1	Urbanization impacts apex predator gene flow
2	but not genetic diversity across an urban-rural divide
3	
4	Trumbo DR <sup>1</sup> , Salerno PE <sup>1</sup> , Logan KA <sup>2</sup> , Alldredge M <sup>3</sup> , Gagne RB <sup>4</sup> , Kozakiewicz CP <sup>5</sup> , Kraberger S <sup>4</sup> ,
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#### 19 Abstract

20 Apex predators are important indicators of intact natural ecosystems. They are also sensitive to 21 urbanization because they require broad home ranges and extensive contiguous habitat to support their 22 prey base. Pumas (*Puma concolor*) can persist near human developed areas, but urbanization may be 23 detrimental to their movement ecology, population structure, and genetic diversity. To investigate 24 potential effects of urbanization in population connectivity of pumas, we performed a landscape genomics 25 study of 134 pumas on the rural Western Slope and more urbanized Front Range of Colorado, USA. Over 26 12,000 single nucleotide polymorphisms were genotyped using double-digest, restriction site-associated 27 DNA sequencing (ddRADseq). We investigated patterns of gene flow and genetic diversity, and tested for 28 correlations between key landscape variables and genetic distance to assess the effects of urbanization and 29 other landscape factors on gene flow. Levels of genetic diversity were similar for the Western Slope and 30 Front Range, but effective population sizes were smaller, genetic distances were higher, and there was 31 more overall population substructure in the more urbanized Front Range. Forest cover was strongly 32 positively associated with puma gene flow on the Western Slope, while impervious surfaces restricted 33 gene flow and more open, natural habitats enhanced gene flow on the Front Range. Landscape genomic 34 analyses revealed differences in puma movement and gene flow patterns in rural versus urban settings. 35 Our results highlight the utility of dense, genome-scale markers to document subtle impacts of 36 urbanization on a wide-ranging carnivore living near a large urban center.

37

38 Keywords: landscape genomics, gene flow, genetic diversity, effective population size, urbanization,
39 *Puma concolor*

#### 41 Introduction

42 Urbanization is a major threat to biodiversity, and in particular to apex predators with broad home 43 ranges (Cohen 2003; Theobald 2005; Crooks et al. 2017). Habitat fragmentation due to urbanization can 44 have important impacts on predator movement, disease, and survival (Markovchick-Nicholls et al 2008; 45 Carver et al. 2016; Fountain-Jones et al. 2017). This reduced connectivity can lead to smaller, more 46 isolated populations, where less gene flow and genetic diversity, as well as smaller effective population 47 sizes (Riley et al. 2006; Vandergast et al. 2007; Ernest et al. 2014) ultimately cause local and regional 48 extirpations through environmental and demographic stochasticity and inbreeding depression (Allendorf et 49 al. 2013). Moreover, increased human recreational activities in wildlife habitats associated with nearby 50 urbanization can change wildlife movement patterns and habitat usage, exacerbating the impacts of 51 fragmentation (McKinney 2002; Lewis et al. 2015). As human populations continue to expand worldwide, 52 urban areas are becoming larger and more extensive on the landscape. However, we do not fully 53 understand how urbanization affects natural ecosystems near wildland-urban interfaces (Radeloff et al. 54 2005; Magle et al. 2012).

55 Large carnivores are important indicators of intact natural ecosystems, as they require an abundant 56 and sustainable prev base, as well as high habitat connectivity to support their broad home ranges (Sergio 57 et al. 2006, 2008). However, understanding the effects of urbanization on large carnivores is difficult due 58 to their low population densities and secretive nature (Logan and Sweanor 2001; Riley et al. 2006; 59 Hornocker and Negri 2009). Camera traps, radio-telemetry, and GPS collars provide valuable information 60 on animal home ranges and population sizes (e.g., Lewis et al. 2015; Blecha et al. 2018), but these studies 61 are expensive, time consuming, and can only monitor a small fraction of the total population for limited 62 time periods. Population and landscape genetics can provide additional, complementary techniques for a 63 more detailed understanding of wildlife populations (Epps et al. 2007; Lowe and Allendorf 2010; 64 Balkenhol et al. 2016). Genetic studies provide an indicator of functional landscape connectivity through 65 measures of gene flow, effective population sizes of breeding individuals, and cost-efficient monitoring of

