likely to perform well in species with strong habitat specialization or patchy distribution (e.g., Schwartz et al. 2009; Wasserman et al. 2010; Shafer et al. 2012). Our results indicate that partial Mantel tests may not be accurate for species that utilize a variety of habitats or are continuously distributed. For these species, estimating resistance can be difficult even with the inclusion of individual movements that relate to single landscape factors.
(e.g., Zeller et al. 2012). In cases where the resistances of landscape factors are unknown and population boundaries are difficult to define, solely applying Mantel tests is unlikely to be an effective method to understand genetic connectivity. Combining Mantel tests and ordination techniques to evaluate specific appropriate hypotheses, synthesizing the results across approaches to facilitate comprehensive interpretation, and using simulations to evaluate significance of test statistics may be the best way to glean meaningful conclusions about gene flow in individual-based landscape genetics.
Figure 1.1. Multifractal landscapes generated for this study where 400 individuals were randomly placed on each landscape. Each landscape consisted of two landscape categories: suitable (light grey) and matrix (black) habitat. The amount of suitable habitat (P) varied from 20% to 80% with five replicates within each P category. Pixels of suitable habitat were given a resistance value of 1 whereas matrix habitat was given one of four values (5, 10, 25, and 50) across simulations for a total of 80 simulated landscapes.
Figure 1.2. Frequency of type I errors in partial Mantel tests, dbRDA, and RDA in simulated landscapes. IBR was incorrectly detected most often in dbRDA tests, and least often in partial Mantel tests. Error bars correspond to standard errors calculated across the 1,600 simulated populations.
**Figure 1.3.** Frequency of type II errors in partial Mantel tests (a), dbRDA (b), and RDA (c) in simulated landscapes. Five replicate landscapes were simulated for each percent suitable habitat (20%, 40%, 60%, 80%), at four matrix habitat resistance levels (R5 – light gray bars; R10 – gray bars; R25 – dark gray bars; R50 – striped bars). One hundred populations were simulated for each of the 80 total landscapes. Type II errors were calculated as the number of populations (out of 100) in which IBR was not detected in our simulated landscapes, and corresponding error bars indicate standard errors calculated across five replicate landscapes. In general, type II errors increased with amount of suitable habitat but were similar across resistance levels for all tests.
Table 1.1. Summary of variables calculated for partial Mantel tests, dbRDA, and RDA analyses. Partial Mantel tests use pair-wise distances for all variables. dbRDA uses genetic distance as the response variable, and individualistic measures of ecological and geographic data as predictors. Genetic data for RDA was derived from a spatial principal components analysis (sPCA). All three tests had three indices of performance (accuracy, type I, and type II error rates) calculated based on 100 simulated populations per landscape.

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<td>Type II error rate</td>
<td>Proportion of populations where tests were not significant for IBR when IBR was present</td>
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Table 1.2. Top six models that explained variation in partial Mantel $r_s$ (a), dbRDA pseudo F-ratios (b), and RDA pseudo F-ratios. Models are ranked according the lowest AIC value, and all those with a $\Delta$AIC < 2.0 are considered top models (bold).

(a)

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(c)

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<th>$\Delta$AIC</th>
<th>Weight</th>
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<td>4.0</td>
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%SH: percent suitable habitat; ResCL: residuals of linear regression between correlation length and percent suitable habitat; ResCoh: residuals of linear regression between patch cohesion and percent suitable habitat
### Table 1.3

Model averaged parameters for the top glm models in partial Mantel tests, dbRDA, and RDA. Only parameters that were shared between models are reported (i.e., models with an ΔAIC < 2.0). When 95% confidence intervals did not include zero, parameters were considered significant (bold).

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%SH: percent suitable habitat; ResCL: residuals of linear regression between correlation length and percent suitable habitat; SE: unconditional standard errors; Upper and Lower CIs: 95% confidence interval boundaries.
Literature Cited


McGarigal K, Cushman SA, Neel MC et al. (2002) FRAGSTATS: Spatial Pattern Analysis Program for Categorical Maps. Computer software program produced by the authors at the University of Massachusetts, Amherst. Available at the following web site: http://www.umass.edu/landeco/fragstats.html.


CHAPTER 2: FINE-SCALE LANDSCAPE GENETICS OF THE AMERICAN BADGER (*TAXIDEA TAXUS*): DISENTANGLING LANDSCAPE EFFECTS AND SAMPLING ARTIFACTS IN AN ELUSIVE SPECIES

**Introduction**

Landscape genetics has become an increasingly important tool for conservation and management by identifying landscape factors that influence genetic connectivity across heterogeneous landscapes. Maintaining genetic connectivity across landscapes is critical for overall population viability (Hanski 1999; Johansson *et al.* 2007), so landscape genetics can provide information to help understand and potentially mitigate the effects of land use change in natural populations (e.g., fragmentation, habitat loss, anthropogenic disturbance; Segelbacher *et al.* 2010). To understand how land use change may impact genetic connectivity within a focal species, researchers must first identify landscape features that are potentially correlated with gene flow. These landscape genetic hypotheses are typically based on variables that predict habitat preferences or occupancy in field-based investigations (e.g., telemetry or presence-absence studies). This method of hypothesis development has been successful in landscape genetics, and multiple studies have shown that important landscape features in field studies such as land cover (Goldberg & Waits 2010; Garroway *et al.* 2011), climatic conditions (Row *et al.* 2014), and anthropogenic barriers (Blanchong *et al.* 2008; Latch *et al.* 2011) also translate to strong correlations with genetic differentiation. When field data corroborates correlations from landscape genetics (e.g., avoided habitats also prevent gene flow; Shafer *et al.* 2012), meaningful conclusions about genetic connectivity can be drawn and used to develop sound conservation and management plans.
Although landscape genetics has certainly proven to be a robust technique for species with extensive field data (e.g., Cushman et al. 2006; Schwartz et al. 2009; Shafer et al. 2012), application to elusive species can be difficult. Two main challenges arise when working with elusive species in landscape genetics: 1) parameterization of landscape genetic hypotheses without relevant life history data, and 2) deciphering between spurious statistical results and true correlations between patterns of gene flow and landscape variables. Ideally, investigators should develop specific hypotheses about how landscape heterogeneity impacts gene flow in a focal species and parameterize models based on *a priori* hypotheses (e.g., Balkenhol et al. 2009; Anderson et al. 2010). For many elusive species, however, we lack information about processes that might affect gene flow such as habitat preferences, distribution, and population structure within a specific study area. Most commonly, expert opinion has been used as a proxy for relevant field data (Zeller et al. 2012). Using expert opinion to parameterize landscape genetic models has been criticized because it is inherently subjective and difficult to assess accuracy without corresponding field data (Spear et al. 2010; Zeller et al. 2012). Without separate field-based data to corroborate landscape genetic results, models based on expert opinion can make distinguishing between an erroneous result and an actual landscape effect difficult for investigators. Spurious correlations in popular statistics like Mantel tests are relatively common even with accurate parameterization (Cushman & Landguth 2010; Graves et al. 2013), so utilizing expert opinion in elusive species requires additional steps to help alleviate potential errors created by improper parameterization. One potential solution to relying on expert opinion for elusive species
is to employ alternative statistical methods that do not require extensive parameterization of landscape genetic models.

Multivariate and regression methods (e.g., Legendre & Legendre 1998; Fortín & Legendre 2010; Wang 2013) could help alleviate challenges associated with parameterizing hypotheses about gene flow in elusive species. Multivariate and regression techniques offer considerable flexibility in the type of landscape and genetic variables that can be evaluated, negating reliance on a priori parameterization of landscape genetic hypotheses. Furthermore, multivariate and regression techniques are considered more robust than correlation statistics in genetics, and therefore, may limit erroneous conclusions due to type I or II errors. Recent studies have demonstrated the utility of multivariate and regression techniques for detecting complex, interacting influences on gene flow within continuously distributed, well-studied species, but have applied these techniques in a population-based framework (i.e., between genetic clusters; Reding et al. 2012 or between sampled areas; Blanchong et al. 2008; Robinson et al. 2012). Extending such methods to elusive species within a single study area or population requires individual-based approaches that can disentangle biologically relevant patterns in gene flow from spurious statistical correlations when there exists little biological data upon which to evaluate conclusions.

All individual-based landscape genetic statistics, even multivariate and regression techniques, are likely to suffer from some degree of error. Statistics aim to partition genetic variation according to specific landscape factors, but errors occur in individual-based studies because there is little genetic differentiation within a single population (i.e., little variance to partition). Errors are especially problematic in studies of elusive species
because statistical errors are difficult to quantify without corroborating field data and spatially biased sampling is almost unavoidable. Spatially biased sampling often causes misleading conclusions about gene flow (e.g., Schwartz & McKelvey 2010; Oyler-McCance et al. 2013), so studies in elusive species would benefit from explicitly testing the role of errors on landscape genetic statistics. Gene flow simulations offer a method to quantify errors in landscape genetic studies by providing a means of replication within a single landscape and control over processes that result in observed genetic variation (e.g., Epperson et al. 2010; Landguth et al. 2010). In particular, simulations can help separate false statistical significance (i.e., type I errors) from actual landscape effects on gene flow because multiple statistics in landscape genetics can have high type I error rates even in idealized sampling schemes (Balkenhol et al. 2009; Graves et al. 2013; Guillot & Rousset 2013). By quantifying type I errors using gene flow simulations, investigators can better understand how landscape heterogeneity impacts gene flow in elusive species, and separate those effects from sampling artifacts.

The American badger (*Taxidea taxus*) is one of the most elusive and poorly understood mesocarnivore species in North America. Badgers inhabit much of central and western North America, but much of their life history is unknown due to their nocturnal, semifossorial life style. Scarce telemetry studies illustrate dramatic differences in home range sizes and habitat selection among regions. For example, in British Columbia, badgers select habitats away from closed canopy forest (Apps et al. 2002), but those in Ohio used fallow fields and other uncultivated habitats within highly agricultural areas (Duquette & Gehrt 2014). Considerable individual variation in habitat selection also occurs within study areas, making it difficult to predict landscape effects on gene
flow. Based on the limited and likely variable life history data throughout most of North America and their elusive nature, this study on badgers suffered from both challenges associated with elusive species: limited life history data to parameterize landscape genetic models and unavoidable spatially-biased sampling.

In this study, we examined the utility of integrating multivariate and regression techniques with simulations for overcoming the challenges associated with limited life history and spatially biased sampling often observed in elusive species. To identify landscape factors that influence badger gene flow, this study used multivariate and regression techniques to test if isolation-by-distance (IBD; Wright 1943), badger’s preference for treeless habitats, and a major topographic barrier within the study area are correlated with genetic variation of badgers in Wisconsin. Multivariate and regression techniques can test for these factors simultaneously, making them highly useful for disentangling multiple influences on gene flow without the need for parameterizing landscape hypotheses. Obtaining a large number of samples from the elusive badger required a statewide citizen-based effort, resulting in spatially biased sampling with samples clumped around populated areas, a factor that could lead to erroneous conclusions about gene flow. Therefore, we also performed gene flow simulations to calculate how often our chosen statistics falsely identified landscape effects (i.e., type I errors) to separate significant results created by sampling and actual landscape effects.

Methods

Study Area
Our study area (110,745 km\(^2\)) encompasses the state of Wisconsin in the Upper Midwest, United States. The landscape exhibits a transition from a mixture of native grasslands and agriculture in the south to more forested habitats as latitude increases. Badger activity has been recorded in every county in Wisconsin based on citizen-based monitoring of badgers from 2009-2014 and Wisconsin Department of Natural Resources mammal surveys from 1987-2008 (Wydeven et al. 1998; Kitchell 2008); both suggest badgers have a relatively continuous distribution throughout Wisconsin. Despite their continuous distribution within Wisconsin, recent genetic evidence suggests that badgers in Wisconsin represent a unique genetic population within North America due to the Mississippi River and Great Lakes (Chapter 3).

Sample Collection

Since badgers are protected from harvest in Wisconsin as a Species with Information Needs (Wisconsin Department of Natural Resources, 2008), 250 individuals were sampled via road-kills (n = 136 tissue samples) and non-invasive hair collections (n = 114 hair samples) from 2005-2013. Hair collection involved attaching a snare (modified from British Columbia Ministry of Environment Ecosystems Branch for the Resources Information Standards Committee 2007) to the entrance of an active burrow and waiting overnight for the animal to pass under the snare. Snares were designed to line the entrance of the burrow, so hair was typically pulled from both the back and sides of an animal. A successful hair collection typically contained 20 to 50 hairs with intact roots, ensuring a sufficient number of hairs for molecular analysis. No burrows were
sampled repeatedly, and we limited hairs to long banded hairs that are typical of adults to help prevent sampling of younger animals (juveniles and kits).

Location information for each individual was recorded in latitude and longitude coordinates. For a few individuals with less precise location information (i.e., Counties, Public Land Survey System locations; n = 42), we utilized the “Create Random Points” function in ArcMap v. 10.1 to assign a spatial coordinate. In Wisconsin, the maximum amount of ambiguity around a point was approximately 10 km (locations as counties). No change in results was detected by excluding individuals with spatial ambiguity from analysis.

**Laboratory Methods**

DNA was extracted using Qiagen DNEasy blood and tissue kits for both tissues and hairs. Extractions used a 1 mm$^3$ piece of tissue or 10 to 20 hairs with intact follicles in a user-refined extraction protocol (Qiagen 2006). Hairs were processed on different days than tissue samples in dedicated laboratory space. We amplified all samples at 12 microsatellite loci (Table S2.1) developed in American badger (Tt-1, Tt-2, Tt-3, and Tt-4; Davis & Strobeck 1998), American mink (*Neovison vison*; Mvis072; Fleming *et al.* 2002), American marten (*Martes americana*; Ma-1; Davis & Strobeck 1998), and European badger (*Meles meles*; Mel112, Mel101, Mel111, Mel108, Mel14, and Mel1: Carpenter *et al.* 2003; Domingo-Roura *et al.* 2003). Multiplex polymerase chain reactions (PCRs) were conducted in sets of 2-4 primers in 10 uL reaction volumes (Table S2.1). Amplified products were genotyped on an ABI 3730 DNA Analyzer at the
University of Wisconsin Biotechnology Center, and alleles were sized using the program GeneMarker (SoftGenetics LLC, State College, PA, USA).

