

Inferring landscape effects on gene flow: a new model selection framework

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Abstract

Populations in fragmented landscapes experience reduced gene flow, lose genetic diversity over time and ultimately face greater extinction risk. Improving connectivity in fragmented landscapes is now a major focus of conservation biology. Designing effective wildlife corridors for this purpose, however, requires an accurate understanding of how landscapes shape gene flow. The preponderance of landscape resistance models generated to date, however, is subjectively parameterized based on expert opinion or proxy measures of gene flow. While the relatively few studies that use genetic data are more rigorous, frameworks they employ frequently yield models only weakly related to the observed patterns of genetic isolation. Here, we describe a new framework that uses expert opinion as a starting point. By systematically varying each model parameter, we sought to either validate the assumptions of expert opinion, or identify a peak of support for a new model more highly related to genetic isolation. This approach also accounts for interactions between variables, allows for nonlinear responses and excludes variables that reduce model performance. We demonstrate its utility on a population of mountain goats inhabiting a fragmented landscape in the Cascade Range, Washington.

Keywords: circuit theory, gene flow, isolation by distance, landscape resistance, mountain goat

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Introduction

Habitat fragmentation can limit the spread of genetic variation across a landscape (gene flow) and reduce the local effective population size (Keyghobadi 2007). Small populations isolated from gene flow lose genetic diversity over time and are therefore susceptible to extinction by genetic processes, including inbreeding depression (Crnokrak & Roff 1999), random fixation of deleterious alleles (Lynch *et al.* 1995; Lande 1998) and loss of adaptive potential (Lande 1995; Swindell & Bouzat 2005). Improving connectivity in fragmented landscapes, and therefore the probability of population persistence, is now a major conservation priority (Crooks & Sanjayan 2006). Designing effective reserve systems and wildlife corridors for this purpose, however, requires a spatially

explicit understanding of how landscapes influence gene flow.

Despite the importance of genetic connectivity to the viability of fragmented populations, few studies employ appropriate empirical data to derive landscape resistance models. Beier *et al.* (2008) recently cited 24 studies that produced maps of corridors, linkages or cost surfaces to guide connectivity conservation decisions. Among these, 15 were based on expert opinion and literature review alone. The lack of empirical data to inform these models makes them highly subjective, given that gene flow is difficult to intuit quantitatively, regardless of the modeller's knowledge of the species (Wang *et al.* 2008). While expert knowledge is important for identifying landscape variables that may shape gene flow and formulating initial hypotheses, unvalidated models parameterized entirely by experts have the potential to misinform connectivity assessments.

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Seven of the 24 studies cited in Beier *et al.* (2008) assigned landscape resistance to gene flow based on habitat models or telemetry data. If a species is able to disperse through poor habitat, however, habitat associations are not necessarily strong indicators of gene flow. Telemetry data may also be misleading because exploratory movements cannot be distinguished from dispersal events that result in survival and reproduction (Koenig *et al.* 1996). As a result, relying on these proxy measures to parameterize landscape resistance models may also lead to misinformed conclusions and management decisions (Horskins *et al.* 2006).

A more rigorous means to infer landscape effects on gene flow is to use genetic data to quantify the degree of genetic isolation between populations or individuals within the study area (Storfer *et al.* 2007; Holderegger & Wagner 2008). Landscape resistance to gene flow between these populations or individuals can then be inferred by identifying a spatial model (also known as a 'resistance surface') among multiple alternative models that best accounts for the observed genetic isolation (reviewed in Spear *et al.* 2010). Indeed, one of the 24 studies cited in Beier *et al.* (2008) used genetic data to relate environmental gradients of slope to gene flow among bighorn sheep (*Ovis canadensis*) in southern California (Epps *et al.* 2007). Innovative landscape genetics approaches such as these and others (e.g. Cushman *et al.* 2006; Garroway *et al.* 2008; Perez-Espona *et al.* 2008; Murphy *et al.* 2009, 2010; Wang *et al.* 2009) are powerful tools that can inform conservation efforts.

Landscape genetic models published to date, however, are often only weakly related to the genetic isolation observed within the study area. One explanation may be the limited capacity of previous model selection frameworks to sample the multidimensional hypothesis space that can be derived from alternative hypotheses of several landscape variables thought to govern gene flow. As it is only practical to test a small subset of this limitless space, researchers have made compromises such as arbitrarily constraining the range of possible resistance values, not accounting for potential interactions between variables, and testing only linear relationships between environmental variables and landscape resistance. These simplifications, as well as the inability of some frameworks to exclude variables that do not improve the fit of the model, reduce the likelihood of identifying a model strongly related to the observed genetic isolation.

Here, we describe a new model selection framework that treats the range of resistance values for each variable as a quasi-unconstrained parameter, allows for interactions among variables, includes nonlinear responses, is capable of excluding variables that reduce model fit and bridges the gap between empirical models and those based on expert opinion alone. Addi-

tionally, this framework incorporates a causal modelling design that evaluates the support for genetic isolation by landscape resistance (IBR; Cushman *et al.* 2006; McRae 2006) relative to null models of isolation by distance (Wright 1943) or isolation by barrier (Ricketts 2001). We demonstrate the utility of this framework by inferring the effect of the Cascade Range, Washington landscape on the genetic isolation of mountain goats (*Oreamnos americanus*).

Methods

Study area

The study area included approximately 36 500 km² of the Cascade Range of Washington, extending along a north-south axis 315 km from Mount Adams to the Canadian border (Fig. 1). The landscape is mostly mountainous and covered with montane forests, except at high elevations, where subalpine parkland, rocky alpine summits and glaciers predominate (Beckey 1987). Elevation varies from near sea level to almost 4400 m. Interstate 90 (I90) crosses the study area on an east-west axis approximately in the centre of the range. In addition, three state highways run east-west across the range (two north and one south of I90), and numerous other highways and secondary roads intrude towards the crest from lower elevation valleys.

Sample collection

We obtained 149 genetic samples from the study area (Fig. 1) by multiple means between 2003 and 2008. The National Park Service (NPS) contributed nine blood samples (six from Mount Rainier National Park and three from the North Cascades National Park) from chemically immobilized animals. Washington Department of Fish and Wildlife (WDFW) also contributed 41 blood samples from chemically immobilized animals. WDFW contributed an additional 68 tissue samples donated by legally permitted hunters. We also collected 29 tissue samples by biopsy darting via helicopter or on foot. Lastly, we obtained one bone marrow and one muscle sample from carcasses found in the field (cause of mortality unknown). The 149 genetic samples collected from the Cascade Range represents 6.2% of the approximately 2400 mountain goats thought to occupy the study area (C. Rice, WDFW, unpublished). All sample collection sites were unique except three sites with two individuals and one site with three individuals (Fig. 1). In collaboration with the NPS and the United States Geological Survey (USGS), we collected an additional 12 tissue and blood samples from the non-native Olympic Range population, from which approximately