66 genetic diversity across broad geographic areas (McRae et al. 2005; Solberg et al. 2006). Moreover, recent 67 high-throughput sequencing technologies enable the genotyping of many more thousands of loci than 68 previously possible, providing higher power to detect the often subtle population genetic structure of wide-69 ranging species such as large carnivores (Luikart et al. 2003; Holderegger et al. 2006). 70 Pumas (Puma concolor; other common names include mountain lions, cougars, panthers, 71 catamounts) are a large, apex predator with one of the broadest latitudinal ranges of any terrestrial 72 carnivore, spanning western North America, Central America, and South America (Hornocker and Negri 73 2009). Pumas are sensitive to urbanization, requiring broad-scale landscape connectivity to persist, and are 74 thus useful indicators for monitoring the effects of urban fragmentation (Beier 1995; Crooks 2002; 75 Maletzke *et al.* 2017). Given sufficient habitat area and landscape connectivity, however, pumas can still 76 persist within and adjacent to urban systems (Wilmers et al. 2013; Riley et al. 2014; Lewis et al. 2015; 77 Zeller et al. 2017; Blecha et al. 2018). Furthermore, the substantial area requirements of large carnivores 78 such as pumas can enhance their role as "umbrella" species, whose protection also benefits co-occurring 79 species through broad-scale habitat preservation (Thorne et al. 2006). 80 The southern Rocky Mountains in western Colorado, USA support natural habitats with high 81 puma densities, as well as many rural and urban human developments (Hornocker and Negri 2009). The 82 Western Slope of the Rocky Mountains primarily consists of large areas of contiguous public wildlands 83 with an abundant prey base for pumas, interspersed with small rural and exurban developments, including 84 the Uncompany Plateau region near the town of Montrose (Western Slope Study Area; Figure 1). In 85 contrast, the Front Range is a rapidly urbanizing, major metropolitan area on the Eastern Slope of the 86 Continental Divide, where urbanization is spreading from lower elevation areas in and around the Denver 87 Metropolitan Area into adjacent wildland habitats in the foothills of the Rocky Mountains. Pumas continue 88 to persist near this wildland-urban interface, including adjacent to the city of Boulder on the western edge 89 of the Denver Metropolitan Area (Front Range Study Area; Figure 1; Lewis et al. 2015; Moss et al. 90 2016a). From 2010 – 2017, Colorado was the 8th fastest growing U.S. state by population (577,829

91 residents added) and the 6th fastest by percentage (11.5% population growth; U.S. Census Bureau 2017), 92 with most of this growth occurring along the eastern edge of the Front Range. Thus, comparative studies 93 of puma movement and gene flow in one of the most populous states in the mid-continental USA, which 94 also supports a robust puma population, can provide insight into the effects of urbanization on this 95 important apex predator.

96 Here, we tested how different landscape factors, including urbanization, enhance or restrict gene 97 flow and genetic diversity in a large apex predator across an urban-rural divide in Colorado, USA. A large 98 sample (n = 134) of pumas were utilized from (a) the rural Western Slope and (b) the more urbanized 99 Front Range (Figure 1). We used double digest restriction site associated DNA sequencing (ddRADseq) to 100 genotype pumas at 12,444 single nucleotide polymorphism (SNP) loci to evaluate the potential differences 101 in gene flow, effective population sizes, genetic diversity, and population structure in these two different 102 landscapes. We tested landscape genomic hypotheses by correlating key landscape factors with puma 103 genetic distance measures. We hypothesized that pumas in the more urbanized Front Range would have 104 (a) smaller effective population sizes, (b) lower levels of genetic diversity, and (c) more landscape factors 105 related to urbanization that restrict gene flow, relative to the rural Western Slope landscape.

106

#### 107 Materials and Methods

#### 108 Samples and sequences

Puma blood and tissue samples were collected as part of ongoing monitoring efforts by Colorado Parks and Wildlife in both the Western Slope and Front Range regions of the southern Rocky Mountains of Colorado, USA (Figure 1; Lewis *et al.* 2015; Carver *et al.* 2016). Samples were collected from 2005-2014 on the Western Slope and 2007-2013 on the Front Range. Western Slope samples consisted of 36 males and 42 females, and Front Range samples consisted of 24 males, 31 females, and 1 puma of unknown sex. Our sampling represents a large proportion of the resident pumas present in both regions during the sampling period, as Lewis *et al.* (2015) estimated 14.4 (S.E. 1.6) and 14.7 (S.E. 1.3) resident

pumas occupying the Western Slope and Front Range study areas at a single time point, respectively, frommotion camera and telemetry data collected in 2009 and 2010.