We utilized a comparative multi-tube approach for genotyping hair samples (Frantz et al. 2003; modified from Navidi et al. 1992 and Taberlet et al. 1996) where each hair extract (two for most samples; n = 85, single extraction; n = 39) was genotyped three (heterozygotes) to seven times (homozygotes) to validate resultant genotypes. Tissue samples were collected from road-killed animals and varied considerably in quality, so we also re-extracted and re-genotyped 50% of tissue samples (n = 68). All resultant homozygotes and 25% of heterozygotes in tissues were re-genotyped as a final check. In total, we identified 6 instances of allelic dropout (all hairs) in the 918 repeated genotypes (0.65% error rate). Any individuals with fewer than 10 genotypes were then culled from the dataset (17 hair samples culled; 233 retained individuals).

**Statistical Analysis**

We used two complementary Bayesian clustering approaches (non-spatial STRUCTURE 2.2.3; Pritchard et al. 2000 and spatial BAPS 5; Corander et al. 2008) to characterize population structure within Wisconsin. In our non-spatial approach, we employed Bayesian clustering in STRUCTURE with five independent runs (100,000 MCMC burn-in, 100,000 permutations) at each hypothesized number of genetic clusters (K) under the admixture, correlated alleles model (Pritchard et al. 2000). The optimal value for K among tested values (K = 1-10) was determined using Evanno et al.’s (2005) ΔK estimator because likelihood values plateaued and variances among runs grew larger at values of K above the optimum (Pritchard et al. 2000; Figure S2.1). Once the optimal
$K$ was identified, five longer runs of 1,000,000 MCMC burn-in, 1,000,000 permutations were conducted to calculate the proportion of each individual’s genome that belongs to each cluster ($q$). Average $q$-values across the 5 runs were calculated in CLUMPP (Jakobsson & Rosenberg 2007), and individuals were assigned to a cluster based on their highest $q$.

In our spatially informed approach, we employed BAPS 5. We tested $K = 1$ through 10 (5 replicates per $K$) using the “Spatial Clustering of Individuals” option in BAPS. Maximum likelihood and highest posterior probability were used to determine the optimal number of genetic clusters in the sample. Admixture between inferred clusters was calculated using 500 simulations based on observed allele frequencies.

For the total sample and the inferred clusters, we calculated population-specific measures of genetic diversity (allelic richness, number of alleles, and heterozygosity, and $F_{IS}$) and genetic differentiation among clusters ($F_{ST}$) in the R package diveRsity (Keenan 2013). Deviations from Hardy-Weinberg (HWE) and Linkage (LE) equilibria were calculated in GENEPOP (Raymond & Rousset 1995) using a corrected alpha for multiple tests ($\alpha = 0.012$; false discovery rate; Benjamini & Yekutieli 2001).

To test for the effect of geographic distance on genetic differentiation, we quantified patterns of IBD within Wisconsin using two complementary approaches. We used simple Mantel tests to test for an association between matrices of pairwise genetic and geographic distances and dbRDA to test for significant effects of geography (latitude and longitude) on the distribution of genetic variation across the study area. Both Mantel tests and dbRDAs were conducted in the R package VEGAN using the functions “mantel” and “capscale” (Oksanen et al. 2008).
In addition to tests for IBD, spatial autocorrelations were used to detect departures from random mating (i.e., panmixia) within 5 km distance categories. Individuals separated by small geographic distances are expected to exhibit positive spatial autocorrelations (i.e., be more genetically similar than expected under panmixia). Statistical significance in spatial autocorrelations was assessed after 1000 permutations using custom code in R. Mantel tests, dbRDA, and spatial autocorrelations were run with the heavily sampled counties subsampled by factors of 5 (0, 5, 10, 15, and 20 individuals were included from Bayfield, Dane, and Iowa Counties) to alleviate any potential sampling bias.

Population structure is often influenced by discrete barriers (isolation-by-barrier; IBB) in addition to geographic distance (IBD). We used a partial Mantel test to determine correlational significance between a genetic distance matrix (Rousset’s \(a\); Rousset 2000) and a barrier matrix while controlling for geographic distance. The barrier matrix was a binary indicator of whether a pair of individuals was on the same (0) or different (1) sides of the Wisconsin River, the most prominent potential barrier to badger gene flow in Wisconsin. Calculations were performed using the R package \texttt{VEGAN}, and statistical significance was assessed via Pearson correlation coefficients after 1000 permutations.

In nature, many different landscape factors can work in tandem to create observed patterns of genetic variation (e.g., Cushman \textit{et al.} 2006). Thus, we sought to incorporate both barriers and ecological variables (collectively called landscape factors) into a cumulative model that explains how all landscape factors influence genetic variation in badgers. Ecological variables included level III ecoregions and land cover (Figure 2.1).
We used level III Ecoregions to define broad scale regions of similar abiotic (i.e., climate and soil) and biotic assemblages (in Wisconsin: Northern Lakes and Forest; North Central Hardwood Forest, Driftless Area, and Southeastern Wisconsin Till; Omernik 1987). Land cover variables were produced via an intersection of two rasters: native soil associations within Wisconsin (Hole 1976) and land cover (NLCD2006; Fry et al. 2011), and then summarized into three land cover categories (Native Open, Forest, and Agriculture) that represent a continuum of habitat suitability for badger. Proportions of each land cover category were calculated within 5 km circular buffers drawn around each badger location. Proportions of Native Open, Forest, and Agriculture were highly correlated ($r = -0.35$ to $-0.65$, all $p < 0.001$), so only one land cover variable was used within each statistical model at a time.

Multivariate techniques like spatial principal components analysis (sPCA; Jombart et al. 2008) are highly effective at detecting both discrete barriers and genetic gradients, which makes them ideal for disentangling complex patterns of gene flow (Jombart et al. 2008). We used the R package adegenet (Jombart 2008) to perform sPCA calculations and significance tests for global and local patterns using 999 permutations. We used an inverse distance-weighting network so that all badgers were considered neighbors. The first two sPCA axes explained the most variation (0.0889 and 0.0571, all others < 0.0459; Figure S2.3), and were retained as dependent variables in landscape genetic analysis (Figure 2.2).

We utilized two approaches to distinguish between landscape factors and geographic distance, both of which can drive patterns in sPCA axes (Jombart et al. 2008). We performed a partial redundancy analysis (RDA), a constrained ordination technique
that is the multivariate analog to simple linear regression (Legendre & Legendre 1998). The two retained sPCA axes were analyzed together as explanatory variables with both latitude and longitude in the conditional matrix (i.e., variables to be controlled for in final models). The barrier (Wisconsin River) and ecological (Ecoregion or Agriculture) variables were included as predictors. We performed a stepwise model selection procedure to identify the variables that best explained genetic differences among individuals. We used the R package \textit{vegan}, with significance assessed with 1000 permutations using function “ordistep” (Oksanen \textit{et al.} 2008).

We used spatially lagged regression models as a second approach to evaluate the evidence for landscape influences on sPCA axes by controlling for the potentially confounding influence of IBD. Spatially lagged regression models evaluate each sPCA axis individually, which provides further evidence for specific influences on patterns found within the sPCA. Spatially lagged regression models control for IBD by including a spatial weighting matrix ($W_{ij}$) and $\rho$, a parameter that accounts for the lack of independence between individuals (Legendre & Legendre 1998). The weighting matrix was constructed based on an inverse weighting calculation as this procedure is thought to best approximate spatial autocorrelation under IBD (Robinson \textit{et al.} 2012). Regression models: $G_i = \rho W_{ij} G_j + \beta X + \varepsilon$ included sPCA scores for a focal individual $i$ ($G_i$) and all other individuals ($G_j$) along with explanatory barrier and landscape variables ($X$), their estimated effects ($\beta$), and residual error ($\varepsilon$). A LaGrange Multiplier test then tests for spatial autocorrelation within residuals of the regressions where a significant test indicates autocorrelation remains within the data. One major advantage of spatial regression is the ability to conduct model selection procedures such as Aikake
Information Criterion (AIC). Model selection followed Burnham & Anderson’s (2002) method where models with a ΔAIC < 2.0 were considered candidate models for explaining a sPCA axis. Parameter estimates were produced through model averaging of all candidate models where significant parameters do not include zero (Burnham & Anderson 2002). All spatial regression methods were performed using the “lagsarlm” function in R package spdep (Bivand et al. 2011).

To assess type I errors within sPCA, partial RDA, and spatially lagged regression models, we simulated 100 populations where only geographic distance influenced gene flow in the program CDPOP v. 1.4 (Landguth & Cushman 2009). Populations were simulated using an inverse square dispersal distribution because strong sex-biased dispersal has not been recorded in badgers (Messick & Hornocker 1981; Hoodicoff 2003; Kierepka et al. 2012). Mating was sexual with replacement and generations were overlapping. The initial population consisted of 1,400 individuals; 1,167 were placed randomly on Wisconsin’s landscape excluding urban and open water habitats and 233 were placed at geographic coordinates identical to our empirical dataset. After 250 generations, we sampled the 233 individuals that had the same geographic locations as the empirical dataset. We performed sPCA, partial RDA, and spatially lagged regression analyses for each simulated population, defining type I errors as false detection of a barrier (Wisconsin River) or ecological (Ecoregion, land cover) factor as a driver of population genetic structure in more than 5 of the 100 simulated populations (alpha = 0.05).

Results
In the BAPS analysis, the optimal solution was $K = 1$ (-11197.19). The STRUCTURE analysis indicated stronger support for $K = 2$ than for $K = 1$ (Figure S2.1a). However, an inspection of the STRUCTURE assignments revealed a gradient pattern (Figure S2.1b). The resultant gradient in q-values and results from BAPS suggests that the inferred structure was likely an artifact of IBD or spatial autocorrelations because both can create false genetic clusters in Bayesian programs (Frantz et al. 2009; Schwartz & McKelvey 2009).

We observed a heterozygote deficiency within Wisconsin over all loci in a global analysis (Table 2.1) and for seven individual loci (all $p < 0.001$). These deviations from HWE are expected if any deviation from panmixia exists within the dataset. All loci were highly polymorphic ranging from 7 to 15 alleles per locus (average = 11.58 alleles/locus), and showed no linkage disequilibrium (all $p > 0.025$).

Results from the dbRDA analysis supported the Bayesian clustering and HWE analyses that indicated a role for geography driving gene flow patterns. However, the dbRDA analysis was significant for only latitude ($F = 2.404$, $p = 0.008$), not longitude ($F = 1.009$, $p = 0.124$). Also, the simple Mantel test did not show a significant correlation between matrices of genetic and geographic distances ($r = 0.012$, $p = 0.0312$). These results suggest that variability in longitude across Wisconsin may be too small to detect IBD.

Despite the equivocal evidence for IBD in dbRDA and Mantel test, we found support for fine-scale spatial autocorrelations. Genetic distances between proximate individuals (25 km or less) were smaller than expected under panmixia in the spatial
autocorrelation (Figure S2.2). In particular, individuals separated by 5 km were highly autocorrelated indicating that individuals found closer together were more genetically similar. One explanation for the strong positive autocorrelations at 5 km is that we sampled relatives in Dane, Iowa, and Bayfield Counties where 59/90 pair-wise comparisons under 5 km occurred. However, most of these samples were road-killed individuals sampled across years or live captured adult animals at separate burrows. We completely removed these counties in another set of spatial autocorrelation tests, and the strong positive autocorrelation remained at 5 km. Therefore, we argue that the positive spatial autocorrelation is not an artifact of sampling relatives.

We found no evidence for IBB resulting from the Wisconsin River in either Bayesian analyses or partial Mantel tests. Visual inspection of population assignments from STRUCTURE revealed a gradient in q-values that lacked a strong genetic break along the Wisconsin River. The partial Mantel test also revealed a lack of barrier effect ($r = -0.0266, p = 0.992$).

Spatial PCA axes revealed two main patterns within Wisconsin: a latitudinal cline across the entire state (Axis 1) and an area of high genetic similarity in central Wisconsin (Axis 2; Figure 2.2). Both axes had signatures of spatial autocorrelation (Moran’s I = 0.572 and 0.301), and Axis 1 explained more variation in genetic diversity than Axis 2 (variance = 0.175 vs. 0.152). Loci Mel1, Tt-2, Mvis072, Mel101, and Mel108 were most informative for Axis 1 and Mel111, Tt-2, and Mel1 were most useful for Axis 2. The Monte-Carlo test confirmed the existence of at least one global pattern (observed: 0.010, $p = 0.001$) but no local pattern (observed: 0.006, $p = 0.362$).
Results from the partial RDA and spatially lagged regressions indicated that the Wisconsin River and Agriculture were correlated with the two sPCA axes. In the partial RDA analysis, both the Wisconsin River (F = 22.44, p < 0.001) and Agriculture (F = 59.416, p < 0.001) were retained after model selection. Ecoregions, in contrast, were not included in the final RDA models. Only Agriculture was found to be associated with sPCA spatially lagged scores for both sPCA axes; Forest or Native Open were not retained as significant variables when used as the land cover variables.

All top spatially lagged regression models for Axis 1 contained the Wisconsin River and model averaging revealed that the parameter estimate did not include zero (parameter estimate = -0.225 ± 0.144; Table 2.2). However, observed models also had relatively high residual spatial autocorrelations (LaGrange Multiplier test: = 1.967-4.768, p = 0.0290-0.116), suggesting that IBD was also associated with Axis 1. All top models for Axis 2 included Agriculture (Agriculture, Agriculture + River, Agriculture + Ecoregion; Table 2.2). Model averaging between these three models indicated that Agriculture was the only significant variable (model averaged parameter: -0.318 ± 0.284). Observed models indicated a lack of remaining spatial autocorrelation in Axis 2 (LaGrange test: 0.379-0.517, p = 0.439-0.538). Like the RDA analysis, when Forest and Native Open were included as land cover variables, they were not significantly associated with sPCA spatially lagged scores of either axis.