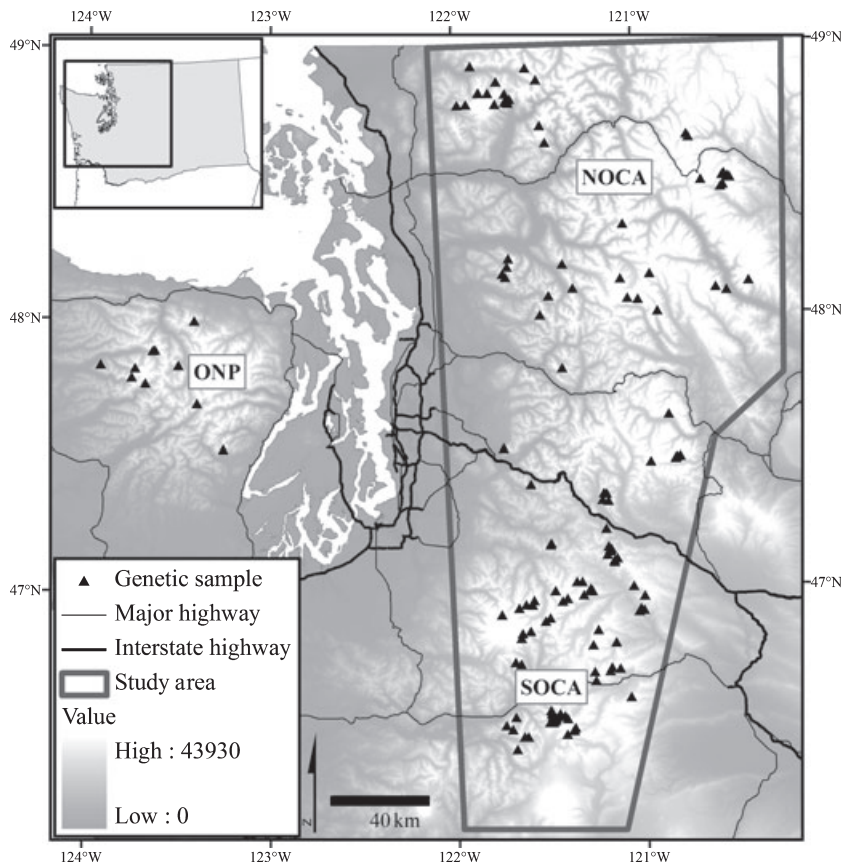


Fig. 1 Study area. The locations of genetic samples (black triangles), major highways (thin black lines), interstate highways (thick black lines), elevation and the landscape resistance model extent (dark grey box) are depicted. The north Cascade (NOCA), south Cascade (SOCA) and Olympic peninsula (ONP) regions are also noted.

130 animals were translocated to the Cascade Range in the 1980s. All procedures were approved by the Animal Care and Use Committee at Western Washington University and permitted by WDFW and the NPS.

Genotyping

We performed all laboratory procedures at the WDFW molecular genetics laboratory in Olympia, Washington. We extracted DNA from tissue, bone marrow or blood with the commercial DNAeasy[®] blood and tissue DNA isolation kit (Qiagen) according to the manufacturer's protocol. We eluted DNA from the column with 100 µL of elution buffer and used polymerase chain reaction (PCR) to amplify 19 previously characterized polymorphic microsatellite markers (Mainguy *et al.* 2005) in six separate uni- or multiplex reactions (multiplex Oam-A [BM121, BM4107, BM6444, TGLA122], Oam-B [OarCP26, OarHH35, RT27], Oam-C [BM1818, BM4630, RT9, URB038], Oam-D [BM203, BMC1009, HUI616, McM527], Oam-E [BM1225, BM4513, HEL10] and TGLA10). PCR conditions are provided in Table S1. We visualized PCR products with an ABI 3730 capillary sequencer (Applied Biosystems) and sized them using the GeneScan 500-Liz size standard (Applied Biosystems) and Genemapper 3.7 (Applied Biosystems).

We used the program MICROCHECKER 2.2.0 (Van Oosterhout *et al.* 2004) to screen for genotyping errors, including allelic dropout, null alleles and stuttering that might hinder detection of heterozygotes. No evidence for allelic dropout or stuttering was detected by MICROCHECKER; however, seven loci (BM203, BMC1009, BM1818, HEL10, BM1225, TGLA10 and HUI616) showed significant homozygote excess, indicating the potential existence of null alleles. Homozygote excess can also arise because of inbreeding, however. After accounting for biased null allele estimates owing to inbreeding ($F = 0.10, 0.12$ and 0.19 in the north Cascade, south Cascade and Olympic peninsula subpopulations identified in the Structure analysis later, respectively) with Null Allele Estimator 1.3 (Van Oosterhout *et al.* 2006), the estimated frequency of null alleles was <0.03 for all seven loci. We therefore elected to retain these loci for further analysis.

Detecting ONP-Cascade admixture because of past translocations

A non-native population of mountain goats was introduced to the Olympic Range from Alaska and British Columbia in the 1920s. In the 1980s, over 100 goats were translocated from the Olympic Range to the

Cascade Range to re-establish occupancy in areas depleted mainly by harvest (Houston *et al.* 1991). Although post-release mortality may have been high, significant admixture between survivors and the native Cascade population owing to translocation had the potential to influence our analysis, as the genotypes of admixed individuals do not solely reflect the Cascade landscape effects on gene flow. To quantify admixture, we used Structure 2.2 (Pritchard *et al.* 2000), which performs Bayesian inference of the most likely number of populations sampled and assigns individuals to populations based on minimizing Hardy–Weinberg and linkage disequilibrium within populations. To infer the number of populations, the program uses a Markov chain Monte Carlo (MCMC) simulation to estimate the posterior probability that the data fit the hypothesis of K populations, $P(X|K)$. We tested values of K ranging from 1 to 10 with 10 independent runs per test (admixture model with correlated allele frequencies, a 100 000 step burn-in followed by 10^6 steps of data collection).

Once we delineated the population structure, we used the fractional probability (Q) of individuals belonging to each subpopulation to identify Cascade individuals strongly admixed with the ONP subpopulation. While all Cascade genotypes might have some low background posterior probability of ONP membership, those with low probability of ONP ancestry are unlikely to bias results, and their omission reduces sample size for analyses of the Cascade population. As a balance between bias and sample size, we required the posterior probability of ONP population membership to be >0.25 to classify a Cascade genotype as 'ONP-Cascade admixed.' We excluded individuals meeting this criterion from further analysis.

Modelling framework overview

We hypothesized that genetic differentiation within the study area was a function of isolation by landscape resistance or isolation-by-resistance (IBR). We a priori considered four landscape variables to be the most important resistors of mountain goat gene flow: distance to escape terrain (D_{et}), roads, landcover type and elevation (a proxy for climate conditions to which mountain goats are adapted). We related each of these variables to landscape resistance with a simple mathematical function characterized by several parameters (see Model Functions) and used each function to reclassify appropriate raster data into a resistance surface. Initially, we parameterized these functions based on expert opinion obtained from the consensus of four biologists with extensive knowledge of mountain goat biology and habitat relations. We then evaluated the relationship of the expert opinion parameterized

resistance surface for each variable to the genetic differentiation observed within the study area (see Model Evaluation). By systematically varying the expert opinion parameter values, we sought to find the univariate optimal hypothesis (see Univariate Optimization) for each variable. We then combined resistance surfaces corresponding to the univariate optimal parameters for each variable into a multivariate model. We again evaluated the optimal parameter values, but in the multivariate context (see Multivariate Optimization). After identifying a multivariate IBR model most related to genetic isolation within the study area, we tested the support for this model in comparison with the null models of isolation by distance (IBD) and IBB within a causal modelling framework (see Causal Modeling).

Model functions

We modelled landscape resistance arising from distance to escape terrain (D_{et}) by reclassifying a raster representing Euclidean distance to escape terrain (defined as slope $\geq 50^\circ$ based on a habitat model derived for coastal mountain goats; Smith 1994) according to the following function:

$$R = (D_{et}/V_{max})^x * R_{max}$$

where R is the resistance for that pixel, x is the response shape exponent, R_{max} is the maximum resistance value (i.e. resistance ranges from 1 to R_{max}) and V_{max} is a constant defining the maximum allowed value of the variable. Thus, as the variable increases up to V_{max} , resistance increases towards R_{max} . If $x = 1$, the increase is linear, and if $x < 1$ or $x > 1$, the increase in resistance is nonlinear. For D_{et} , we set $V_{max} = 600$ m based on the observation that 95% of all telemetry observations collected by WDFW from mountain goats within the study area were within this distance of escape terrain (C. Rice, WDFW, unpublished). Any pixel >600 m from escape terrain was assigned a resistance equal to R_{max} .