118 Genomic DNA was extracted from tissue or blood using QIAGEN DNeasy Blood & Tissue kits 119 (QIAGEN Inc., Valencia, CA). We genotyped a total of 78 individuals from the Western Slope and 56 120 individuals from the Front Range using the ddRADseq protocol described in Peterson et al. (2012) and 121 sequenced on Illumina HiSeq 2500 and 4000 machines (Illumina, San Diego, California) using 100bp 122 single-end sequencing at the University of Oregon Genomics Facility (gc3f.uoregon.edu). We tested 9 123 different combinations of restriction enzymes on puma samples for digestion efficiency and evaluated the 124 size ranges of fragment distributions using an Agilent Tapestation 2200 (Agilent Genomics, Santa Clara, 125 California). We chose the digest enzymes EcoRI-HF (6bp recognition) and NlaIII (4bp recognition) and a 126 target fragment size range of 300–400 bp (excluding adapters). We used a Blue Pippin with a 2%, internal 127 standard, 100-600 bp gel cartridge (Sage Science, Beverly, Massachusetts) for size selection and a 128 biotinylated P2 adapter with DynaBeads<sup>®</sup> (Peterson *et al.* 2012) to purify the polymerase chain reaction 129 (PCR) template for the final enrichment. PCR was performed for 12 cycles and five reactions were tested 130 for each pool of individuals. We initially genotyped 16 individuals multiplexed into an Illumina 2500 131 HiSeq lane to estimate maximum multiplexing based on a target of >12X coverage per locus. After 132 assessment of locus coverage, we proceeded to multiplex 48 and 70 individually-barcoded samples on 133 Illumina 2500 and 4000 HiSeq lanes, respectively, using the Peterson et al. (2012) flex adaptors.

134

135 Bioinformatics pipeline and filters

136 We evaluated read quality for each sequencing lane using FastQC

137 (bioinformatics.babraham.ac.uk) and assembled our SNP dataset *de novo* using Stacks v 1.41 (Catchen *et* 

138 *al.* 2013). Details on Stacks code and parameter settings used are on the GitHub repository;

139 github.com/pesalerno/PUMAgenomics. We demultiplexed and filtered sequencing reads using the

140 program *process\_radtags* in Stacks. Due to sensitivity of downstream genotyping with different Stacks

141 parameter settings (Mastretta-Yanes et al. 2015; Paris et al. 2017), we incorporated individual sample 142 replicates in library preparations. In each library, we included 3 within and 3 between library replicates, 143 which were used for estimating genotyping error rates for different combinations of parameters used to 144 construct loci with the *denovo* map.pl Stacks pipeline. We ran 11 different *de novo* assemblies varying 4 145 different Stacks parameters that affect locus, allele, and SNP error rates and the number of loci genotyped, 146 consisting of (1) minimum number of identical, raw reads required to create a stack (-m), (2) number of 147 mismatches allowed between loci when processing a single individual (-M), (3) number of mismatches 148 allowed between loci when building the catalog (-n), and (4) maximum number of stacks at a single de 149 novo locus (-max locus stacks) (Table S1; Mastretta-Yanes et al. 2015). Locus error rate was calculated 150 as the number of loci present in only one of the samples of a replicate pair divided by the total number of 151 loci, allele error rate was the number of allele mismatches between replicate pairs divided by the number 152 of loci, and SNP error rate was the proportion of SNP mismatches between replicate pairs.

153 After identifying the most supported parameter settings that minimized locus, allele, and SNP 154 error rates, while maximizing the number of SNPs ( $-m = 3, -M = 4, -n = 4, max_locus_stacks = 3$ ; Table 155 S1), we exported the SNP matrix with the *populations* program in Stacks (Catchen *et al.* 2013), retaining 156 SNPs that were present in at least 20% of individuals by population, and retaining a single random SNP 157 per locus. This matrix was further filtered for missing data in Plink v. 1.07, first by locus, then by 158 individual, and then by minor allele frequency (MAF) using multiple combinations of thresholds for 159 reducing missing data in the matrix (see github.com/pesalerno/PUMAgenomics). After evaluating missing 160 data from SNP matrices, we retained the matrix with a more stringent locus filter (excluding loci missing 161 >25% individuals) and a less stringent filter on minor allele frequency (excluding loci with MAF < 0.01). 162 We additionally filtered loci that were found at position 95 (the last position of our reads) due to a higher 163 number of SNPs present in this position, suggesting increased error rates due to low sequence quality 164 towards the end of the sequencing read. In order to compare landscape resistances with putatively neutral 165 loci, we used a Principal Components Analysis (PCA) to identify loci showing strong signatures of