Simulations revealed similar type I error rates for partial RDA and spatially lagged regression (Figure 2.3). The Wisconsin River (52-57/100 populations) and Ecoregions (68-76/100 populations) were falsely detected as ecological variables affecting gene flow much more frequently than our 5 tests out of 100 cut-off. A likely
cause of high error rates was that residual autocorrelation (i.e., IBD) remained within the data (LaGrange Multiplier test: all p < 0.062) in almost all cases of significance. This high false significance rate in simulated populations suggests that our finding of an association between the Wisconsin River and spatially lagged scores from Axis 1 in our empirical dataset was likely an error. A lack of barrier effect for the Wisconsin River was also supported by the Bayesian analyses and partial Mantel test. In contrast, Agriculture was falsely detected as an ecological variable affecting gene flow in less than 5% of populations regardless of the test (3-5/100 populations; Figure 2.3). Therefore, our finding of an association between Agriculture and the spatially lagged regression scores was unlikely to be a type I error in our empirical dataset, and represents an ecological variable that likely impacts gene flow.

Discussion

There are many challenges associated with elusive species that can affect conclusions about how features of the landscape influence gene flow. In badgers, both limited life history data and spatially biased sampling were obstacles that could potentially confound our interpretation of landscape genetic statistics. Our study demonstrates that combining multivariate and regression statistics with simulations to quantify errors can help disentangle potential errors from landscape effects on gene flow in an individual-based framework. RDA and spatially lagged regression did not require parameterized hypotheses, which was critical for this study given the nearly complete lack of relevant life history data. With simulations, we were able to quantify the
confounding effects of spatially biased sampling to separate type I errors from biologically relevant landscape effects on gene flow. Following error assessment, our genetic dataset indicated that geographic distance is the strongest influence on badger gene flow within Wisconsin, with agriculture playing a lesser role.

Geographic distance was the primary driver of gene flow in badgers, as evidenced by dbRDA and spatial autocorrelations. Both axes in the sPCA had strong signatures of spatial autocorrelation, particularly in axis 1, further demonstrating that geographic distance influenced gene flow in badgers. The non-significant simple Mantel test and dbRDA for longitude were inconsistent with the sPCA results, which suggests that geographic distance is only important at local scales. Positive spatial autocorrelations were particularly pronounced under 5 km in our dataset, which could reflect either a behavioral mechanism or sampling artifact. Restricted dispersal can occur in mammals due to philopatry, particularly in females (Greenwood 1980), but little evidence has been found for philopatry in American badgers (Messick & Hornocker 1981; Kierepka et al. 2012). However, dispersal regimes can vary according to habitat quality where dispersal is more restricted in suitable habitat (e.g., Broquet et al. 2006; Frantz et al. 2009), so high spatial autocorrelations may indicate badgers may exhibit some degree of restricted dispersal in Wisconsin. Badgers were highly genetically similar in areas with more suitable habitat, but these areas were also the most heavily sampled. Within these heavily sampled counties, proximate pairs of individuals were often collected as road-killed animals, making it difficult to determine whether individuals were killed within suitable habitat or during dispersal though matrix habitat. Regardless of the mechanism, positive spatial autocorrelations at fine scales is a fairly ubiquitous factor influencing spatial
patterns of genetic variation in other highly mobile carnivores (e.g., Schwartz et al. 2006; Cegelski et al. 2006; Zalewski et al. 2009; Croteau et al. 2010), so it is not surprising to find that geographic distance exhibits a strong influence on gene flow in badgers as well.

After controlling for the influence of geographic distance across Wisconsin, our multivariate and regression statistics detected the Wisconsin River as a potential influence on patterns in sPCA axis 1. However, considerable residual autocorrelation remained in the spatially lagged regressions and differentiation in sPCA axis 1 appeared to be greatest between the southeastern and northwestern areas of Wisconsin (i.e., consistent with geographic distance). Simulated populations in which geographic distance was the only influence on gene flow also frequently detected a barrier effect of the Wisconsin River, indicating that the statistically significant effect we observed in our dataset is likely a sampling artifact. Badger activity in the largely forested northcentral areas of Wisconsin was rarely reported, so most sampled individuals coded as west of the Wisconsin River were from northwestern Wisconsin. Also, the Wisconsin River occurs in the center of our study area, so removing the effect of geography in the spatial regression and RDA was difficult as evidenced by the high residual autocorrelation. Therefore, the gap in sampling west of the Wisconsin River combined with the underlying isolating effects of geographic distance appear sufficient to create the statistically significant associations between patterns of genetic variation and the Wisconsin River.

Unlike the Wisconsin River, type I error rates were low for Agriculture, which suggests agricultural landscapes influence gene flow within Wisconsin badgers. In this case, agriculture appeared to facilitate gene flow more readily than other habitats as
evidenced by the high genetic similarity of individuals (Axis 2 of sPCA) within agricultural habitats. Optimal habitats often facilitate gene flow (Cushman et al. 2006; Schwartz et al. 2009), but badgers generally avoid agriculture in other portions of their range (Messick & Hornocker 1981; Warner & Ver Steeg 1995; Duquette & Gehrt 2014). Avoidance behavior suggests that agriculture is not an optimal habitat for badgers, so understanding the exact role of agriculture in driving gene flow across Wisconsin is not straightforward without corroborating field data on dispersal.

Several mechanisms could explain why agricultural habitat is correlated with genetic variation in badgers. First, agriculture could be significant because of its tight correlation with Native Open, the likely preferred badger habitat. Native Open was not significantly associated with either sPCA axis, but sampling within native open habitats was relatively sparse compared to agriculture. Robust sampling occurred within the northwestern native open habitats unlike the native open habitats within the largely agricultural southern Wisconsin. In southern Wisconsin, native open habitats were often interspersed with agriculture, but spatial data may have lacked the resolution to capture fine-scale habitats (e.g., native open habitats along fencerows) necessary for dispersal (Anderson et al. 2010). Badgers in other agricultural landscapes utilized linear corridors along agricultural areas for movement (Duquette & Gehrt 2014), so agriculture as a whole may appear to facilitate gene flow in Wisconsin. Alternatively, density-dependent dispersal could promote gene flow from suitable habitats into agriculture (e.g., Carr et al. 2007). As resident badgers tend to remain in the same area over consecutive years (Lindzey 1978; Messick & Hornocker 1981), a lack of unoccupied areas within suitable habitats (i.e., native open) may force transient and juvenile individuals into agriculture, a
sub-optimal habitat. Both of these putative mechanisms would produce the pattern observed in sPCA Axis 2 where individuals in high intensity agricultural habitats show high genetic similarity to those in surrounding suitable habitats (e.g., Native Open or pasturelands in central and southwestern Wisconsin, respectively). Both mechanisms also would produce elevated $F_{IS}$ values in badgers in agricultural habitat relative to other habitats, a pattern that was present in our study but not significant. Determining the exact role of agriculture in badger gene flow is difficult because we do not know whether animals sampled in agriculture were transient dispersers or had established home ranges. Despite an inability to distinguish between potential mechanisms causing landscape genetic patterns in badgers, genetic data can be used to develop focused hypotheses (i.e., how badgers disperse in agricultural habitats) that with targeted sampling, can test how these specific landscape features impact connectivity.

When interpreted properly through rigorous error testing, regression and multivariate statistics are well suited to elusive species like badgers in individual-based landscape genetics. One benefit of multivariate and regression techniques is that they offer considerable flexibility in both landscape and genetic variables. To date, population-based studies have successfully incorporated a wide array of variables into multivariate and regression techniques (e.g., Balkenhol et al. 2009; Reding et al. 2012; Robinson et al. 2012), but many of those population-specific variables are not applicable to individual-based approaches. Our study with badgers emphasizes that multivariate and regression techniques are equally useful in individual-based studies with appropriate choices in landscape and genetic variables. We selected our categorical land cover variables based on the limited telemetry studies in badgers (e.g., Apps et al. 2002;
Duquette & Gehrt 2014) because no movement or habitat preference data is available in Wisconsin or nearby areas. Deriving pair-wise connectivity metrics for badgers would have had considerable uncertainty given the substantial individual and population variation in demography and habitat associations throughout their range (Messick & Hornocker 1981; Apps et al. 2002; Duquette & Gehrt 2014). Even in well-studied species, landscape genetic patterns vary between landscapes, primarily due to the presence or spatial arrangement of important factors (e.g., Short Bull et al. 2011). RDA and spatially lagged regressions do not necessitate extrapolation to produce pair-wise connectivity metrics, and can still detect subtle impacts on gene flow as seen with agriculture in badgers. Therefore, their flexibility in variable type makes multivariate and regression techniques attractive for a myriad of landscape genetic studies, including those focused on species with limited life history information.

In addition to flexibility, multivariate and regression techniques may be particularly useful in conservation because both techniques have high power to detect fine-scale landscape genetic patterns (Balkenhol et al. 2009; Fortín & Legendre 2010). In this study, both RDA and spatially lagged regressions detected agriculture’s subtle impact on gene flow despite the strong influence of geographic distance on genetic variation. Based on their ability to detect landscape genetic patterns in simulated spatial gradients with considerable noise (Fortín & Legendre 2010), multivariate and regression techniques likely would perform well in situations with complex patterns in gene flow (e.g., Bowen et al. 2005; Kamler et al. 2013) and biased sampling. Although multivariate and regression techniques are flexible, statistically robust, and provide a means of multi-model inference, error assessment is also critical as all statistics utilized in this study
were also vulnerable to type I errors. When combined with simulations to quantify potential errors, multivariate and regression techniques can be a powerful tool to inform conservation and management efforts in poorly understood species.

Overall, this study provides important information for future management of the American badger in Wisconsin, a population that is genetically distinct due to the Mississippi River and Great Lakes (Chapter 3). Badger gene flow within Wisconsin is largely unrestricted despite heterogeneous habitat composition and the presence of large riverine barriers. Future studies would be helpful to assess potential mechanisms that could explain the relationship between agriculture and genetic variation observed in this study (i.e., conduits of dispersal or density dependent dispersal). This study demonstrates the utility of multivariate and regression methods within individual-based landscape genetics, which to date, were largely restricted to population-based investigations of well-studied species. With multivariate and regression methods and explicit error assessment, individual-based landscape genetic approaches can provide valuable insights into how landscape heterogeneity impacts genetic variation despite limited life history and biased sampling.
Figure 2.1. Predictor variables for each georeferenced badger describe its location according to three landscape features (Wisconsin River, level III Ecoregion, and land cover). For the Wisconsin River (blue line), each individual was coded as either east or west of the river (a). All badger locations fell within one of four level III Ecoregions within Wisconsin (b): Driftless Area (DA), Northcentral Hardwood Forests (NCHF), Northern Lakes and Forest (NLF), and Southwestern Wisconsin Till Plain (SWTP). Land cover data for each badger was calculated as percent land cover within a circular buffer surrounding each badger’s location (most predominant land cover class given by point’s color; c).
Figure 2.2. Spatially lagged scores for the first two sPCA axes for Wisconsin badgers. Scores from Axis 1 (a) and 2 (b) were correlated with the Wisconsin River and percent agriculture respectively. Dark colors (black and dark grey) represent negative sPCA scores while positive values are light in color (white and light grey). More extreme values in sPCA axes are displayed with larger squares.
Figure 2.3. Percent of false significant tests (type I errors) for the Wisconsin River, level III Ecoregions, and Agriculture calculated in 100 simulated IBD populations. Both partial RDAs and spatially lagged regressions (sReg) for each axis separately incorrectly identified the Wisconsin River and level III Ecoregions as influences on gene flow more often than Agriculture.
Figure S2.1. Results of Structure analyses from $K = 1$ to $K = 10$. Maximum likelihoods (grey squares) and $\Delta K$ values (black diamonds) occurred at $K = 2$ (a). Error bars correspond to standard deviations around each likelihood across five runs per $K$. A heat map was constructed from averaged q-values for $K = 2$ based on inverse distance weighting (b). The heat map revealed no conspicuous barriers, but rather exhibited a gradient from northwest to southeast.
Figure S2.2. Spatial autocorrelation for all badgers within Wisconsin. Dotted lines correspond to 95% confidence intervals derived from 1000 permutations of Rousset’s $a$ within each 5 km distance class. Each point represents average Rousset’s $a$ at a distance class, and those below the 95% confidence intervals (< 20 km) had lower genetic distance than expected by random mating.
**Figure S2.3.** Eigenvalues (a) and screeplot of sPCA (b) displaying the variance explained and spatial autocorrelation within each sPCA axis. Axis 1 and 2 were retained for further analysis because they had the highest eigenvalues and explained the most variance within the dataset.
Table 2.1. Locus-specific summary of genetic variation for n = 233 badgers in Wisconsin. Metrics included are the number of alleles per locus (A), observed heterozygosity ($H_O$), expected heterozygosity ($H_E$), inbreeding coefficients ($F_{IS}$), and 95% confidence intervals for $F_{IS}$ for each locus. Asterisks indicate significant deviations from HWE at a corrected $\alpha = 0.012$.