We modelled landscape resistance arising from roads by reclassifying a raster with four levels (0, 50, 4000 and 28 000 vehicles per day) of annual average daily traffic volume (AADT) corresponding to typical use of no road, secondary/forest service road, highway and interstate highway within the study area (WSDOT 2006). We related these four levels of road traffic volume to landscape resistance according to the function:

$$R = (AADT/V_{max})^x * R_{max}$$

where x and R_{max} represent the response shape exponent and maximum resistance, respectively, and V_{max}

is a constant (28 000) representing the vehicle use per day for the highest traffic volume road category. Thus, as vehicle traffic on a road increases towards V_{max} , resistance increases towards R_{max} at a rate governed by x .

We obtained landcover data from the Northwest GAP Analysis Project (NWGAP 2007) and reclassified it into six broad landcover types. We then ranked these landcover types in order of increasing landscape resistance to mountain goat gene flow based on input from an expert panel of biologists. We assigned alpine/rock the lowest rank, followed by subalpine forest/parkland, ice, montane forest, mesic forest/disturbed and water. We then generated a resistance surface for landcover according to the function:

$$R = (Rank/V_{max})^x * R_{max}$$

where x and R_{max} represent the response shape exponent and maximum resistance, respectively, and V_{max} is a constant (6) representing the highest landcover resistance rank. Thus, as the landcover rank increases towards V_{max} , resistance increases towards R_{max} at a rate governed by x .

In addition to the earlier landcover types, some urban and agricultural areas were present within the study area. We reasoned that, because these categories are only present in small patches, and mountain goats strongly avoid such highly modified human habitats, gene flow would be directed around rather than through these areas. Consequently, we assigned these areas a prohibitively high resistance fixed at 100 000.

We modelled landscape resistance arising from elevation by reclassifying a digital elevation model of the study area according to a Gaussian function, which is appropriate for mountain goats as they are adapted to an optimal elevation range straddled by suboptimal elevations where lowlands and glaciated summits predominate (Festa-Bianchet & Côté 2008). This function takes the form:

$$R = R_{max} + 1 - R_{max} * e^{\frac{-(elevation - E_{opt})^2}{2 * E_{SD}^2}}$$

where R_{max} , E_{opt} and E_{SD} represent the maximum resistance, optimal elevation and the standard deviation about the optimal elevation, respectively. Thus, as elevation increases or decreases away from E_{opt} , resistance increases to R_{max} at a rate governed by E_{SD} (Fig. S1). We reasoned that mountain goat gene flow would never traverse the summits of Cascade strato-volcanoes and therefore assigned this elevation class (>3300 m) a prohibitively high resistance fixed at 100 000.

Model evaluation

We constructed a principal components analysis (PCA)-based genetic data matrix G with n rows $\times$ m columns, where n is the number of individuals in the analysis and m is the number of alleles present within the population ($m = 97$). Each element in the matrix $G(i,j)$ is populated for individual i by the number of occurrences for the j th allele. We then computed the eigenvectors of G in R 2.8 (R Development Core Team 2008). Finally, we used the Ecodist package in R 2.8 (Goslee & Urban 2007) to generate an $n \times n$ pairwise genetic distance matrix (Y) based on the distance between individuals along the first eigenvector (Patterson *et al.* 2006). As a comparison to PCA-based genetic distance, we also calculated genetic distance based on the proportion of shared alleles (D_{PS} ; Bowcock *et al.* 1994) and Rousset's a (Rousset 2000) in FSTAT 2.93 (Goudet 1995).

Next, we used Circuitscape 3.0.1 (McRae 2006) to generate a matrix (X) of pairwise resistance distances between all genetic sampling locations for each landscape resistance surface. Circuitscape uses circuit and graph theory to quantify current and total resistance between pairs of points (McRae & Beier 2007). We used a connection scheme where gene flow was only allowed between the nearest four cells (i.e. no diagonal connections because they allow connectivity across linear features like highways without additional resistance when the highway pixels are not aligned north-south or east-west), and resistance between any two cells was based on the average of the resistance assigned to both cells. To achieve a reasonable computing time for each Circuitscape run ($\bar{x} \approx 2$ hr), we decreased the resolution of each resistance surface from 30 to 450 m by aggregating 15×15 blocks of 30 m pixels into a single pixel based on the average resistance of all the cells within the block. We also created landscape distance matrices representing the null models of IBD and IBB. We based the barrier model on the hypothesis that Interstate 90 subdivides the Cascade population into two panmictic subpopulations (NOCA and SOCA) with no migration. Specifically, we assigned a pairwise landscape distance of zero for samples within a subpopulation and a pairwise landscape distance of 100 000 (approximating a complete barrier to migration) for samples between subpopulations.

To evaluate each model, we used Mantel's tests (Mantel 1967) in the R software package Ecodist (Goslee & Urban 2007) to calculate the correlation between genetic and landscape distance. We identified the most supported model as the one with the highest significant correlation if it also met the additional criteria described in the Univariate and Multivariate Optimization sections as follows.

Univariate optimization

First, we determined the correlation between the univariate expert opinion models and genetic distance (i.e. X_{Det} , X_{Road} , X_{Land} or $X_{Elev} \sim Y$). By systematically increasing or decreasing the function parameters (x or R_{max} for landcover, roads and distance to escape terrain, or R_{max} , E_{opt} and E_{SD} for elevation) and re-evaluating the correlation, we determined whether altering the expert opinion model parameters improved the correlation with genetic isolation for each variable. We generated alternative parameter values (favouring the direction that increases the correlation) until we observed a unimodal peak of support (i.e. the correlation decreased steadily if the parameter was less than or greater than the optimal value) for all parameters or they reached a natural limit (i.e. the optimal parameter value was zero or infinity).

Given the number of hypotheses tested (25–45 per variable), it is possible alternative parameter values were better supported than the expert opinion model simply by chance. As a unimodal peak of support for a single alternative hypothesis would be unlikely to arise randomly, however, we used this as a criterion for accepting an alternative hypothesis over the expert opinion model. We also required that the best-supported hypothesis has a significant positive correlation with genetic distance after partialling out the effects of the IBD or IBB null models. With these criteria, we excluded variables that likely reduce the fit of the IBR model relative to the null models.

Multivariate optimization

We generated a multivariate model resistance surface by summing the univariate optimized resistance surfaces meeting the criteria for inclusion in the multivariate model. To account for interactions between variables in the multivariate context, we optimized the model's performance by altering parameter values for each variable. Testing a large factorial of alternative parameter values for each variable in relation to every other variable would have required an impractical number of models to test. Instead, we optimized the function parameters for one variable while holding the other variable parameters constant. We re-tested the same set of parameter combinations performed in the univariate optimization step; however, in some cases, we expanded the range of parameter values tested as needed to reach a unimodal peak of support in the multivariate context. If optimal parameters changed in the multivariate context, and this change was still characterized by a unimodal peak or natural limit, we held the new optimal model constant and varied the parameters

of the next variable. We optimized the variables in order of decreasing correlation from the univariate optimization. If the optimal parameters changed for any of the variables in the multivariate context, we optimized each variable again in a second round of optimization.

Causal modelling

Once we identified the most highly supported IBR model following the univariate and multivariate optimizations, we compared it to the null models of IBD and IBB within a causal modelling framework (Cushman *et al.* 2006; Cushman & Landguth 2010). Specifically, we established the expectation that the causal model would retain a significant relationship with genetic isolation after partialling out the effects of the other competing models. Conversely, we expected that partialling out the causal model's effect would leave no significant relationship to genetic isolation explained by the competing models. We evaluated these expectations for the IBR, IBD and IBB models with partial Mantel's tests (Smouse *et al.* 1986) in the R software package Ecodist (Goslee & Urban 2007).