166 selection relative to neutral background genomic variation with the program PCAdapt (Luu et al. 2016). 167 We found twelve, putatively adaptive, outlier loci using a false discovery rate of 10%, so we filtered these 168 outliers out for downstream landscape genomic analyses to avoid confounding neutral demographic 169 patterns with patterns generated by loci under selection. 170 171 Population genomics and structure 172 Population genomic statistics were calculated for the two sampling regions, the Western Slope and 173 Front Range (Figure 1). Observed and expected heterozygosity ( $H_{obs}$  and  $H_{exp}$ ), nucleotide diversity ( $\pi$ ), 174 inbreeding coefficient ( $F_{IS}$ ), and population genetic differentiation ( $F_{ST}$ ) were calculated using the 175 *populations* program in Stacks with SNP loci that passed previous filters, excluding a single individual 176 (sample\_1382) that did not pass the 75% missing data threshold. We estimated allelic richness ( $A_r$ ) using 177 HP-RARE 1.0 (Kalinowski 2005), which corrects for variance in sample sizes using rarefaction. Two 178 complementary, individual-based genetic distances were calculated: proportion of shared alleles distance 179 (D<sub>ps</sub>; Bowcok et al. 1994) using the adegenet R v. 3.3.3 package and relatedness distance (r; Smouse and 180 Peakall 1999) using the PopGenReport R package. We then calculated mean genetic distance among 181 individuals for each region, corrected for geographic distance (i.e., genetic distance per km), since 182 individuals that are farther apart are expected to have higher genetic distances due to neutral isolation by 183 distance population processes (Wright 1942; Balkenhol *et al.* 2016). Effective population sizes ( $N_e$ ) were 184 estimated using the linkage disequilibrium method in NeEstimator v. 2.01 (Do et al. 2014), using the 185 correction for chromosome number (Waples et al. 2016), which has been shown to be a robust method for 186 inferring  $N_e$  using SNP datasets and large sample sizes (Waples 2016; Waples *et al.* 2016). We evaluated 187 overall genetic structure as well as genetic differentiation among the two sampling sites (Western Slope 188 and Front Range) using PCA and Discriminant Analysis of Principal Components (DAPC) in the R 189 package adegenet (Jombart 2008) and Admixture ancestry analysis (Alexander et al. 2009). We used the 190 function assignplot to identify individuals that were putative migrants or admixed based on the individual

DAPC assignment probabilities. We used the find.clusters command in adegenet and minimized crossvalidation error in Admixture to estimate the number of populations (i.e., K).

193

194 Landscape genomics

195 Geographic Information Systems (GIS) data were collected for different landscape factors that we 196 hypothesized would affect puma dispersal and gene flow in Colorado. Table 1 provides details on GIS 197 data sources, spatial resolution, and ecological justification for each landscape factor. Study area extents 198 were calculated and landscape variables were compared across regions by buffering individual data points 199 by a typical female puma dispersal distance of 34.6 km (Logan and Sweanor 2001), dissolving 200 overlapping buffers, and calculating zonal statistics within each region (Western Slope and Front Range) 201 using ArcGIS v. 10.1 (ESRI, Redlands, California). Landscape data were converted into resistance 202 surfaces using the Reclassify and Raster Calculator tools in ArcGIS. The following hypothesized 203 relationships of landscape factors with puma gene flow were modeled: percent impervious surface cover 204 (negative effect on gene flow), land cover (forested, open-natural, and developed: positive, neutral, and 205 negative effects on gene flow, respectively), percent tree canopy cover (positive effect), vegetation density 206 (positive effect), river and stream riparian corridors (positive effect), roads (negative effect), minimum 207 temperature of the coldest month (negative effect), annual precipitation (positive effect), topographic 208 roughness (positive effect), and elevation (negative effect). Additionally, we included an isolation by 209 geographic distance model, which would be supported if none of the landscape variables had an effect on 210 gene flow except for straight line, Euclidean distance between individuals (Wright 1942; Balkenhol et al. 211 2016). Table S2 describes methods and justification for converting raw landscape variables to resistance 212 surfaces.