<table>
<thead>
<tr>
<th>Locus</th>
<th>A</th>
<th>$H_O$</th>
<th>$H_E$</th>
<th>$F_{IS}$</th>
<th>$F_{IS}$ Lower CI</th>
<th>$F_{IS}$ Upper CI</th>
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<td>11</td>
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<td>0.85</td>
<td>0.198*</td>
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<td>10</td>
<td>0.81</td>
<td>0.83</td>
<td>0.022</td>
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<td>0.083</td>
</tr>
<tr>
<td>Mel1</td>
<td>13</td>
<td>0.81</td>
<td>0.87</td>
<td>0.071*</td>
<td>0.012</td>
<td>0.126</td>
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<tr>
<td>Tt-1</td>
<td>13</td>
<td>0.79</td>
<td>0.81</td>
<td>0.024</td>
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<td>0.084</td>
</tr>
<tr>
<td>Tt-2</td>
<td>7</td>
<td>0.59</td>
<td>0.68</td>
<td>0.133*</td>
<td>0.044</td>
<td>0.219</td>
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<td>Ma-1</td>
<td>9</td>
<td>0.68</td>
<td>0.68</td>
<td>0.000</td>
<td>-0.079</td>
<td>0.083</td>
</tr>
<tr>
<td>Tt-3</td>
<td>15</td>
<td>0.77</td>
<td>0.88</td>
<td>0.126*</td>
<td>0.063</td>
<td>0.187</td>
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<td>Tt-4</td>
<td>14</td>
<td>0.83</td>
<td>0.88</td>
<td>0.064*</td>
<td>0.008</td>
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<td>Mvis072</td>
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<td>0.64</td>
<td>0.81</td>
<td>0.209*</td>
<td>0.136</td>
<td>0.283</td>
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<td>-0.008</td>
<td>0.102</td>
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<tr>
<td>Mel108</td>
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<td>0.78</td>
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<tr>
<td>Mel112</td>
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<td>0.81</td>
<td>0.261*</td>
<td>0.184</td>
<td>0.337</td>
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<tr>
<td>Overall</td>
<td>11.83</td>
<td>0.73</td>
<td>0.81</td>
<td>0.103*</td>
<td>0.080</td>
<td>0.122</td>
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Table 2.2. Model selection results for spatially lagged regressions on sPCA Axes 1 and 2. Top models (ΔAIC < 2.0; in bold) for Axis 1 all included the Wisconsin River (River) whereas models for Axis 2 incorporated Agriculture (Ag). Ecoregion (Eco) was not significant for either axis following model averaging.

<table>
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<tr>
<th>Model</th>
<th>AIC</th>
<th>ΔAIC</th>
<th>w_j</th>
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<tr>
<td><strong>Axis 1</strong></td>
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<td></td>
<td></td>
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<tr>
<td>Eco+River</td>
<td>169.66</td>
<td>0.00</td>
<td>0.499</td>
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<tr>
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<tr>
<td>River</td>
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<td>0.200</td>
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<tr>
<td>Ag+River</td>
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<td>3.28</td>
<td>0.097</td>
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<td>Ag</td>
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<td>29.74</td>
<td>0.000</td>
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<tr>
<td>Ag+Eco</td>
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<td>31.49</td>
<td>0.000</td>
</tr>
<tr>
<td>Eco</td>
<td>203.11</td>
<td>33.45</td>
<td>0.000</td>
</tr>
<tr>
<td>Null</td>
<td>204.56</td>
<td>34.90</td>
<td>0.000</td>
</tr>
<tr>
<td><strong>Axis 2</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
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<td>Ag</td>
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<td>0.482</td>
</tr>
<tr>
<td>Ag+River</td>
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<td>1.52</td>
<td>0.225</td>
</tr>
<tr>
<td>Ag+Eco</td>
<td>218.52</td>
<td>1.84</td>
<td>0.192</td>
</tr>
<tr>
<td>Ag+Eco+River</td>
<td>219.90</td>
<td>3.22</td>
<td>0.096</td>
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<tr>
<td>Eco</td>
<td>227.58</td>
<td>10.90</td>
<td>0.002</td>
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<tr>
<td>Null</td>
<td>229.16</td>
<td>12.48</td>
<td>0.000</td>
</tr>
<tr>
<td>River</td>
<td>229.93</td>
<td>13.25</td>
<td>0.000</td>
</tr>
</tbody>
</table>
Table S2.1. Details of the microsatellite loci to be used in this study. Each multiplex set consisted of three loci. Locus amplification conditions used in this study are listed as footnotes.

<table>
<thead>
<tr>
<th>Locus Name</th>
<th>Multiplex Set</th>
<th>Primer Sequence</th>
<th>Primer (pmol)</th>
<th>Size Range</th>
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<td>1</td>
<td>F: 6-FAM-TGCATAACGCTCCCTGAAAG&lt;br&gt;R: GTGGTAGATGCTGGGATAGT</td>
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<tr>
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<td>1</td>
<td>F: HEX-GACACAGCAAACCTTTCTTCC&lt;br&gt;R: CATCTGGAAGTAGGCAATAAT</td>
<td>5</td>
<td>169-191</td>
</tr>
<tr>
<td>Mel1</td>
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<td>5</td>
<td>240-276</td>
</tr>
<tr>
<td>Mel101</td>
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<td>F: 6-FAM-ACGGTCCAACCATGATGAT&lt;br&gt;R: ACAATGGGAAGGTTCTCTCTT</td>
<td>8</td>
<td>119-149</td>
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<tr>
<td>Mel108</td>
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<td>F: 6-FAM-GTCTGGAGCCCATGTT&lt;br&gt;R: TCTTTGGAATGGAAGTTATGG</td>
<td>5</td>
<td>330-360</td>
</tr>
<tr>
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<td>2</td>
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<td>10</td>
<td>387-411</td>
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<tr>
<td>Tt-1</td>
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<td>Tt-2</td>
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<td>194-212</td>
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<td>Ma-1</td>
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<td>F: 6-FAM-ATTTATGTGCTGGGTCTA&lt;br&gt;R: TTATGCGTCTCTGTTTGCTCA</td>
<td>5</td>
<td>190-206</td>
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<td>Tt-3</td>
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<td>F: HEX-GGTGAGACCTGGAAATAGAAA&lt;br&gt;R: GCTAACCACACCTACGCAATGAT</td>
<td>8</td>
<td>147-179</td>
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<tr>
<td>Tt-4</td>
<td>4</td>
<td>F: 6-FAM-GGTGAGACCTGGAAATAGAAA&lt;br&gt;R: GCTAACCACACCTACGCAATGAT</td>
<td>5</td>
<td>164-196</td>
</tr>
<tr>
<td>Mvis072</td>
<td>4</td>
<td>F: HEX-CTGCAAGCTTGAATGGGAG&lt;br&gt;R: CCACACACTGGAAGTTTACGCA</td>
<td>8</td>
<td>297-339</td>
</tr>
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</table>

*Thermocycler profile for multiplex sets 1 and 2: Initial denaturation step at 94°C for 5 min; 21 cycles of 94°C for 1 min, 65°C for 1 min (-0.5°C per cycle), 72°C for 1 min; 15
cycles of 94°C for 1 min, 50°C for 1 min, 72°C for 1 min; final extension at 72°C for 2 min; soak at 60°C for 45 min.

bThermocycler profile for multiplex sets 3 and 4: Initial denaturation step at 94°C for 5 min; 30 cycles of 94°C for 30 s, 58°C for 30 s, 72°C for 30 s; final extension at 72°C for 10 min; soak at 60°C for 45 min.
Literature Cited


edge of their current distribution in North America with implications for genetic viability. *Conservation Genetics, 7*, 197–211.


Resources, 10, 831–844.


QIAGEN (2006) User developed protocol: purification of total DNA from nails, hair, or feathers using the DNeasy® Blood and Tissue Kit. QIAGEN-DYO4-Aug06. (accessed on 5 August 2010).


CHAPTER 3: RANGE-WIDE GENE FLOW IN A WIDESPREAD, SPECIALIST CARNIVORE, THE AMERICAN BADGER (TAXIDEA TAXUS)

Introduction

Contemporary species distributions are the direct result of the geological, climatic, and ecological conditions that influenced evolution across landscapes. In North America, evidence suggests that climatic oscillations during the Pleistocene represent a major evolutionary force in North America, particularly for temperate species. Taxa in northern latitudes were displaced by glacialations and restricted to one or more refugia, which often results in genetic variation that reflects population fluctuations and isolation within refugia during the Last Glacial Maximum (LGM; Hewitt 2000). Comparative studies in biogeographic hotspots such as the Pacific Northwest (Shafer et al. 2010) have identified common ecological factors that impact how co-distributed species responded to Pleistocene glaciation. Among these ecological factors, habitat specialization appears to play a key role in location and number of refugia occupied as well as patterns of recolonization after glaciation, a finding that is consistent with Pleistocene fossil records (e.g., Graham et al. 1996).

Pleistocene glaciations displaced many temperate species, and resultant distributional ranges and genetic structure in taxa with strong habitat preferences often closely mirror the changes that occurred within their preferred habitat type. For example, many forest-associated species exhibit eastern and western clades within North America because ice sheets, topographic features like the Rocky Mountains, and aridification
within the Great Plains separated areas of suitable forest (e.g., Wooding & Ward 1997, Weir & Schluter 2004, Runck & Cook 2005). While the Great Plains acted as a barrier to forest species, many grassland specialists were restricted to central North America and then expanded when habitats became more suitable during the Holocene (e.g., Johnson et al. 2003; Oyler-McCance et al. 2005; Wisely et al. 2008). According to fossil records during the climatic oscillations of the Quaternary, at the LGM, assemblages of species within habitats were highly predictable because they were largely defined by similar habitat preferences (Graham et al. 1996). For strict specialists, the distribution of preferred habitats likely continued to be the dominant driver of dynamics until contemporary time scales because strict specialists cannot persist outside their suitable habitat. Consequently, many strict specialists exhibit genetic structure that reflects both historical and contemporary changes in their preferred habitat (Curtois et al. 2003; Wisely et al. 2008; Shafer et al. 2011; Trumbo et al. 2013).

Outside strict specialists, previously defined assemblages of organisms during the LGM became highly variable following glacial recession (Graham et al. 1996), presumably because other species-specific traits contributed to post-glacial expansions into newly available habitats. For example, high dispersal capabilities allow individuals to traverse potential barriers and colonize new habitats quickly, so many highly mobile species exhibit patterns associated with rapid expansion following glacial recession (Vila et al. 1999; Latch et al. 2009; Reding et al. 2012). High dispersal capabilities also can facilitate varying levels of homogenization of previously isolated refugial populations (Latch et al. 2009; Smith et al. 2011) and even limited genetic structure in specialists within heterogeneous landscapes (e.g., Garroway et al. 2011; Centeno-Cuadros et al.
However, high dispersal capabilities do not always prevent genetic differentiation, particularly when accelerated genetic drift (i.e., low effective population size; Schwartz et al. 2009; Weckworth et al. 2014) or limited realized dispersal (e.g., natal habitat biased dispersal; Sacks et al. 2005, isolation-by-ecology; Musiani et al. 2007; Louis et al. 2014) resist the homogenizing effects of gene flow. Therefore, even if species retain signatures of past isolation within glacial refugia that are common among species with similar habitat preferences (e.g., Aubry et al. 2009; Barton et al. 2012; Reding et al. 2012; Latch et al. 2014), predicting overall genetic structure is difficult due to the species-specific responses that have occurred according to changes in habitat following the LGM into contemporary timescales.

One group that exemplifies the difficulty in understanding how habitat specialization influences gene flow are mammalian carnivores because they typically have large geographic ranges and high dispersal capabilities combined with low population densities and substantial variation in morphology and behavior throughout their range. Large geographic ranges and high dispersal capabilities predict extensive gene flow whereas low densities and morphological/behavioral variation suggest high rates of genetic drift and differential selection, respectively. Many carnivores are subject to intense management, so understanding genetic structure in carnivores has received considerable attention in the literature, particularly in forest or arctic-associated species (e.g., Schwartz et al. 2009, Short Bull et al. 2011; Koen et al. 2012; Reding et al. 2012; Row et al. 2012). From these studies, a wide array of responses to habitat heterogeneity have been recorded including differentiated ecotypes (Musiani et al. 2007), landscape variables driving genetic differentiation (Schwartz et al. 2009; Short Bull et al. 2011;
Reding et al. 2012), and continent-wide patterns of isolation-by-distance (Tammeleht et al. 2010; Row et al. 2012). Unlike forest and arctic-associated carnivores, grassland species have received considerably less attention in genetic studies (but see Wisely et al. 2008), yet they can provide insight into how changes during and following the LGM such as forest encroachment and massive conversion to agriculture have contributed to genetic variation.

This study focused on the American badger (Taxidea taxus), a semifossorial, grassland-associated mesocarnivore that has a large geographic range encompassing most of central and western North America where suitable grassland habitats occur. Along with strong preferences for grassland habitats, their semifossorial lifestyle makes badgers dietary specialists as the vast majority of their diet is composed of available burrowing mammals like prairie dogs and ground squirrels (Messick and Hornocker 1981; Goodrich and Buskirk 1998; Sovada et al. 1999; Azevedo et al. 2006). Evidence for differentiation in badgers across their range is reflected in subspecific designations as morphological and behavioral differences define four subspecies (T. t. berlandieri, T. t. jacksoni, T. t. jeffersonii, and T. t. taxus; Long 1972; Figure 3.1). Each subspecies roughly corresponds to grassland habitat types (prairie, scrub-steppe, and oak savannas) or large physical barriers (Rocky Mountains, Grand Canyon; Long 1972). Taken together, specialist traits and apparent differences according to habitat (i.e., subspecific designation) predict limited gene flow between habitat types.

Although American badgers possess traits that suggest habitat-based differentiation, their large dispersal capabilities and geographic range are more typical of a generalist. Their large geographic range greatly exceeds other grassland specialists and
includes a number of hypothesized glacial refugia for mammals (Great Plains; Wisely et al. 2008, Pacific Northwest; e.g., Latch et al. 2009, Shafer et al. 2010; Great Basin and California; Howard & Swenson 2005). Pleistocene badger fossils have been found throughout North America (Long 1972), suggesting that badgers may not have been as limited in distribution as other highly specialized species during the LGM. Badgers are also highly mobile as demonstrated by their seasonally large home ranges and dispersal distances (125 km+; Messick & Hornocker 1981). Therefore, high mobility and corresponding gene flow in badgers could theoretically override any isolation created by habitat preferences or Pleistocene glaciation.