Results

Genotyping

The 161 genotypes were 97.4% complete. We re-amplified and genotyped number samples (at all loci) to estimate the error rate, which was <0.6%. Eighteen of the nineteen loci genotyped were polymorphic. We excluded the monomorphic locus URB038 from further analysis. We found no evidence for linkage disequilibrium (LD) among any of the loci after Bonferroni correction for multiple comparisons ($\alpha = 0.05$), but we observed significant departures from Hardy-Weinberg equilibrium (HWE) in 13 loci when all samples were evaluated as a single population. When a population is structured, however, disequilibrium is predicted by the Wahlund effect (Wahlund 1928). After dividing the data into Olympic, Cascade north of I90 and Cascade south of I90 regions, none of the loci showed significant departures from HWE after Bonferroni correction for multiple comparisons.

Population structure and ONP admixture

We found the strongest evidence for three populations (Fig. S2) based on the criteria proposed by the software authors; namely the value of K at which $P(X|K)$ asymptotes and also meets the expectation that subpopulation memberships are not uniform across individuals (Pritchard *et al.* 2000). The three subpopulation clusters

(Fig. S2) generally correspond to the Olympic peninsula (ONP), Cascades north of I90 (NOCA) and Cascades south of I90 (SOCA). We found no evidence for additional population substructure within the two Cascade subpopulations (Fig. S2).

All 12 ONP individuals genotyped had a >95% probability of belonging to a single ONP cluster. We observed a generally low but variable degree of admixture between the Cascade (NOCA and SOCA) and ONP subpopulations next to past translocation sites (Fig. S3). Among the 149 Cascade Range genotypes, 14 had a probability >0.25 of belonging to the ONP population and were therefore excluded from further analyses (Fig. S3). To evaluate the potential for the cut-off value to influence the genetic distances between individuals, we changed the cut-off value from 0.25 to 0.05. This reduced the sample size by 29 but did not substantially change genetic distances between individuals ($r = 0.963$). We therefore elected to keep the cut-off threshold at 0.25.

Expert opinion univariate models

The four univariate expert opinion models accounting for resistance owing to D_{et} , roads, elevation and landcover all exhibited high correlations with PCA-based genetic distance (Fig. 2). The expert opinion hypotheses for resistance owing to D_{et} and elevation, however, were not significant after partialling out the effects of IBD (Table 1).

Univariate optimization

The univariate optimized parameter values differed from the expert opinion values in all cases (Fig. 2). With the exception of D_{et} resistance, the univariate optimized alternative hypotheses retained a significant positive correlation with PCA-based genetic distance after partialling out the effects either IBD or IBB (Table 1). Roads, landcover and elevation were therefore retained in the multivariate optimization, while distance to escape terrain resistance was excluded. The univariate optimal parameters for all variables were also characterized by a unimodal peak of support or reached a natural limit (Fig. 2).

Multivariate optimization

We optimized parameter values in multivariate models in order of decreasing correlation with genetic distance. Multivariate optimization proceeded in the order of roads, landcover and elevation resistance one variable at a time while holding the other variable parameters constant. Only the multivariate optimal parameter values for roads and landcover differed from the univari-

ate optimal parameters (Fig. 3). Because variable parameters changed in the multivariate analysis, we performed a second round of optimization in the same order. In this second round, the optimal parameter values for roads changed slightly (Fig. 3), while the optimal parameters for landcover and elevation remained unchanged (Fig. S4). The final estimated resistance for each landscape variable category is listed in Table 2 and depicted in a study-wide representation in Fig. 4a.

Based on the optimized IBR resistance surface, the total circuit theory resistance between the most northerly sample location and the most southerly sample location was 25.8. The circuit theory resistance between points spanning the NOCA subpopulation was 14.2, while resistance between points spanning the SOCA subpopulation was 9.5. The circuit theory resistance between points flanking interstate 90 was 12.5 (Fig. 4b). Highly resistive landscape features appeared to constrain current in several areas (Fig. 4b).

Causal modelling

All three conceptual models of genetic isolation exhibited very high correlations with PCA-based genetic distance ($r = 0.723$, 0.686 and 0.645 for the optimized IBR, IBD and IBB models, respectively, P -value <0.0001, Table 3). Log transforming the IBD model to account for the multidimensional nature of isolation by distance did not improve the correlation ($r = 0.538$). The optimized IBR model (IBR_{opt}) appeared better supported than the null models as it retained a significant positive relationship with PCA-based genetic distance even after partialling out the effects of either IBD or IBB (Table 3). Partialling out the effects of IBR_{opt} from IBD and IBB resulted in weak or nonsignificant correlations, providing further support for IBR_{opt} as the causal model.

If we substitute the expert opinion model (IBR_{expert} – the sum of the four univariate expert opinion models) in place of IBR_{opt} in the causal modelling framework, it retained a significant correlation with PCA-based genetic distance after partialling out the effect of IBB, but not IBD. Furthermore, IBD and IBB still retained significant correlation after partialling out the effect of IBR_{expert} (Table 3). Thus, only IBD met the criteria for a causal model when compared to IBB and the IBR_{expert} model.

To explore the effect of the choice of genetic distance metric on the causal modelling outcome, we present the causal modelling results recalculated based on the proportion of shared alleles (D_{PS} ; Bowcock *et al.* 1994) and Rousset's a 2000 rather than PCA-based distance in Tables S2 and S3, respectively. While the absolute value of the correlations differed, causal modelling supported IBR_{opt} over IBD or IBB for all genetic distance metrics evaluated.