In this study, we provide a continent-wide assessment of genetic structure within the American badger. Our primary goal was to investigate if specialization on grassland habitats impacted dynamics during Quaternary glaciation cycles and contemporary gene flow. If specialization on grasslands drives gene flow patterns in badgers, badgers would be expected to have experienced a restricted distribution in the Great Plains (sensu Johnson et al. 2003; Oyler-McCance et al. 2005; Wisely et al. 2008) with subsequent rapid recolonization following glacial recession. Also, subspecies designations should be correlated with genetic differentiation if specialization impacts contemporary gene flow. Conversely, if high mobility overrides any effect of specialization, badgers are expected to exhibit limited genetic differentiation except when confronted with large topographic barriers. To test these predictions, we sampled 917 badgers from across North America representing all four subspecies to evaluate the effect of topographic barriers, geographic distance and ecological variables on genetic differentiation. This broad scale genetic survey can aid in understanding how ecological traits such as specialization and mobility
interact to produce gene flow patterns in grassland species. Additionally, specialists like badgers are vulnerable to fragmentation (Crooks 2002) and grasslands have undergone intense conversion to agriculture, so observed genetic patterns may also provide insight into how fragmentation has affected connectivity in badger populations.

Methods

Sample Collection

We collected 917 American badgers from 2001-2013, representing all four described subspecies and the majority of the North American range (Figure 3.1). Samples included tissue (n = 520) and skin (n = 397) collected from road-killed and fur-trapped animals (Table S3.1). All samples were georeferenced, either as latitude/longitude coordinates recorded at the time of sample collection or estimated based on written descriptions of the sampling locations. Exact locations for location descriptions (nearest city, state/province, Public Land Survey System [PLSS]) were calculated in ArcGIS 10.1 using the “create random points” function either within a political boundary (PLSS, state, county, or province) or within 20km circular buffers around a city center (buffer radius = 5, 10, 20, or 50 km yielded concordant results).

Laboratory Methods

DNA was extracted from all samples and all sample types using the Qiagen DNEasy Blood and Tissue Kit. A 600 bp segment of the mitochondrial displacement loop (d-loop) was amplified using newly designed primers (Tax-dloopF 5’-
CGGGGTCTTTGTGACTCTTCT-3’ and Tax-dloopR 5’-CAGCACCCAAAGCTGATATTC-3’). PCR amplifications occurred in 10 uL reactions with 10 ng of genomic DNA, 5 pmol of each primer, 0.2 mM of each dNTP, 0.75 U of PerfectTaq DNA polymerase, and 1 X PerfectTaq PCR Buffer. Thermocycler conditions for mtDNA included a 2-min initial denaturation step at 94°C, 30 cycles of 30 second denaturation (94°C), annealing (60°C), and extension (72°C) steps, and final extension at 72°C for 5 min. Each 10 uL sequencing reaction contained 50 ng of template, 0.3 pmol of forward primer, 0.5 uL of ABI Big Dye in 2 X reaction buffer. Thermocycler conditions followed the manufacturer’s recommendations, except 60 cycles were used to improve signal strength. Cleaning followed a modified low sodium precipitation method (Latch & Rhodes 2005), and then cleaned products were sequenced on an ABI3730 DNA Analyzer at the University of Wisconsin Biotechnology Center. All sequences were analyzed using the program GENEIOUS PRO v. 5.2.

For quality control, we re-extracted and re-sequenced all samples yielding unique haplotypes (n = 54), all samples containing a 25 bp deletion (n = 56), and a randomly selected subset of remaining badgers (n = 103). We observed four sequencing errors (4/213 = 1.88% error rate); three in individuals with unique haplotypes and one in an individual with a common haplotype. A representative sequence for each unique haplotype was deposited in Genbank. Our final mtDNA dataset contained 785 badgers with mtDNA sequences, which excluded all individuals with the 25 bp deletion region (n = 14 haplotypes with deletion).

We amplified all samples at 12 microsatellite loci developed in American badger (Taxidea taxus; Tt-1, 2, 3, and 4; Davis & Strobeck 1998), American mink (Neovison
vison; Mvis072; Fleming et al. 2002), American marten (Martes americana; Ma-1; Davis & Strobeck 1998), and European badger (Meles meles; Mel112, Mel101, Mel111, Mel108, Mel14, and Mel1: Carpenter et al. 2003; Domingo-Roura et al. 2003). PCRs were conducted in four multiplex reactions containing 3 primers each (conditions described in Chapter 2). Each 10 uL reaction contained 10 ng of genomic DNA, 3-8 pmol each primer, 0.2 mM of each dNTP, 0.75 U of PerfectTaq DNA polymerase, and 1 X PerfectTaq PCR Buffer. Amplified products were genotyped on an ABI3730 DNA Analyzer at the University of Wisconsin Biotechnology Center, and alleles were sized using the program GeneMarker (SoftGenetics LLC, State College, PA, USA).

For quality control, we re-genotyped all individuals that had rare alleles (i.e., occurred in fewer than 2% of samples; n = 17) and those in mtDNA quality control (n = 213) to calculate error rates and confirm rare alleles. These repeated PCRs and amplifications confirmed all rare alleles and detected one mismatch due to probable allelic dropout (error rate = 1/2760; 0.0036%). All individuals (n = 917) were successfully genotyped at 10 or more loci (11/11,004 total genotypes missing; 0.10% missing).

**Historical Gene Flow**

To investigate historical patterns in gene flow, we generated phylogenetic trees using three algorithms (Maximum Parsimony; MP, Maximum Likelihood; ML, and Bayesian) implemented in GENEIOUS. The GENEIOUS plug-in for PAUP (Swofford 2003) and MrBayes (Ronquist et al. 2012) performed calculations identical to their parent programs with pre-defined parameters. We defined gaps as a fifth state and used the 50%
majority rule consensus trees for all three algorithms. jModelTest 2 (Darriba et al. 2012) suggested that the GTR+G+I mutation model (Tavaré 1986) with a gamma shape of 0.49 best fit our data. A Meles meles d-loop sequences (AM711900.1) served as the outgroup for all analyses. Support nodes within MP and ML (bootstraps) as well as Bayesian trees (posterior probabilities) were calculated after 1000 permutations.

Traditional tree-building techniques can yield unresolved intraspecific trees (Posada & Crandall 2001), so a haplotype network was constructed to better visualize relationships among haplotypes. The network was assembled in the program TCS (Clement et al. 2000) based on maximum parsimony. We used a 95% connection limit with gaps as a 5th state. The resultant haplotype network connected all haplotypes into a single network.

To complement tree-building and network analyses, we also tested for differentiation in mtDNA using a spatial analysis of molecular variance (SAMOVA; Dupanloup et al. 2002). SAMOVA identifies the number of groups of populations (K) that maximizes \( \phi_{CT} \) or total genetic variance explained by differentiation between groups. SAMOVA requires individuals be grouped into populations, so we grouped badgers into a priori populations (n = 19 or 23; Table S3.1). For half of the Canadian samples, we had only location of fur deposit, not the specific trapping location, so SAMOVAs were run with either all Canadian badgers pooled as ‘Canada’ or only including individuals that had specific trapping locations (Saskatchewan = 27, Manitoba = 5, Alberta = 5). To generate geographic coordinates for each population, we drew 100% minimum convex polygons around geo-referenced individuals within each state or Canada via the Minimum Bounding Geometry function in ArcGIS. Latitude and longitude points were
calculated for the centroid of each polygon, and used as the geographic coordinates for each population within the SAMOVA. We tested $K = 2$ through 10 using the program SAMOVA v. 1.0 (Dupanloup et al. 2002).

We used Arlequin 3.5 (Excoffier & Lischer 2010) to calculate basic diversity metrics for the total dataset, four subspecies, and SAMOVA groups. Diversity metrics for each group included haplotype diversity ($h$), nucleotide diversity ($\pi$), and number of pair-wise differences. Differentiation ($\phi_{ST}$) between SAMOVA groups and subspecies were also quantified in Arlequin.

To test for evidence of postglacial demographic expansion, we used Arlequin to calculate three complementary statistics. Fu’s $F_S$ (Fu 1997) and Tajima’s $D$ (Tajima 1989) were calculated for the total dataset, subspecies, and SAMOVA groups. Significantly negative values of $F_S$ and $D$ indicate rapid demographic expansion. We further evaluated evidence for postglacial expansion by constructing mismatch distributions of nucleotide differences for all individuals, subspecies, and SAMOVA groups. Under constant population size, mismatch distributions exhibit a ragged pattern whereas recently expanded populations have a unimodal distribution (Slatkin & Hudson 1991; Rogers & Harpending 1992). To assess fit of the observed nucleotide differences to expected distributions under sudden demographic or spatial expansion, we calculated two statistics: raggedness index (Rogers & Harpending 1992) and sum of squared differences (SSD; Schneider & Excoffier 1999) in Arlequin.

Another method to examine recolonization dynamics following glacial recession is through simulations that attempt to reconstruct demographic histories (e.g., Heled & Drummond 2008). However, the requirement for specific priors, particularly lineage
specific mutation rates or fossil calibration points, would have yielded considerable uncertainty in badgers. When we calculated mutation rates for badgers, a highly divergent, monotypic lineage in Mustelidae (approximately 20 mya diverged from common ancestor of remaining mustelids; Koepfli et al. 2007), mutation rates were very low (e.g., 3.835 X 10^{-9} bp/generation using Meles clade) compared to other estimates of d-loop in mammals (e.g., Koblmüller et al. 2012). Regions of d-loop typically have a higher mutation rate compared to the rest of the mitogenome and tend to have species-specific mutation rates (Pesole et al. 1999). Therefore, we hypothesize that mutations within d-loop obscured the distinctiveness of the American badgers through homoplasy, similar to d-loop in fisher (Martes pennanti; Knaus et al. 2011). Inaccurate mutation rates has been recorded when using ancient divergences to calibrate nodes for demographic reconstruction (e.g., Ho et al. 2005; Navascués & Emerson 2009), and using different mutation rates or fossils for calibration can give radically divergent conclusions (e.g., Ho et al. 2005; Shapiro & Ho 2014). Based on the likely inaccurate mutation rate for d-loop and little differentiation among haplotypes, we chose not to estimate divergence times and demographic parameters in this dataset.

Contemporary Gene Flow

Two Bayesian programs, non-spatial STRUCTURE 2.2.3 (Pritchard et al. 2000) and spatial BAPS 5 (Corrander et al. 2008), were run to examine contemporary population structure of badgers in North America. In our non-spatial approach, we performed clustering in STRUCTURE for all 917 individuals for 5 iterations of each value of $K$ from $K = 1 – 15$. Each run consisted of 100,000 replicates of the MCMC after a burn-in of
100,000 replicates. We used both likelihoods and Evanno *et al.* (2005)’s $\Delta K$ method to determine the most likely $K$ since likelihoods plateaued and exhibited higher variances after the optimum $K$ is reached (Pritchard *et al.* 2000; Pritchard & Wen 2003). Once the optimum $K$ was identified, longer runs (1,000,000 MCMC burn-in, 1,000,000 permutations) were performed at $K-1$, $K$, and $K+1$ to verify results from the shorter runs. To assign individuals to inferred clusters, we performed a final set of 10 runs (1,000,000 burn-in, 1,000,000 stored replicates) at the optimal $K$. Each individual was assigned to the cluster in which it had the highest average probability of membership, based on averaged $q$-values (i.e., proportion of genome that belonged to each cluster) calculated in *Clumpp* (Jakobsson & Rosenberg 2007). The entire clustering process was repeated iteratively for each of the inferred clusters, in order to identify any additional substructure within the main genetic clusters.

In our spatially explicit approach, we employed Bayesian clustering in *BAPS 5* using the “spatial clustering of individuals” model. We performed five iterations for each value of $K$ for $K = 1$-15. The maximum likelihood and highest posterior probability were used to determine the optimum number of genetic clusters in the sample, and each individual was assigned to a cluster. We then performed an admixture analysis based on the results of the mixture clustering, using 100 iterations, 20 reference individuals per population, and 5 iterations per reference individual within clusters.

The Bayesian algorithms yielded the same $K$ following iterative runs in *Structure*, but the locations of genetic discontinuities between clusters were more defined in *BAPS*. Therefore, we utilized results from *BAPS* to examine genetic diversity and differentiation between putative clusters. Deviations from Hardy-Weinberg and
Linkage Equilibrium were quantified in GENEPOP (Raymond & Rousset 1995) for each cluster and the total dataset following corrections for multiple tests (false-discovery rate; Benjamini & Yekutieli 2001). We used the R package diveRsity (Keenan 2013) to calculate both genetic differentiation between clusters ($F_{ST}$) and genetic diversity metrics (heterozygosity, allelic richness, and $F_{IS}$).

A number of processes likely influence genetic variation in badgers across North America. One such process is isolation-by-distance (IBD; Wright 1941), a phenomenon where genetic differentiation is correlated with geographic distance. To test for IBD, we utilized simple Mantel tests that quantify the correlation (i.e. Pearson’s $r$) between genetic distances and Euclidean distances. Pair-wise genetic distances (Rousset’s $a$; Rousset 2000) were calculated in SPAGeDI (Hardy & Vekemans 2002) and Euclidean distances were calculated in ArcMap. Simple Mantel tests were performed in the R package vegan (function “mantel”; Oksanen et al. 2008) for the entire dataset and for each group identified in the Bayesian clustering analyses, and statistical significance was assessed after 1000 permutations.

In addition to IBD, we tested how three topographic barriers (Rocky Mountains, Mississippi River, and Lake Michigan) and subspecific designations contribute to observed patterns in genetic variation via three tests. Partial Mantel tests, principal components analysis (PCA), and spatial principal components analysis (sPCA) provide complementary methods to test for barrier effects. The first test, partial Mantel tests, control for the effects of geographic distances to evaluate correlations between each topographic barrier or subspecific designation and pair-wise genetic distances. For subspecies, predictor variables were coded as dummy variables (0 or 1) based on whether
individuals were of the same (0) or different (1) subspecies. We only tested differences between adjacent subspecies, resulting in four partial Mantel tests (T. t. jacksoni/taxus, T. t. taxus/jeffersonii, T. t. taxus/berlandieri, T. t. jeffersonii/berlandieri; Figure 3.1 for subspecies boundaries). For each partial Mantel test, only individuals within 300 km of the putative barrier or subspecies boundary were included to control for any confounding effects of geography or other barriers. Statistical significance was evaluated via Pearson’s correlation coefficients ($r$) after 1000 permutations in the R package vegan (function partial.mantel; Oksanen et al. 2008).