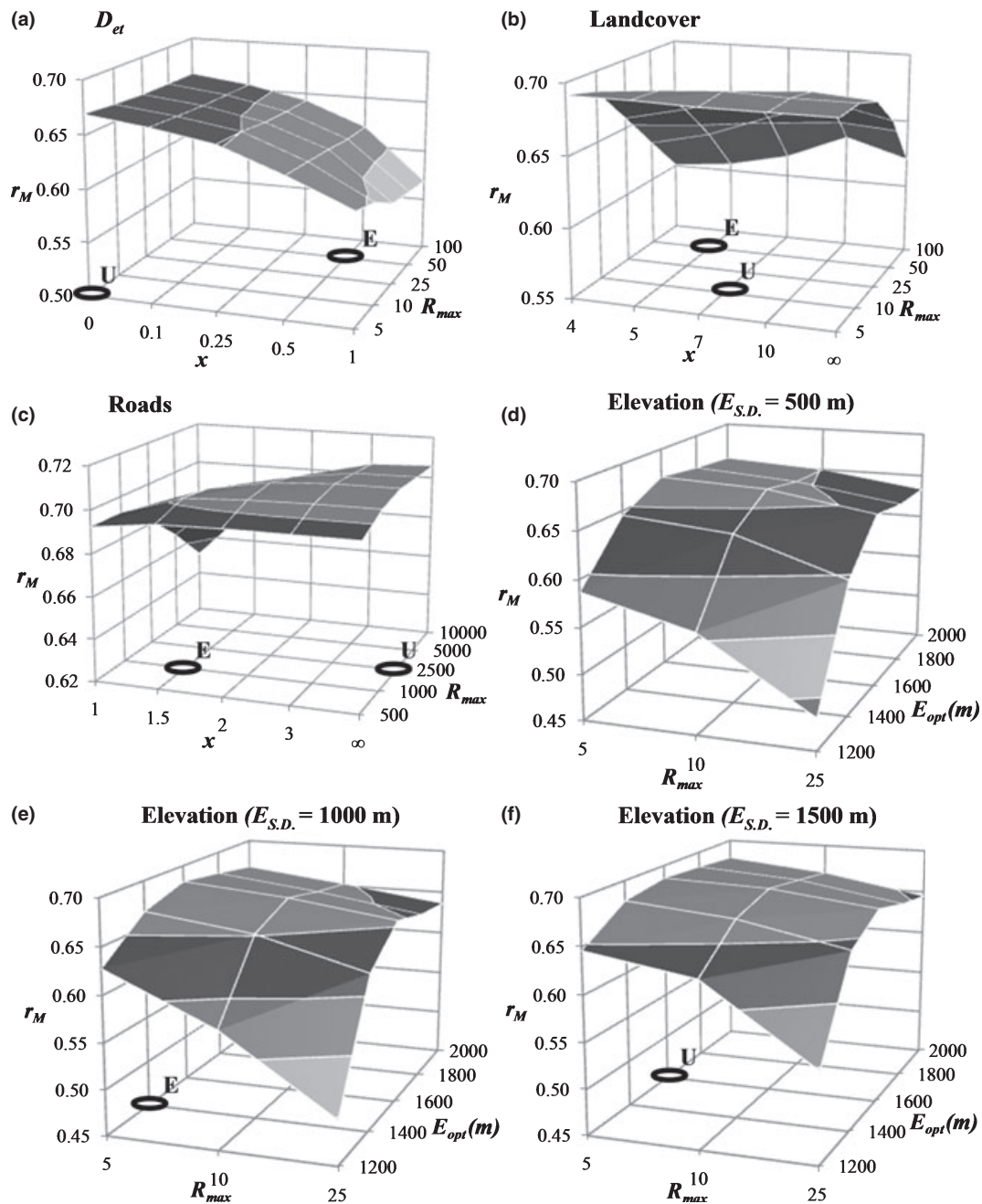


Fig. 2 Univariate IBR model optimization. The 3D contours represent the Mantel's correlation (r_M) with genetic isolation (y -axis) interpolated across the range of parameter values shown. Parameter values governing the response shape (x) and maximum resistance (R_{max}) are noted on the x -axis and z -axis, respectively, for various hypotheses relating distance to escape terrain (D_{et} ; a), landcover (b) and roads (c) to genetic isolation. We also evaluated hypotheses relating elevation to genetic isolation (panel d, e, and f). Elevation resistance parameters included the elevation standard deviation ($E_{S.D.}$; one value shown per plot), optimal elevation (E_{opt} ; z -axis on each plot), and maximum resistance (R_{max} ; x -axis on each plot). The x -axis and z -axis labels for all plots represent parameter values we evaluated (i.e. the floor of each plot represents the factorial combination of parameter values). The expert opinion hypothesis (circle labelled with E) and the univariate optimized alternative hypothesis with the highest correlation (circle labelled with U) are shown.

Table 1 Univariate IBR model relationship to genetic isolation after partialling out the effect of null models

Model	Partial r	Monte Carlo P value
Expert opinion models		
$G \sim D_{et} \mid \text{IBD}$	0.005	0.4436
$G \sim D_{et} \mid \text{IBB}$	0.250	<0.0001
$G \sim \text{Roads} \mid \text{IBD}$	0.202	<0.0001
$G \sim \text{Roads} \mid \text{IBB}$	0.391	<0.0001
$G \sim \text{Landcover} \mid \text{IBD}$	0.124	0.0006
$G \sim \text{Landcover} \mid \text{IBB}$	0.304	<0.0001
$G \sim \text{Elevation} \mid \text{IBD}$	0.002	0.4760
$G \sim \text{Elevation} \mid \text{IBB}$	0.404	<0.0001
Univariate – optimized models		
$G \sim D_{et} \mid \text{IBD}$	0.039	0.1171
$G \sim D_{et} \mid \text{IBB}$	0.363	<0.0001
$G \sim \text{Roads} \mid \text{IBD}$	0.236	<0.0001
$G \sim \text{Roads} \mid \text{IBB}$	0.389	<0.0001
$G \sim \text{Landcover} \mid \text{IBD}$	0.147	<0.0001
$G \sim \text{Landcover} \mid \text{IBB}$	0.427	<0.0001
$G \sim \text{Elevation} \mid \text{IBD}$	0.091	0.0050
$G \sim \text{Elevation} \mid \text{IBB}$	0.420	<0.0001

Partial Mantel's test correlation with genetic distance (G) for each expert opinion or univariate optimal model of landscape resistance after partialling out the effects of the null models of isolation by distance (IBD) or isolation by barrier (IBB) are listed. The P values are based on 9999 Monte Carlo randomizations of the row/column order.

Discussion

Landscapes shape gene flow in a variety of ways. Habitats highly connected by dispersal may give rise to simple patterns of isolation by distance (e.g. Koopman *et al.* 2007), strong barriers may impose a discrete subpopulation structure (e.g. Allentoft *et al.* 2009), or gradients of environmental variables influencing dispersal may yield complex patterns of isolation by resistance (e.g. Cushman *et al.* 2006). In this study, we present a novel analytical framework designed to evaluate the relative support for isolation by distance, barriers and multiple alternative landscape resistance hypotheses with the goal of identifying the model most related to the observed pattern of genetic isolation. As with all model selection processes, reducing complex ecological phenomena to relatively simple models necessarily involves decisions that balance complexity with practical realities. These decisions profoundly shape the framework's ability to identify a landscape model strongly related to the process of gene flow.

The multivariate hypothesis space

Two important aspects of a model selection framework include the number of dimensions comprising the mul-

tivariate hypothesis space and the density with which that space is sampled. More dimensions sampled at higher density permits evaluation of more complex or more highly optimized landscape resistance models. In some cases, greater complexity may not be necessary. For example, Schwartz *et al.* (2009) identified a landscape resistance model related to wolverine gene flow based on a single variable (spring snow depth) in one dimension (resistance assigned to low snow depth pixels) after sampling the hypothesis space seven times. Cushman *et al.* (2006) explored the fit of a more complex model for black bear gene flow in northern Idaho based on four variables (slope, elevation, landcover and roads) in a total of four dimensions (each variable varied by the degree of resistance assigned) after sampling the hypothesis space 108 times. In this study, we considered mountain goat gene flow to be a function of several potentially interacting environmental variables that may exhibit nonlinear relationships with landscape resistance. We also considered the potential for the range of resistance values (from 1 to R_{max}) to differ between variables. In total, our hypothesis space contained nine dimensions (R_{max} and x for landcover, roads and D_{et} plus R_{max} , E_{opt} and E_{SD} for elevation), which we sampled 310 times (120 times in the univariate optimization and 95 times in each of two multivariate optimization rounds). The added complexity of this modelling framework appeared warranted, given the interactions, strongly nonlinear optimal relationships to genetic isolation, and differences in R_{max} between variables we observed in the multivariate optimized IBR model. For species that respond in similarly complex ways to their environment, failure to account for this complexity in a modelling framework may result in models that are less informative to connectivity conservation efforts.

A role for expert opinion in landscape genetic models

Although the increased number of dimensions in our model selection framework likely improved the fit of the model selected, it also presented a challenge. Landscape genetics studies published to date have systematically sampled the hypothesis space at predefined intervals. Systematically testing the full factorial of multiple hypotheses in each of our nine dimensions would require an impractical number of models to evaluate. Instead, we specified only a single starting hypothesis (based on expert opinion), and, through the univariate and multivariate optimization steps, sampled alternative hypotheses in the direction favouring improved model fit. In this way, we greatly reduced the number of hypotheses tested. Allowing feedback from the model optimization process to define the hypothesis