To complement the partial Mantel tests, we used PCAs to visualize the potential roles of all topographic barriers (Mississippi River, Rocky Mountains, and Lake Michigan), water barriers only (Mississippi River and Lake Michigan), and subspecific designations on gene flow. We summarized patterns in genetic diversity into linear axes called principal components in the dudi.pca function within the R package adegenet (Jombart et al. 2008). We accounted for missing data by filling in empty genotypes with mean allele frequencies using the “scaleGen” function. For each grouping (topographic barriers, water barriers, or subspecies), 95% inertia ellipses drawn around each group permit visual inspection of the differentiation among groups. In total, we drew 95% ellipses around individuals according to three groupings: topographic barriers (Mississippi River, Rocky Mountains, and Lake Michigan), water barriers only (Mississippi River and Lake Michigan), and subspecific designations (T. t. berlandieri, T.t. jacksoni, T. t. jeffersonii, and T. t. taxus).

sPCA provides principal components scores that summarize both the non-spatial genetic variation and the spatial autocorrelation structure among individual genotypes.
(Jombart et al. 2008). In this method, highly positive eigenvalues reflect axes with large variances and positive spatial autocorrelation (i.e., global patterns; Jombart et al. 2008). We performed a sPCA in adegenet using a distance-based connection network to ensure all individuals were connected by at least one segment. Badgers were unlikely to be completely genetically isolated, so this network is likely most realistic. To assess which axes are important for explaining genetic variation, we examined two outputs generated from the sPCA. Axes with the highest eigenvalues are considered the most important because they explain the most variance, so eigenvalues were plotted to examine differences among eigenvalues for each axis. Another visualization that aids in interpretation of sPCA results is a screeplot, a graph of spatial autocorrelation vs. variance explained for each axis. Axes that are well differentiated from other axes explain the most variation in genetic structure. Spatially lagged scores from axes that had the highest eigenvalues and were separated from most others in the screeplot were plotted in ArcMap to examine any geographic structuring within the sPCA. We conducted a permutation procedure with 1000 randomizations to test for a significant global ("global.rtest") pattern in the data.

Topographic barriers are likely to be important in shaping badger gene flow, but they may occur in tandem with other landscape or climate variables as observed in other carnivores (e.g., Garroway et al. 2011; Reding et al. 2012). Therefore, we constructed a full model of how landscape heterogeneity impacts gene flow by including landscape (soil, land cover, and precipitation) variables in addition to the three topographic barriers, subspecies, and geographic coordinates. The first set of landscape variables included soil parameters likely to be important for semifossorial badgers: soil texture (% clay, sand, or
silt) and soil order (5 total within sampled area: United States Department of Agriculture 1999) for each sample location. We also derived land cover (NCLD2006; Fry et al. 2011) and precipitation (average annual precipitation from WorldClim v. 1.4; Hijmans et al. 2005) data for each sample location. Subspecific designations roughly correspond to major habitat breaks defined by level I ecoregions, regions of similar biotic and abiotic factors within North America (Obernick 1987). Therefore, subspecific designations were included as predictor variables (dummy variables 1-4) as well as the three topographic barriers (Rocky Mountains, Mississippi River, and Lake Michigan) and geographic coordinates (latitude and longitude). In total, ten landscape variables (soil texture, soil order, land cover, average annual precipitation, subspecies, Rocky Mountains, Mississippi River, Lake Michigan, latitude, and longitude) served as predictors within distance-based redundancy analysis (dbRDA) models.

We used dbRDA, a multivariate technique analogous to linear regression, to evaluate the relationship between landscape predictor variables and pair-wise genetic distances (Rousset’s $a$). An initial marginal test analyzed all landscape variables in DISTLM v. 5 (Anderson 2004) while subsequent partial dbRDAs calculated the impact of every variable individually. Due to multiple tests, alpha was corrected (alpha = 0.015) for the partial dbRDAs when assessing significance. The program DISTLM FORWARD 3 (Anderson 2003) eliminated non-significant variables via forward selection to produce the best combination of predictors that explained genetic differentiation. By utilizing both partial tests that isolate each variable and forward selection that adds variables sequentially, we were able to control any effects of multicollinearity within our predictor variables.
Results

Historical Gene Flow

In total, 95 haplotypes were identified among 785 individual badgers (54 polymorphic sites, 53 substitutions). The mean number of pairwise differences among sequences was 3.660 (SD = 1.856) and the overall nucleotide diversity was 0.0062 (SD = 0.0035). No *a priori* groupings (four subspecies and two SAMOVA groups) exhibited significant $\phi_{ST}$ values (all $\phi_{ST} = 0.000$, $p = 0.999$), suggesting limited differentiation in mtDNA in the total sample.

The haplotype network showed some spatial structuring of mtDNA haplotypes. The overall pattern was dominated by at least two common haplotypes surrounded by similar haplotypes, producing star-like patterns (Figure 3.2). The most common haplotype (Haplotype 1, $n = 257$, 32.7%) occurred throughout North America whereas the second most common (Haplotype 3; $n = 149$, 19.0%) was primarily found in northern latitudes west of Wisconsin (i.e., North Dakota, Montana, and Canada). The last two common haplotypes (Haplotype 2, $n = 96$, 12.2%; and Haplotype 4, $n = 30$, 3.8%) were a single mutation step away from Haplotypes 1 and 3 respectively. Haplotype 2 was largely restricted to Michigan and Ohio (76/96 occurrences) whereas Haplotype 4 was primarily found west of the Rocky Mountains (28/30 occurrences; Figure S3.1). Both Haplotypes 1 and 2 were shared between all four subspecies while Haplotype 3 only occurred in *T. t. jacksoni* and *T. t. taxus* (Figure 3.2a). Haplotype 4 was largely concentrated within *T. t. berlandieri* and *T. t. jeffersonii*. All other haplotypes had a frequency of 10 individuals (1.3%) or less. Despite Haplotypes 1 and 2 exhibiting star-
like patterns often consistent with recent bottlenecks followed by rapid expansion, little differentiation was observed between mitochondrial haplotypes. All tree-building approaches produced phylogenograms that had a comb-like topology where only the outgroup (*Meles meles*) received high support (bootstraps > 80, posterior probabilities > 0.80), further supporting limited differentiation among haplotypes.

The SAMOVA found the optimum split in haplotypes occurred at $K = 2$ regardless if Canadian provinces were pooled or separate, which corresponded to the Lower Peninsula of Michigan (LP), Indiana, and Ohio versus all other individuals (Table 3.1). Additional evidence for $K = 2$ being the best grouping to explain variance in our mtDNA was that the fixation index ($\phi_{CS}$) peaked at $K = 2$, and in simulated datasets, $\phi_{CS}$ reached its maximum at the appropriate $K$ (Dupanloup *et al.* 2002). The cause of differentiation was driven by the high frequency of Haplotype 3 within the LP, Indiana and Ohio (Figure 3.2b).

All three tests for population expansion supported rapid population expansion in the entire dataset, each subspecies, and western SAMOVA groups. Fu’s $F_s$ ($F_s = -24.197, p < 0.001$) and Tajima’s $D$ ($D = -1.395, p = 0.043$) were significantly negative in the total dataset whereas only Fu’s $F_s$ remained significant in subspecies and SAMOVA groups (all $p < 0.002$; Table 3.1). Both raggedness indices (all $r < 0.0928, p > 0.052$) and SSD (all SSD < 0.313, $p > 0.19$) suggested that the observed mismatch distributions followed predicted distributions under rapid or spatial expansion for all except the eastern SAMOVA group (Figure 3.3). Raggedness and SSD for the eastern SAMOVA group rejected both rapid and spatial expansion ($r = 0.209, p = 0.002$; SSD = 0.028, $p = 0.008$). However, the number of haplotypes within the Eastern SAMOVA group was very small.
(n = 5), suggesting that the statistically significant results in raggedness and SSD could be due to low sample size. Metrics based on mismatch distributions tend to be less powerful than Fu’s $F_S$ (Ramos-Onsins & Rozas 2002), so the significant Fu’s $F_S$ indicates that the significance in raggedness and SSD was due to low sample size.

**Contemporary Gene Flow**

The optimal solution in BAPS was $K = 3$ for our badger dataset corresponding to three groups: i) Lower Peninsula of Michigan (LP), ii.) Wisconsin, Upper Peninsula of Michigan, and parts of Iowa (WI), and iii.) west of the Mississippi River (West; Figure 3.4). STRUCTURE initially detected two clusters that corresponded to the eastern United States and Canada and all other individuals (Figure S3.2). Iterative runs of the eastern cluster detected a split between the LP and all other individuals whereas the western cluster exhibited no additional substructure (Figure S3.3). The overall pattern of the three clusters with admixture within the upper Midwestern USA and Canada were similar between STRUCTURE and BAPS; disagreements on assignments largely occurred within contact zones between clusters (e.g., within Wisconsin, Minnesota, and Iowa). Therefore, the disagreement in individual assignments likely reflects either a weak barrier (Latch *et al.* 2006) or IBD because both programs can yield discrete clusters when only IBD is present (Frantz *et al.* 2009). Assignments in LP, however, were largely consistent between programs where all but three (BAPS) to five (STRUCTURE) individuals were assigned to the LP cluster.

Microsatellite diversity averaged 13.83 alleles/locus (range: 9-20 alleles). A heterozygote deficiency was recorded in the total dataset ($F_{IS} = 0.090, p < 0.001$),
indicating the presence of genetic structure within our dataset. One source of structure that was evident was IBD across North America (simple Mantel test $r = 0.137$, $p < 0.001$). Furthermore, all three clusters detected in Bayesian programs (BAPS or STRUCTURE) exhibited low but significant levels of differentiation ($F_{ST} = 0.023$-$0.072$; all $p < 0.001$). LP was more strongly differentiated from West ($F_{ST} = 0.072$, SE = 0.021) and WI (0.052, SE = 0.012) than West versus WI (0.023, SE = 0.011). In addition to the stronger differentiation of LP from West and WI, LP also had significantly lower genetic diversity than the other two clusters (paired t-tests: all $t > 5.17$, $p < 0.001$).

LP, West, and WI clusters each exhibited heterozygote deficiencies similar to the full dataset ($F_{IS} = 0.089$-$0.092$, $p < 0.001$). Like the full dataset, all clusters had significant IBD according to simple Mantel tests ($r = 0.054$-$0.13$, all $p < 0.001$). The internal IBD within clusters combined with observed admixture between clusters, particularly between West and WI, are likely causes for the observed heterozygote deficiencies.

We recorded evidence for barrier effects in partial Mantel tests, PCA, and sPCA. Partial Mantel tests detected both Lake Michigan ($r = 0.152$, $p = 0.001$) and Mississippi River ($r = 0.065$, $p = 0.001$) as barriers to dispersal. In contrast, the Rocky Mountains and subspecific pairings were not significant (all $r < 0.023$, $p > 0.067$).

PCA also supported that Lake Michigan and Mississippi River were barriers to dispersal because groupings based on large aquatic barriers (i.e., Lake Michigan and Mississippi River) fit the data better than groupings based on subspecific definitions (Figure 3.5). Badgers east of Lake Michigan were the most genetically distinct according to the scatterplot for the first two PCA axes (Figure 3.5a). Considerable overlap in 95%
inertia ellipses were observed between those east and west of the Mississippi River, but the two groups exhibited some differentiation as compared to subspecies. Little resolution was observed among the four subspecies except for *T. t. jacksoni*, which was largely due to the presence of those east of Lake Michigan (Figure 3.5b).

Results from the sPCA were largely concordant with previous tests that showed genetic variation is dictated by both IBD and topographic barriers. The first two sPCA axes explained the most variance and had the highest eigenvalues (eigenvalues = 0.17 and 0.049; all others < 0.037), so they were considered most important in explaining genetic variation (Figure 3.6a). Both main sPCA axes exhibited high levels of spatial autocorrelation (Moran’s I = 0.75 and 0.32 for axes 1 and 2 respectively; Figure 3.6b), indicating IBD. The first axis explained the most variance in the dataset (0.251) and exhibited an east-west cline with the LP having the most extreme values (Figure 3.6c). Axis 2 explained less variance than Axis 1 (0.150), but in general, individuals in the Upper Midwest were differentiated from the rest of the dataset (Figure 3.6d). Monte-Carlo tests indicated that at least one global structure (i.e., those with positive eigenvalues) observed in the sPCA was significant (p = 0.0001).

After forward selection, the dbRDA detected four factors that influence gene flow. The resultant model explained 10.11% of the total variation in genetic distances across North America. In the partial dbRDAs (i.e., each variable was isolated by partialling out all others), Lake Michigan explained the most variation (1.86%) in the data with Latitude as the second most important factor (1.61%). Longitude (0.91%) and Mississippi River (0.61%) were the final two significant factors (Table 3.5). None of the
landscape variables we tested (soil texture, soil order, land cover, precipitation) were significant in the partial tests.

**Discussion**

Badgers have strict habitat preferences that are predicted to limit gene flow outside of grassland habitats with soils suitable for digging, and to create genetic structure that is tightly associated with preferred habitats. However, badgers also exhibit high mobility and have a vast distribution that are expected to facilitate gene flow regardless of habitat and resist accumulation of spatial genetic structure. Therefore, it was unclear how grassland specialization and high mobility would interact to shape patterns of genetic variation across North America. Our data revealed broad-scale spatial genetic structure within both mitochondrial and microsatellite datasets. Differentiation was weak within mtDNA, but we recorded geographic structuring among haplotypes and a split between the Lower Peninsula, Ohio, and Indiana and the rest of the dataset. Microsatellites had more resolution than mtDNA where both individual-based and population-based approaches detected IBD and additional substructure within the Midwestern United States. Based on the geographic structuring in mtDNA and strong signatures of IBD in all microsatellite analyses, the primary influence on gene flow was geographic distance. Prominent water barriers, especially Lake Michigan, were also important in explaining patterns of genetic variation. The strongest genetic differentiation in both markers occurred between areas east and west of Lake Michigan whereas only microsatellites detected the Mississippi River as barrier to gene flow. With
IBD and water barriers being the prominent influences on gene flow, high mobility appears to be a more powerful influence than ecological specialization on broad-scale genetic patterns in badgers.

**Historical Gene Flow**

Mitochondrial DNA revealed high diversity, little differentiation among haplotypes, and weak geographic structuring throughout North America in badgers. Little differentiation among haplotypes and the presence of a few common haplotypes surrounded by rare haplotypes (i.e., star-like topology; Slatkin & Hudson 1991) within the haplotype network suggest that badgers did not occupy multiple, isolated glacial refugia during the LGM. Persistence in multiple glacial refugia is fairly typical for high dispersal species (e.g., Shafer et al. 2010; Stewart et al. 2010), but isolation within separate refugia typically yields well-defined groups of haplotypes that are separated by multiple mutational steps (e.g., Aubry et al. 2009; Latch et al. 2009; Lait & Burg 2013; van Els et al. 2014). The pattern of mitochondrial variation we observed more closely resembles patterns described for grassland species that were restricted to the Great Plains (Johnson et al. 2003; Oyler-McCance et al. 2005; Wisely et al. 2008; Koblmüller et al. 2012) or other highly mobile species that were not fragmented during the LGM (Carmichael et al. 2007; Teacher et al. 2011; Pulgarin-Restrepo & Burg 2012).

Dynamics following glacial recession in badgers appear to be more complex than other grassland specialists because the SAMOVA suggested geographic structuring in haplotypes and Quaternary fossils of badgers have been found in numerous locations outside the Great Plains (Long 1972). Our mitochondrial dataset was consistent with
badgers being somewhat widespread below the ice sheets and then multiple invasions into glaciated areas from different locations or time periods. Support for the widespread distribution of badgers during the LGM was also seen in the limited genetic differentiation, high genetic diversity and star-shaped haplotype networks because these characteristics are consistent with high effective population size (Crandall & Templeton 1993) without barriers to gene flow. The limited phylogeographic structure found in badgers greatly resembles European populations of red fox (*Vulpes vulpes*), a highly mobile, generalist carnivore thought to remained continuous during the LGM (Teacher et al. 2011). High gene flow in badgers, in contrast, was surprising given that grasslands underwent substantial changes during and following the LGM (Axelrod 1985; Whitlock 2000). Grasslands contracted during the late Pleistocene due to climate cooling (Axelrod 1985) and some areas (e.g., Columbia Basin and eastern Beringia; Whitlock 2000) completely shifted to forested habitats. Invasion of forests into grasslands did not necessarily mean range contraction for badgers, however, because many areas maintained mixes of forest and grasslands that are still suitable habitats for badgers (e.g., Duquette & Gehrt 2014). Thus, the variable and heterogeneous habitats below the ice sheets may have remained suitable for badgers, allowing for broad-scale movement and retention of genetic diversity.

Despite the dynamic history of grasslands and numerous topographic barriers in North America, the SAMOVA only split the dataset into samples east and west of the Lake Michigan. Western populations appear to have expanded north from southern areas such as Utah and the Great Plains due to strong signatures of population expansion in demographic tests and high diversity below ice sheets. Expanded populations, in
contrast, typically exhibit decreased genetic diversities and persistence of common haplotypes at high frequencies due to repeated founder effects (e.g., Hewitt 2000). Ethier et al. (2012) recorded much more substructure in northern latitudes than this study, which would occur if northern populations experienced bottlenecks associated with colonization from southern populations. The decreased genetic diversity and high rates of drift would create higher genetic differentiation even without physical barriers to gene flow. When we included southern localities that had substantially higher diversity, we did not find any discrete groups despite similar geographic structuring of Haplotypes 2 and 4. Therefore, our data indicates that most of western North America maintained high effective population sizes and gene flow that prevented strong differentiation in mtDNA, and founder effects occurred in northern latitudes resulting in high frequencies of Haplotype 2.

Mechanisms shaping geographic structuring of haplotypes within the Midwest were apparent, with populations exhibiting clear signatures of one or more population expansions. Previously glaciated regions west of Lake Michigan (e.g., Wisconsin and Minnesota) exhibited high frequencies of Haplotype 1, consistent with expansion from the Great Plains. The presence of Haplotype 3 in Ohio and the LP, in contrast, suggests two potential colonization scenarios for the LP. First, badgers could have colonized eastern North America once and Lake Michigan could have isolated badgers in the LP, resulting in the split between eastern (i.e., LP, Ohio, and Indiana) and western badgers observed in the SAMOVA. Haplotype 3 is one transition away from Haplotype 1, the haplotype most common in surrounding states (i.e., Minnesota, Wisconsin, and Iowa), so Haplotype 3 could have arisen post colonization and then Lake Michigan prevented the
spread of Haplotype 3 back into western North America. Alternatively, a separate colonization in the LP caused the high frequency of Haplotype 3 in the LP. Guilday (1968) suggested that eastern North America was invaded twice by prairie-associated species based on fossils in both pre-and post-Wisconsin deposits in Kentucky, Maryland, and Pennsylvania. Multiple colonization routes into the Great Lakes region has been recorded in herpetofauna (Austin et al. 2002; Zamudio & Savage 2003; Placykr Jr et al. 2007), resulting in disparate frequencies of haplotypes between the LP and surrounding states much like Haplotype 3 in this study. The few mammal studies in the Great Lakes region are largely limited to forest-associated species (Rowe et al. 2004; Taylor & Hoffman 2010; Reding et al. 2012), recent invaders (Koblmüller et al. 2012), or those not found in the Lower Peninsula (Koblmüller et al. 2009; vonHoldt et al. 2011), so it is difficult to determine if multiple colonization routes also occurred in mammals including badgers. Both mechanisms, a founder effect followed by isolation or two colonization routes, would result in the high frequency of Haplotype 3 and the corresponding split between eastern (i.e., LP, Ohio, and Indiana) and western North America. Therefore, further studies could help differentiate between these two mechanisms via hypothesis testing of separate recolonization scenarios (e.g., Frantz et al. 2014) with alternative markers that allow divergence time estimation.

Contemporary Gene Flow

Grasslands have undergone massive changes following the LGM including widespread conversion to agriculture. Conversion to agriculture has reduced the amount of grasslands and increased fragmentation among grassland patches, resulting in
demographic declines and reduced movement between remaining patches in many grassland-associated taxa (e.g., Coppedge et al. 2001; Steffan-Dewenter & Tscharntke 2002; Wisely et al. 2008). Genetic differentiation can occur if individuals cannot reach distant grassland patches, and resultant genetic variation would be correlated with the presence of grassland habitat. Our microsatellite dataset did not follow this expectation because we detected both IBD and topographic barriers only in all analyses. Instead, patterns of genetic structure within our microsatellite markers were remarkably similar to our mitochondrial dataset (i.e., geographic structuring and strong divergence within the Great Lakes region). Therefore, high dispersal appears to be the primary force in driving patterns of genetic variation across North America.

The overall lack of structuring according to ecological factors in badgers suggests that their high mobility counteracts any isolating impacts of habitat heterogeneity, particularly in western North America. Much of our dataset was contained within the West cluster, a single genetic population characterized by IBD. Mobile specialists can experience extensive connectivity in suitable habitat, particularly in the absence of topographic barriers (e.g., Carmichael et al. 2007; Marthinsen et al. 2009; Row et al. 2012). However, the large West cluster in badgers was surprising because it encompassed many potentially isolating barriers, including those that defined population boundaries in previous studies (e.g., mountain ranges; Kyle et al. 2004; Ethier et al. 2012). We likely found limited genetic subdivision in western North America because detection of genetic differentiation heavily depends on the balance between genetic drift and gene flow, two evolutionary processes that promote and resist genetic differentiation respectively. The rate of genetic drift in a population is directly correlated with effective
population size (Wright 1931), and previous studies included endangered Canadian populations whose total census sizes are 200 or less (COSEWIC 2012). Even in these small populations, mountain ranges were not complete barriers to gene flow (Kyle et al. 2004; Ethier et al. 2012), so the extensive gene flow within western North America appears to overcome any isolating effects of the Rocky Mountains or ecological variables. Therefore, badgers appear to maintain large population sizes outside isolated, peripheral areas, which combined with their high mobility resists genetic differentiation.

Badgers in eastern North America formed two genetic clusters and had lower genetic diversity than western areas, which at least in part result from large water barriers (i.e., Mississippi River and Lake Michigan) preventing gene flow. Both the Mississippi River and Lake Michigan appear to disrupt enough gene flow to allow genetic drift to cause detectable genetic differentiation. Considerable admixture and disagreement between Bayesian clustering programs was observed between the West and Wisconsin clusters, so greater gene flow likely occurs across the Mississippi River than Lake Michigan. Another factor that may enhance genetic differentiation observed in eastern North America is that eastern badgers occur at a range periphery. Genetic drift is stronger in populations with small effective populations sizes, and effective population sizes in peripheral populations can be much lower than core areas (Vucetich & Waite 2003). Eastern North America certainly contains less grassland habitat than western North America, which may cause lower population sizes and high fluctuations typical of peripheral populations (Hengeveld & Haeck 1982). Based on the high habitat heterogeneity at the eastern periphery, genetic differentiation due to ecological variables should have been detectable if gene flow was restricted as observed in other highly
mobile species (e.g., Schwartz et al. 2009; Garroway et al. 2011; Weckworth et al. 2013; Balkenhol et al. 2014). However, we only detected genetic differentiation according to topographic barriers, so badgers appear able to maintain high gene flow except when faced with large geographic barriers.

Lake Michigan prevents gene flow from western populations (i.e., Upper Peninsula and Wisconsin) as evidenced by the strong assignments of most LP individuals to the LP cluster. The LP, therefore, mainly receives migrants from southern areas that have low density badger populations (e.g., Warner & Ver Steeg 1995; Duquette & Gehrt 2014). A lack of connectivity with the genetically diverse populations to the west facilitated the loss of genetic diversity in LP badgers following postglacial recolonization. Other peninsular populations with low genetic diversity have been recorded in highly mobile species, and most have found genetic variation patterns consistent with a combination of historic founder events and subsequent isolation within a peninsula (Tammeleht et al. 2010; Reding et al. 2012; Frantz et al. 2014). Our data supports a historic founder event and subsequent isolation because no demographic declines have been recorded in the LP and both genetic datasets exhibited low diversity. Very little life history information is available for badgers in the LP, but previous genetic work in the LP suggested three potential mechanisms for heterozygote deficiencies: Wahlund Effect, mixing of populations, and inbreeding (Kierepka et al. 2012). We also recorded a heterozygote deficiency within the LP, but with the strong assignments to a single cluster in both Bayesian programs, badgers in the LP appear to experience some degree of inbreeding. Further assessment of the LP should focus on determining if badgers in the LP warrant conservation actions like other peripheral populations in British Columbia.
and Ontario. The potential for inbreeding as a mechanism reducing genetic variation in LP badgers, and other peripheral populations, should be further explored by pairing it with demographic and fitness related data to assess any future conservation needs.

Conclusions

Our broad-scale genetic investigation of gene flow in badgers revealed limited genetic structure across much of North America. IBD largely defined genetic structure within microsatellites, but we still detected signatures of historical biogeographic events. Badgers in western North America appear to maintain a robust population size based on the high genetic diversity and limited differentiation west of the Mississippi River. In contrast, eastern North America had genetic differentiation due to the Mississippi River and Lake Michigan as well as lower genetic diversity, so we believe eastern badgers have lower or more variable population sizes than western grassland-dominated areas. In particular, the LP was genetically distinct in all analyses due to both low genetic diversity and barrier effects of the Lake Michigan, which suggests a combination of recolonization following glacial retreat and peninsular geography driving genetic variation. Taken together, high mobility appears to prevent genetic differentiation according ecological factors in badgers, but differences in genetic diversity and differentiation between western and eastern North America may reflect variation in habitat quality impacting the strength of genetic drift. Therefore, even if high mobility in badgers is the dominant driver of gene flow across North America, their specialist traits may influence genetic variation through differences in effective population sizes according to habitat quality.
With limited genetic structure in badgers observed throughout much of North America, patterns of mitochondrial and microsatellite variation conflict with all current morphological subspecific designations. Previous studies have found evidence for a split between *T. t. taxus* and *jeffersonii* in peripheral, endangered populations (Kyle *et al.* 2004; Ethier *et al.* 2012), but with the inclusion of more southern locations in this study, no such divergence was found. Discordance between morphological subspecies and genetic variation in North America is relatively common (e.g., Zink 2004; Cullingham *et al.* 2008; Godbout *et al.* 2008; Hull *et al.* 2008; Sabatino & Routman 2009; Kodandaramaiah *et al.* 2012; Latch *et al.* 2014), which may argue for taxonomic revision of badgers. However, proper delineation of subspecies should consider multiple datasets including genetics, morphology, and behavior (Haig *et al.* 2006), and our dataset based on neutral genetic markers does not provide a complete picture of how subspecific boundaries may reflect evolutionary divergence. In particular, neutral datasets may not reflect variation at adaptive loci, and morphological differences could reflect selection pressures that coincide with subspecific boundaries. Therefore, any decision to re-evaluate taxonomic classification should incorporate potential adaptive drivers of morphological differentiation in badgers, especially since no common thread appears to exist in how carnivores react to landscape heterogeneity.