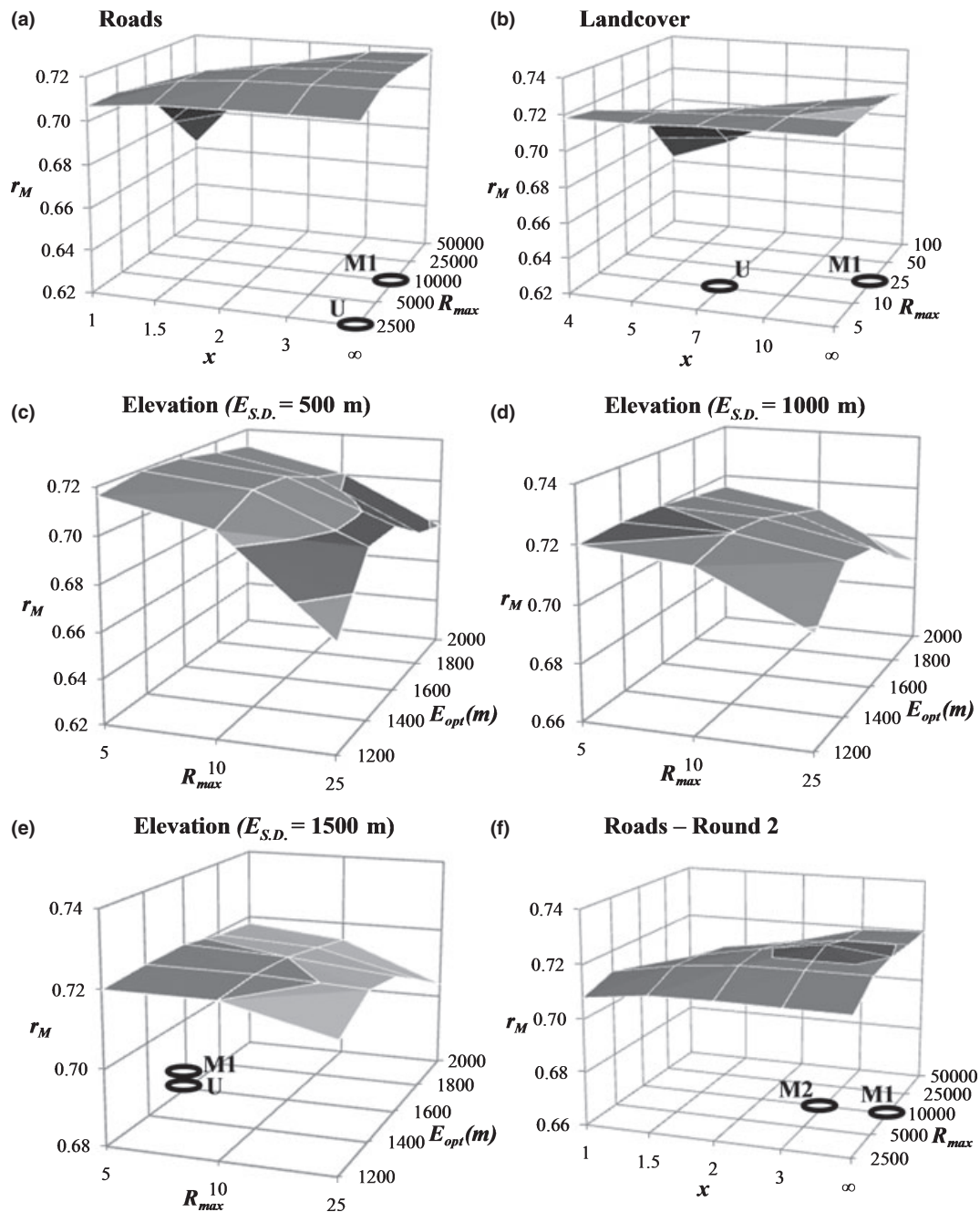


Fig. 3 Multivariate IBR model optimization. The 3D contour plot represents the Mantel's correlation (r_M) with genetic isolation (y -axis) interpolated across the range of parameter values shown. We first optimized road resistance parameters (a), including the response shape (x) and maximum resistance (R_{max}) noted on the x -axis and z -axis, respectively. Next, we optimized landcover resistance parameters (b), including the response shape (x) and maximum resistance (R_{max}) noted on the x -axis and z -axis, respectively. Finally, we optimized elevation resistance parameters (c, d, and e), which included the elevation standard deviation ($E_{S.D.}$; one value shown per plot), optimal elevation (E_{opt} ; z -axis on each plot) and maximum resistance (R_{max} ; x -axis on each plot). The x -axis and z -axis labels for all plots represent parameter values we evaluated (i.e. the floor of each plot represents the factorial combination of parameter values). The univariate optimal parameter values (circled and labelled U) and the multivariate optimized alternative parameter values with the highest correlation (circled and labelled M1) are shown. Because the parameter values changed in the multivariate context for one or more variables, we performed a second round of multivariate optimization in the same order. In the second round, only the road resistance parameters changed slightly (f; Second round optimal values labelled M2), while the optimal parameters for landcover and elevation did not change (see Fig. S4).

Variable – Pixel type	Expert opinion	Univariate optimized	Multivariate optimized
Elevation – >3300 m*	100 000	100 000	100 000
Elevation – <3300 m†	1–5	1–5	1–5
Landcover – Urban/Ag.*	100 000	100 000	100 000
Landcover – Water	50	10	25
Landcover – Mesic/disturbed forest	16	2	1
Landcover – Montane forest	2	1	1
Landcover – Ice	1	1	1
Landcover – Subalpine	1	1	1
Landcover – Alpine	1	1	1
Roads – Interstate	1000	2500	10 000
Roads – Highway	89	1	80
Roads – Secondary	1	1	1
D_{et} – 0 to 600 + m‡	1–50	-	—

*Urban and agriculture landcover pixels, as well as elevations > 3300 m were considered impermeable to gene flow and assigned a fixed resistance value (not optimized) of 100 000.

†Elevation resistance was one near the optimal elevation and increased according to the standard deviation of elevation to R_{max} at higher and lower elevations.

‡ D_{et} resistance was one nearest to escape terrain and increased with distance as a function of x and R_{max} up to 600 m. Beyond 600 m, D_{et} resistance was constant at R_{max} .

Table 2 Landscape resistance values. Resistance values (per pixel) for the expert opinion, univariate optimized and multivariate optimized IBR models are listed

space also added tremendous flexibility in finding the optimal model. Potentially any combination of parameter values could be evaluated with this approach. This flexibility likely improved our ability to identify a model strongly related to gene flow, as we did not anticipate the highly nonlinear responses detected for all variables, nor the magnitude of resistance owing to interstate highways. Starting with expert opinion parameters also allowed us to directly test the assumptions of expert opinion, grounding this approach in classical hypothesis testing and bridging the gap between empirical studies and those based on expert knowledge alone.

Univariate and multivariate IBR model optimization

We chose to divide the IBR model optimization process into two steps. In the first step, univariate optimization of model parameters provided three important pieces of information. First, it allowed for a comparison of models generated from the univariate optimum parameter values to null models of isolation by distance or barrier. With this information, we excluded variables from the multivariate analysis that were not likely to improve support for the IBR model relative to the null models. Second, the univariate optimization provided a plot of model performance as a function of alternative parameter values (Fig. 2). We used this to exclude variables whose univariate optimum model appeared correlated with genetic isolation by chance (i.e. not characterized by a unimodal peak of support), a con-

cern given the number of models we tested. Eliminating variables from the multivariate analysis if their parameters did not meet the earlier criteria also reduced the dimensionality of the multivariate optimization step. This, in turn, simplified the task of finding a multivariate optimum set of parameter values while likely improving the fit of the multivariate model. Third, the univariate optimization step provided a ranking of each variable's importance (based on the strength of their correlation with genetic isolation). This information determined the order of the multivariate optimization step. Optimizing the variable with the greatest univariate correlation first was a practical approach that likely decreased the number of rounds of optimization required to reach a stable model compared to an unordered process.