Badgers join the growing number of highly mobile species that demonstrate the difficulty in predicting how dispersal capability and habitat preferences impact gene flow in widespread taxa. Badgers represent somewhat of a unique case among specialists because we observed limited genetic differentiation across multiple habitat types, a finding that usually occurs only within suitable habitat (Carmichael *et al.* 2007; McRae &
Beier 2007; Koen et al. 2012; Row et al. 2012). Therefore, reliance on grasslands and fossorial rodents does not preclude badgers from modified or sub-optimal habitats, but maintaining corridors (e.g., remnant habitats in high agricultural landscapes; Duquette & Gehrt 2014) may be critical for continued high gene flow within peripheral populations. In conclusion, this broad-scale genetic survey of badgers revealed that dispersal capabilities and specialization influence both gene flow and effective population size, and thus, can produce disparate genetic patterns across a large geographic range.
Figure 3.1. Sampling locations and subspecific designations of $n = 917$ badgers. Designations are morphological subspecies as defined in Long (1972).
Figure 3.2. Haplotype network for 95 d-loop haplotypes detected in $n = 785$ badgers. Each circle represents a single haplotype and the size of the circle corresponds to its frequency. Each line represents a single mutational step, and black circles indicate missing genotypes. The network was color-coded by the morphological subspecies (a) or inferred subspecies (b). The most common haplotypes (H1, H2, H3, and H4) are labeled.
Figure 3.3. Mismatch distributions for the total dataset (a), subspecies (b-e), and SAMOVA groups (f-g). Observed distributions are represented as gray bars and expected distributions under rapid demographic expansion are black solid lines. 95% confidence intervals for the expected distribution are given as dotted black lines. All groups followed expected patterns under rapid demographic expansion (all p < 0.008) except for the eastern SAMOVA group (p < 0.002).
Figure 3.4. Distribution of three genetic clusters inferred in BAPS 5. The three clusters (West: black; Wisconsin: grey, and LP: white) are largely separated by two major water barriers: the Lake Michigan and the Mississippi River. The Mississippi River is represented by the bold line.
Figure 3.5. Plot of first two PCA axes when grouped by large aquatic barriers (a) or subspecies (b). Each point represents a single badger's genotype that is color-coded according to either the aquatic barriers (east of Lake Michigan, west of Lake Michigan/East of Mississippi River, and west of the Mississippi River) or subspecies (*T. t. berlandieri, jacksoni, jeffersonii, or taxus*). Ovals around points correspond to 95% confidence ellipses that denote where 95% of individuals within each group occur. Each square on the background grid corresponds to 0.5.
Figure 3.6. Results of the sPCA analysis. Eigenvalue plots (a) and the screeplot (b) indicate that the first two axes are most important in explaining genetic variation. The screeplot plots the relationship between variance explained by each axis and spatial autocorrelation (Moran’s I) within each axis. Spatially lagged scores from Axes 1 (c) and 2 (d) depict two distinct patterns in genetic variation. Squares represent each individual where larger squares have stronger positive (white) or negative (black) values whereas smaller, grey (light grey = positive, dark grey = negative) squares are less strongly differentiated.
**Figure S3.1.** Geographic distribution of the four main haplotypes and those with the 26 bp deletion region that were not included within the mtDNA analyses. Rare haplotypes had a frequency of 10 or less individuals (dark grey circles). Haplotype 1 (red squares) is found throughout North America, but its frequency is the highest in the Upper Midwest. Haplotype 2 (blue circles) primarily occurs within northern latitudes, particularly in Montana, Canada, and North Dakota. Haplotype 3 (yellow diamonds) is one base pair away from Haplotype 1, and is chiefly located in the Lower Peninsula of Michigan and Ohio. Haplotype 4 (light blue triangles) is one mutation away from Haplotype 2, and almost always occurred west of the Rocky Mountains. Like Haplotype 4, individuals with the 26 bp deletion region (green squares) are primarily found west of the Rocky Mountains. The haplotype network of the 95 haplotypes that did not include the deletion region is provided where the four main haplotypes are colored.
Figure S3.2. Results of the overall clustering analysis within STRUCTURE. ΔK indicated the most likely number of clusters was 2 as shown by the large peak at $K = 2$ within the plot (a). The spatial arrangement of q-values (b) indicated a general cline from east to west. We interpolated q-values using inverse-weight distance to create a genetic surface across our sampling area and individuals are colored according to their q-values. Those with higher q-values (large blue circles) were more strongly assigned to the eastern cluster.
**Figure S3.3.** Results of the iterative runs of the two main clusters detected in the overall clustering analysis within **STRUCTURE**. The $\Delta K$ plot for the eastern cluster (a) indicated further substructure whereas the plot for the western cluster (b) did not strongly support any additional genetic substructure. When mapped (c), the iterative run in the eastern cluster ($n = 210$ individuals) suggested the presence of two additional clusters that separated the Lower Peninsula of Michigan from the rest of the individuals. Interpolated $q$-values, created via inverse-weight distance interpolation, revealed a strong divergence between the Lower Peninsula and Wisconsin (high assignment to the Lower Peninsula is denoted with large blue circles).
Table 3.1. Results of SAMOVA for $K = 2$ where the first group contained the Lower Peninsula of Michigan, Indiana, and Ohio and the second group contained all other individuals. All variance components were significant after 1000 permutations (all $p < 0.001$).

<table>
<thead>
<tr>
<th>Source</th>
<th>Df</th>
<th>SS</th>
<th>Components</th>
<th>% Var</th>
</tr>
</thead>
<tbody>
<tr>
<td>Among Groups</td>
<td>1</td>
<td>138.955</td>
<td>0.564</td>
<td>25.08</td>
</tr>
<tr>
<td>Among Pops Within Groups</td>
<td>17</td>
<td>129.667</td>
<td>0.161</td>
<td>7.18</td>
</tr>
<tr>
<td>Within Pops</td>
<td>765</td>
<td>1164.412</td>
<td>1.522</td>
<td>67.74</td>
</tr>
<tr>
<td>Total</td>
<td>783</td>
<td>1433.04</td>
<td>2.247</td>
<td>100.00</td>
</tr>
</tbody>
</table>
Table 3.2. Diversity metrics for n = 785 badgers, for each of the four subspecies, and for the two SAMOVA groups. All diversity measures include standard errors.

<table>
<thead>
<tr>
<th>Grouping</th>
<th>N</th>
<th>H       ± SE</th>
<th>π         ± SE</th>
<th>Pairwise diff ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total</strong></td>
<td>785</td>
<td>0.831 ± 0.099</td>
<td>0.006 ± 0.003</td>
<td>3.660 ± 1.855</td>
</tr>
<tr>
<td><strong>Subspecies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T. t. berlandieri</td>
<td>24</td>
<td>0.873 ± 0.047</td>
<td>0.008 ± 0.005</td>
<td>4.681 ± 2.376</td>
</tr>
<tr>
<td>T. t. jacksoni</td>
<td>222</td>
<td>0.325 ± 0.034</td>
<td>0.003 ± 0.002</td>
<td>1.856 ± 1.068</td>
</tr>
<tr>
<td>T. t. jeffersonii</td>
<td>54</td>
<td>0.926 ± 0.040</td>
<td>0.008 ± 0.005</td>
<td>4.774 ± 2.370</td>
</tr>
<tr>
<td>T. t. taxus</td>
<td>485</td>
<td>0.786 ± 0.045</td>
<td>0.007 ± 0.004</td>
<td>3.827 ± 1.928</td>
</tr>
<tr>
<td><strong>SAMOVA grouping</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>East SAMOVA</td>
<td>134</td>
<td>0.523 ± 0.023</td>
<td>0.001 ± 0.001</td>
<td>0.800 ± 0.582</td>
</tr>
<tr>
<td>West SAMOVA</td>
<td>651</td>
<td>0.813 ± 0.011</td>
<td>0.007 ± 0.004</td>
<td>3.319 ± 1.924</td>
</tr>
</tbody>
</table>

N: Sample sizes for each group; h: haplotype diversity; π: nucleotide diversity
Table 3.3. Mismatch statistics for the four subspecies and for the two SAMOVA groups. Fu’s Fs and Tajima’s D indicate demographic expansion when they are significantly negative whereas significant SSD and raggedness indices suggest deviations from patterns expected under demographic expansion. P-values for each test are given in parentheses; significant values (P < 0.05) are in bold.

<table>
<thead>
<tr>
<th>Group</th>
<th>N</th>
<th>Fu’s Fs</th>
<th>Tajima’s D</th>
<th>SSD</th>
<th>Raggedness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>785</td>
<td>-25.075</td>
<td>-1.352</td>
<td>0.036</td>
<td>0.025</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.015)</td>
<td>(0.422)</td>
<td>(0.112)</td>
</tr>
<tr>
<td>T. t. berlandieri</td>
<td>24</td>
<td>-25.464</td>
<td>-0.516</td>
<td>0.106</td>
<td>0.033</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.328)</td>
<td>(0.125)</td>
<td>(0.423)</td>
</tr>
<tr>
<td>T. t. jacksoni</td>
<td>222</td>
<td>-27.350</td>
<td>-0.895</td>
<td>0.096</td>
<td>0.031</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.125)</td>
<td>(0.098)</td>
<td>(0.282)</td>
</tr>
<tr>
<td>T. t. jeffersonii</td>
<td>54</td>
<td>-25.618</td>
<td>-0.729</td>
<td>0.020</td>
<td>0.014</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.221)</td>
<td>(0.324)</td>
<td>(0.428)</td>
</tr>
<tr>
<td>T. t. taxus</td>
<td>485</td>
<td>-25.220</td>
<td>-1.005</td>
<td>0.035</td>
<td>0.022</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.120)</td>
<td>(0.183)</td>
<td>(0.212)</td>
</tr>
<tr>
<td>East SAMOVA</td>
<td>134</td>
<td>0.048</td>
<td>-1.384</td>
<td>0.029</td>
<td>0.211</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.563)</td>
<td>(0.065)</td>
<td>(0.001)</td>
<td>(0.002)</td>
</tr>
<tr>
<td>West SAMOVA</td>
<td>651</td>
<td>-25.096</td>
<td>-1.346</td>
<td>0.023</td>
<td>0.038</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.001)</td>
<td>(0.059)</td>
<td>(0.288)</td>
<td>(0.346)</td>
</tr>
</tbody>
</table>
Table 3.4. Genetic diversity measures for three genetic clusters detected in BAPS. All three populations had significant heterozygote deficiencies (bold), and LP had lower genetic diversity than WI and West (all t > 2.341, p < 0.02). P-values are given in parentheses for F_{IS} values.

<table>
<thead>
<tr>
<th></th>
<th>N</th>
<th>A_R</th>
<th>H_O</th>
<th>H_E</th>
<th>F_{IS}</th>
</tr>
</thead>
<tbody>
<tr>
<td>LP</td>
<td>115</td>
<td>7.99</td>
<td>0.641</td>
<td>0.705</td>
<td><strong>0.089 (0.001)</strong></td>
</tr>
<tr>
<td>WI</td>
<td>155</td>
<td>10.22</td>
<td>0.731</td>
<td>0.804</td>
<td><strong>0.092 (0.001)</strong></td>
</tr>
<tr>
<td>West</td>
<td>649</td>
<td>11.16</td>
<td>0.747</td>
<td>0.816</td>
<td><strong>0.086 (0.001)</strong></td>
</tr>
</tbody>
</table>

A_R: allelic richness corrected for smallest sample size of 115; H_O: observed heterozygosity; H_E: expected heterozygosity; F_{IS}: Weir & Cockerham (1984)'s inbreeding coefficient
**Table 3.5.** Results from partial dbRDA analyses to evaluate the relationship between genetic distance and topographic barriers, geographic coordinates, and landscape variables. The final model after sequential selection included Latitude, Longitude, Lake Michigan, and the Mississippi River (bold), and together explained 10.11% of the total genetic variation in the dataset. Variance explained by each singular variable with all other variables partialled out are provided (% var).

<table>
<thead>
<tr>
<th>Variable</th>
<th>F-ratio</th>
<th>p-value</th>
<th>% var</th>
</tr>
</thead>
<tbody>
<tr>
<td>Latitude</td>
<td>16.31</td>
<td>0.0001*</td>
<td>1.41</td>
</tr>
<tr>
<td>Longitude</td>
<td>10.54</td>
<td>0.0001*</td>
<td>0.91</td>
</tr>
<tr>
<td>Lake Michigan</td>
<td>21.45</td>
<td>0.0001*</td>
<td>1.86</td>
</tr>
<tr>
<td>Mississippi River</td>
<td>7.92</td>
<td>0.0002*</td>
<td>0.69</td>
</tr>
<tr>
<td>Rocky Mountains</td>
<td>3.95</td>
<td>0.0260*</td>
<td>0.34</td>
</tr>
<tr>
<td>Subspecies</td>
<td>1.86</td>
<td>0.1500</td>
<td>0.16</td>
</tr>
<tr>
<td>Soil Order</td>
<td>5.09</td>
<td>0.0240*</td>
<td>0.44</td>
</tr>
<tr>
<td>Soil Texture</td>
<td>0.37</td>
<td>0.9020</td>
<td>0.09</td>
</tr>
<tr>
<td>Precipitation</td>
<td>4.12</td>
<td>0.0300*</td>
<td>0.41</td>
</tr>
<tr>
<td>Land Cover</td>
<td>3.25</td>
<td>0.0510</td>
<td>0.22</td>
</tr>
</tbody>
</table>

*corrected alpha = 0.015
Table S3.1. Information about all the badgers, location, and source. Location corresponds to the state where badgers were sampled and for those that had small sample sizes (< 10 individuals), their SAMOVA group is listed in parentheses. All other SAMOVA groups are identical to their capture state or Canada. Sample sizes (N) correspond to the number of individuals provided by each collector and organization.

<table>
<thead>
<tr>
<th>Location</th>
<th>N</th>
<th>Organization</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alberta (Canada)</td>
<td>5</td>
<td>North American Fur Auction</td>
</tr>
<tr>
<td>Arizona (New Mexico)</td>
<td>2</td>
<td>Arizona Fish and Game Department</td>
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CURRICULUM VITAE

Education

B.S., Michigan State University-December 2006
Major: Zoology, Honors

M.S., Central Michigan University-May 2009
Major: Biology, Conservation Biology

Dissertation Title: Landscape genetics of the American badger: understanding challenges with elusive species