Combining univariate optimized models yielded a starting point for the multivariate optimization step, where we sought to account for interactions between variables and thereby optimize the IBR model in the multivariate context. In this study, the optimal parameters changed for several of the factors during multivariate optimization, resulting in a slight increase in the model's correlation with genetic isolation. The slightly higher correlation masks rather large changes for some model parameters, particularly for roads. The minor change in correlation for such a large change in parameter values is a result of the relative rarity of road pixels in the landscape. Interstate highways represent only a very small proportion of the total study area, requiring a correspondingly greater

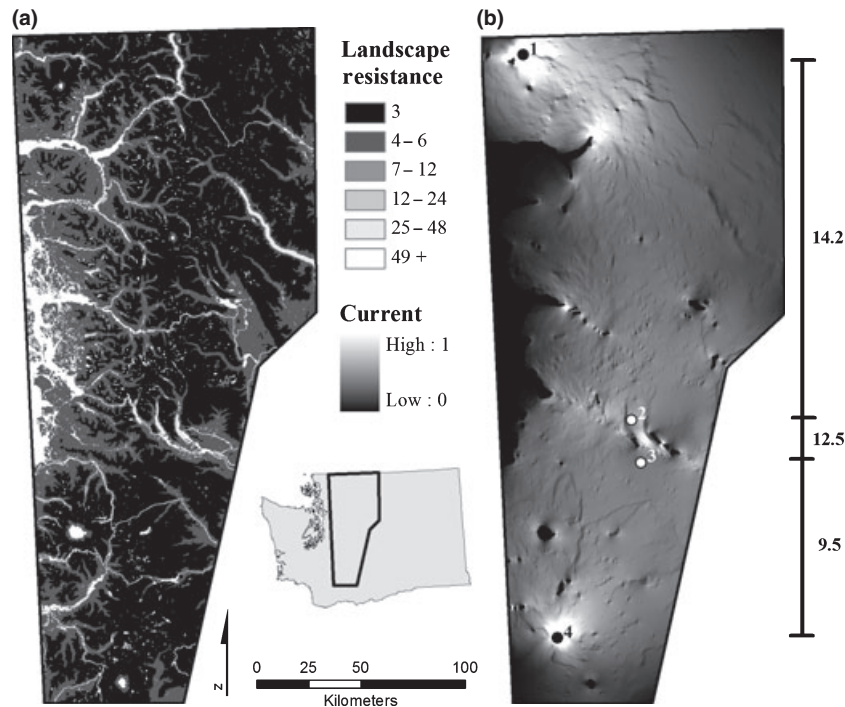


Fig. 4 (a) Optimized IBR resistance model. The multivariate optimized IBR model is depicted, with the highest resistance in white and lowest resistance in black. The values represent the sum of resistance owing to roads, landcover and elevation. (b) Circuit theory current (analogous to gene flow). The predicted current flow after injecting one ampere of current into the most northerly sample location (point 1) and connecting the most southerly sample location (point 4) to ground is shown (based on the optimized IBR resistance model). Zero current flows through areas where landscape resistance is very high (shown in black). Moderate levels of current flow through areas where gene flow is relatively unconstrained. In areas where highly resistive features constrain flow, current increases (approaching white where current is maximum). Because current is constrained by the location where it is injected and the ground location where it exits, current is also high near points 1 and 4. The total circuit theory resistance between points 1 and 4 is 25.8. Circuit theory resistances for subsets of the study area, including intermediate points (point 2 and 3) flanking interstate 90 are shown to the right of the current flow map.

resistance assigned to this pixel type to influence the effective landscape resistance between sampling locations. Thus, a particular feature's resistance and rarity in the landscape interact to shape the overall effect on the model.

Five important limitations of this model optimization framework should be noted: (i) In cases where a species' response to one or more environmental variables is multimodal, exploring the hypothesis space from a single starting point could potentially identify a local peak of support for a particular IBR hypothesis rather than a global peak of support. Particularly in these instances, the model selected may depend heavily on the initial starting point. (ii) As we explored the hypothesis space, we sampled relatively coarse intervals for each parameter. Consequently, we did not finely resolve the optimal parameter values. (iii) We evaluated alternative hypotheses in the multivariate optimization step one variable at a time while holding other variables constant. As the combinations tested do not represent the full factorial of all hypotheses for every

variable, it is possible a more optimal multivariate hypothesis went untested, or that a different order of variable optimization could produce a different result. (iv) It may not be practical to optimize all aspects of the IBR model (e.g. we assigned fixed ranks to different landcover types and a fixed resistance to certain habitats thought to be absolute barriers to mountain goat gene flow). Thus, the optimized parameters of the IBR model and its performance relative to null models could be influenced by the value assigned to these unoptimized components. (v) While optimization clearly improved the fit of the model to the observed genetic isolation, the model may be overfit. An overfit model may provide a spatially explicit prediction of landscape resistance but fail to offer a robust ecological inference regarding the actual relationship between landscape variables and their resistance to gene flow. Exploring this issue with model cross-validation using independent data (as in Braunisch *et al.* 2010) and simulations (as in Cushman & Landguth 2010) would be a useful addition to this framework in future studies.

Table 3 Causal modelling. The Mantel's test correlation with genetic distance (G) and the associated *P* value are given for each causal model of genetic isolation (isolation by landscape resistance, IBR, for both the multivariate expert opinion and multivariate optimized models; isolation by distance, IBD; and isolation by barrier, IBB)

Model	<i>r</i>	Monte Carlo <i>P</i> value
Simple mantel		
G ~ IBR _{opt}	0.723	<0.0001
G ~ IBR _{expert}	0.676	<0.0001
G ~ IBD	0.686	<0.0001
G ~ IBB	0.645	<0.0001
Partial mantel		
G ~ IBR _{opt} IBD	0.303	<0.0001
G ~ IBR _{opt} IBB	0.426	<0.0001
G ~ IBR _{expert} IBD	0.000	0.4950
G ~ IBR _{expert} IBB	0.327	<0.0001
G ~ IBD IBR _{opt}	-0.006	0.5916
G ~ IBD IBR _{expert}	0.181	<0.0001
G ~ IBD IBB	0.419	<0.0001
G ~ IBB IBR _{opt}	0.076	0.0007
G ~ IBB IBR _{expert}	0.375	<0.0001
G ~ IBB IBD	0.300	<0.0001

The partial correlations with genetic distance, controlling for the alternative models, are also shown. *P* values are based on 9999 Monte Carlo randomizations of the row/column order.

Causal modelling

Comparison of the optimized IBR model, IBD and IBB based on simple Mantel's tests alone would lead one to conclude they were all strong models. Yet causal modelling clearly supported only the optimized IBR hypothesis. This demonstrates the need to rigorously evaluate the full spectrum of potential causal models of genetic isolation, rather than only a single model (Cushman & Landguth 2010). Importantly, using causal modelling to assess the expert opinion IBR model resulted in the highest support for IBD. This highlights the importance of the model optimization process, even if the increase in correlation is relatively small. It also demonstrates how a priori assumptions regarding the relationship between landscapes and genetic isolation may not reflect actual ecological relationships and argues for caution when basing landscape and wildlife management decisions on expert opinion models alone.

A key statistical tool used in the causal modelling approach we employed is the partial Mantel's test. These tests have been controversial on the grounds that they may inflate type I error (Raufaste & Rousset 2001; Castellano & Balleto 2002). Yet Cushman & Landguth's (2010) evaluation of partial Mantel's tests within a causal modelling framework demonstrated very high

statistical power to reject alternative models and accept the correct model with low type I and type II error rates. Notably, this evaluation also demonstrated that partial Mantel's tests were able to correctly identify the causal model even when there was a high degree of collinearity among the alternative causal models.

Individual vs. population-based models

Another important aspect of landscape genetic studies is the unit of observation. Classical population genetic theory focuses on the unit of the population or subpopulation. This requires the subjective task of assigning sampled individuals to discrete groups, even if the population is continuously distributed (Schwartz & McKelvey 2009). Furthermore, population-based metrics of genetic distance such as linearized F_{ST} assume an island model of population structure (Slatkin 1995), which may not be representative of the true structure. Here, we used an individual-based approach which obviates the need for assigning individuals to a population while dramatically increasing the number of observations with which to make correlations. This, in turn, reduces the number of samples required to achieve high statistical power, which is particularly advantageous for rare or difficult to sample species.

Genetic distance and landscape distance metrics

Yet another important component of a landscape genetic model selection framework is the metric used to capture landscape and genetic distances. A commonly used estimate of effective landscape distance is based on the accumulated resistance along a least cost path (LCP) (e.g. Arnaud 2003; Chetkiewicz & Boyce 2009). Important limitations to LCP distances are the unfounded assumptions that dispersing animals have perfect knowledge and perception of landscape resistance, only use a single corridor, and encounter equal resistance whether the corridor is wide or constrained (Adriaensen *et al.* 2003; McRae 2006). In this study, we used a recently developed effective landscape distance metric based on circuit theory. In contrast to the LCP approach, circuit theory assumes a random walk dispersal behaviour, simultaneously integrates the contribution of all possible pathways to gene flow and attributes greater resistance to narrow corridors than wide corridors (McRae 2006). This more robust theoretical foundation may explain why landscape distances derived from circuit theory were more strongly related to genetic isolation in comparison with LCP distances in both simulated and real landscapes (McRae 2006; McRae & Beier 2007), though the limited nature of these comparisons has not fully resolved this issue.

While the circuit theory distance metric provides theoretical advantages, it comes at a cost of greater computer processing requirements. Considering the number of locations in our sample set and the scale of our study area, we found a pixel size of 450 m to be the maximum practical resolution. This may not have been appropriate to model landscape features that occur at a finer spatial scale. Escape terrain in particular may occur in narrow cliff bands that were not captured at our coarser resolution and may explain why the most supported model of IBR for this important mountain goat habitat variable did not outperform the null models. The latest versions of Circuitscape offer improved processing efficiency (B. McRae, personal communication) and continued increases in computer memory use efficiency as well as processor speed will make this issue less relevant to future studies.

To quantify genetic distance, we used a metric based on principal components analysis. A major advantage of this approach is that alleles that capture the greatest proportion of the genetic variation within a population have a correspondingly greater contribution to genetic distance than common alleles, which capture relatively little of a population's genetic variation. In theory, this makes PCA a more sensitive method to detect genetic dissimilarity than other methods, such as the proportion of shared alleles (DPS; Bowcock *et al.* 1994) or (Rousset's *a* 2000) where all alleles contribute equally. In a limited comparison, we found correlations with landscape resistance models to be higher overall with the PCA-based metric, though the causal modelling outcome was the same regardless of which genetic distance we used (Figs S3 and S4). While the theoretical advantages of PCA-based genetic distance are compelling, the statistical basis for its use in landscape genetics and its performance relative to other metrics requires further study.

Mountain goat gene flow in the cascade range, Washington

Surprisingly, resistance to gene flow within the study area did not appear to increase with distance to escape terrain. While the grain of our analysis may not have been capable of detecting such an effect (as discussed previously), there may be other explanations for why this important habitat variable was not accounted for in the optimized IBR model. For example, optimizing dispersal paths to maximize proximity to escape terrain may not be possible for individuals dispersing through unfamiliar habitat. Alternatively, as escape terrain is abundant within the study area, this important habitat variable may not be a limiting factor for dispersal and therefore offer no additional resistance to gene flow.

Elevation appeared to play a modest role in shaping gene flow within the study area. Cascade Range mountain goats frequently migrate from lower elevation winter habitat to higher elevation summer habitat in late spring and summer (Rice 2008). During this time, mountain goats may disperse to new patches of habitat, potentially resulting in gene flow. Dispersal and migration to summer habitat is facilitated by the receding winter snow pack, which presumably decreases the energetic costs to movement and increases foraging success (Festa-Bianchet & Côté 2008). Though mountain goat habitat use as a function of elevation is variable throughout the study area, the least resistive elevation range of the optimized IBR model (from 1000 to 2200 m) generally encompasses both winter and summer habitat (C. Rice, WDFW, personal communication). Thus, the lowest elevation resistance aligns with the elevations likely to be used by dispersing mountain goats when the snow pack recedes. At higher elevations, increased resistance (up to 5) likely corresponds to greater snow depth, poorer foraging success and greater energetic costs. At lower elevations, resistance may increase because of the generally poor adaptation of this species to low elevation habitats.

Similar to elevation, landcover also appeared to play a modest role in shaping gene flow within the study area. Though mountain goats are commonly considered alpine habitat specialists, subalpine, montane and mesic forests offered no additional resistance to gene flow in the optimized IBR model. This is congruent with the adaptation of coastal North American mountain goats to forested environments, where they may spend much of the winter on steep slopes beneath the canopy (Festa-Bianchet & Côté 2008). Water was the only optimized landcover type that appeared to resist mountain goat gene flow. The resistance attributed to water (25) was far less than that of crossing a highway (80) or interstate (10 000), suggesting at least smaller water bodies like those found within the study area are not major impediments to gene flow. Indeed, mountain goats are capable swimmers and have been observed crossing major lakes and rivers. In addition to the resistance owing to water pixels, we a priori assigned a high fixed resistance prohibiting gene flow through urban and agriculture pixels. These landcover types are associated with high human population densities and highly modified landscapes, both of which are strongly avoided by mountain goats. Such areas rarely occur within the study area, minimizing their influence on the IBR model. In a few instances, however, urban and agricultural areas are configured such that they constrain gene flow. This is evident in a few locations on Fig. 4B, where current is increased in narrow corridors of suitable dispersal habitat between small towns, and about

50 km south of Mount Baker (Point 1), where agriculture and development likely forces gene flow in a wide arc to the east.

Together, the influence of elevation gradients and landcover patterns we inferred based on the optimized IBR model suggests a clinal population structure only moderately more resistive to gene flow than IBD (where the resistance of all pixels is one). In contrast, I90 appears to greatly reduce gene flow and impose a sharp genetic discontinuity on this otherwise clinal population. Major interstates are notorious resistors of gene flow for many species (Coffin 2007). Its significance to the genetic structure of this population is supported by the high resistance assigned to it in the optimized IBR model and the sharp boundary coinciding with I90 we detected using Structure (Fig. S3). Despite the small area, this feature occupies (nearly invisible in Fig. 4), its resistance is on par with the total resistance spanning the north or south Cascades. Yet I90 does not appear to be a complete barrier to gene flow, as we detected recent admixture between the north and south Cascades (Fig. S1), and the IBB model (which posits complete isolation between the north and south) was not supported by causal modelling. In contrast to interstates, the resistance attributed to highways within the study area suggests this road class does not greatly impede gene flow. Indeed, mountain goats have regularly been observed crossing these highways. Similarly, mountain goat gene flow appears to move easily over minor roads (with no additional resistance based on the optimized IBR model).

These findings have important consequences for the conservation of mountain goats in the Cascade Range, Washington. The population has declined by up to 70% over the past 50 years, likely because of unsustainable harvest rates (Rice & Gay 2010). Though harvest has been dramatically reduced for over two decades, the population has not shown significant recovery across most of the range, and large areas of formerly occupied habitat remain sparsely occupied (C. Rice, WDFW, unpublished). Recovery and recolonization of habitat will likely depend heavily on dispersal from the few remaining large herds. Yet recent landscape changes, particularly the high resistance values associated with anthropogenic modifications, suggest greater resistance to dispersal now than was present in the historic landscape. This combination of demographic decline and reduced gene flow would be predicted to greatly reduce the local effective population size throughout the range. As a result, we would expect increased rates of inbreeding, reduced allelic diversity and concomitant loss of fitness and adaptive potential over time. The quantitative and spatially explicit understanding of landscape resistance gained from the optimized IBR

model is currently being used to inform habitat connectivity planning in an attempt to improve the population viability of mountain goats in the Cascade Range, Washington.

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Supporting information

Additional supporting information may be found in the online version of this article.

Table S1 PCR Conditions and multiplex groups

Table S2 Causal modeling results based on proportion of shared alleles (D_{PS}) genetic distance

Table S3 Causal modeling results based on Rousset's a genetic distance

Fig. S1 A Gaussian function relating elevation to landscape resistance. This function has three parameters: R_{max} (resistance ranges from 1 to R_{max}), E_{opt} (the optimal elevation where resistance is lowest), and E_{SD} (the standard deviation about the optimal elevation defining the shape of the response curve).

Fig. S2 Population structure based on Structure.

Fig. S3 Structure population assignments and fractional probability of subpopulation membership (Q).

Fig. S4 The second round of multivariate IBR model optimization for landcover and elevation following the road optimization (depicted in Fig. 3) is shown.

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