Two genetic distance measures were used as response variables in landscape genomic analyses: proportion of shared alleles distance (D<sub>ps</sub>; Bowcok *et al.* 1994) and relatedness distance (r; Smouse and Peakall 1999). Environmental resistances among individuals were calculated using Circuitscape (McRae

216 2006) for each landscape resistance surface (McRae 2006; Row et al. 2017). Circuitscape resistances are a 217 useful tool in landscape genetics because they summarize all potential movement pathways 218 simultaneously, as opposed to least cost paths that evaluate only a single idealized pathway, and thus 219 assume the study organism has complete knowledge of the landscape and always chooses the ideal 220 pathway (McRae 2006; Balkenhol et al. 2015). Landscape variables were tested for multicollinearity, both 221 prior to and after calculating environmental resistances in Circuitscape, to ensure Pearson's r correlations 222 < 0.7 and variance inflation factor (VIF) scores < 5 in final landscape genomics models, as collinearity can 223 cause instability in parameter estimation in regression models (Tables S3 and S4; Warren et al. 2010; 224 Dormann et al. 2012; Rowe et al. 2017). 225 Two complementary methods were used to estimate the effects of environmental resistances on 226 genetic distances: multiple regression on distance matrices (MRDM; Legendre et al. 1994) using 227 PERMUTE v.3.4 and maximum likelihood of population effects (MLPE; Clark et al. 2002; van Strien et 228 al. 2012; Row et al. 2017) using the lme4 R package. MRDM is a permutational, distance matrix-based 229 approach that has been traditionally used in landscape genetic analyses, whereas MLPE is a newer linear 230 mixed effects modeling technique that models pairwise comparisons as a random effect and environmental 231 resistances as fixed effects (Balkenhol et al. 2016). Recent evaluations of landscape genetic approaches 232 found linear mixed effects modeling using MLPE to be more accurate, although both approaches 233 performed well (Shirk et al. 2017). Therefore, we included the traditional MRDM approach as well as 234 MLPE in order to utilize multiple, complementary techniques for inferring associations between landscape 235 features and gene flow. For MRDM and MLPE, genetic distances were the response variable and 236 environmental resistances were explanatory variables. Additionally for MLPE, a random effect matrix of 237 individual comparisons was included to control for the non-independent, pairwise structure of the data, 238 and landscape resistances were standardized to units of standard deviation centered on the mean (van 239 Strien et al. 2012; Row et al. 2017). Models were ranked using the Bayesian information criterion (BIC), 240 and top models within 5 BIC units are reported (Richards 2015).

241

- 242 **Results**
- 243 Genotyping and filtering SNP matrices

244 Initial Stacks processing retained a single random SNP per 95 bp read and SNPs present in at least 245 20% of individuals by population, resulting in a matrix of 98,813 SNPs. These SNPs were further filtered 246 in Plink by removing loci that were present in less than 75% of individuals, which resulted in a matrix of 247 20,355 SNPs. Only a single individual was excluded based on our >75% missing loci per individual 248 threshold. After excluding SNPs present in the 95<sup>th</sup> sequencing base position and with minor allele 249 frequency <0.01, we retained 12,456 SNPs. PCAdapt detected twelve outlier loci, putatively under 250 selection, while accounting for population structure (K=2). After removing these putatively adaptive loci, 251 the final neutral dataset contained 12,444 SNPs (Table S1; github.com/pesalerno/PUMAgenomics).

252

#### 253 *Population genomics and structure*

254 The two study areas encompass similar geographic extents: 11,889 km<sup>2</sup> for the Western Slope and 255 11,958 km<sup>2</sup> for the Front Range (Table 2). Measures of genetic diversity ( $H_{obs}$ ,  $H_{exp}$ ,  $\pi$ ,  $A_r$ ,) and inbreeding 256  $(F_{IS})$  were similar for the Western Slope and Front Range (Table 2). However, the effective population 257 size  $(N_e)$  was smaller, mean genetic distances among individuals (D<sub>PS</sub>/km and r/km) were higher, and there 258 was more overall population substructure in the more urbanized Front Range (Table 2, Figure 2). We also 259 calculated  $N_e$  using subsets of individuals (i.e., pre and post-2010 individuals in the Front Range, pre and 260 post-2011 individuals in the Western Slope), since multiple overlapping generations may bias effective 261 population size estimates low or high (Waples 2016; Waples et al. 2016). N<sub>e</sub> remained consistently higher 262 in the Western Slope, although it differed between the earlier and later sampling periods there, and 263 indicated the population may be expanding (Table S5). We found a detectable signature of population 264 differentiation between the Western Slope and Front Range regions based on a PCA and DAPC, and 265 Admixture ancestry analysis indicated K=2 was the best supported value of K by minimizing cross

266	validation error (Figure 2; Alexander et al. 2009). The proportion of correct individual assignment to
267	populations based on DAPC (Figure 2b), which attempts to minimize within population distances and
268	maximize between population distances (Jombart 2008), was high for most individuals in both the
269	Western Slope (0.98) and the Front Range (0.96). However, the DAPC assignplot also identified admixed
270	individuals and putative migrants between regions, including a female and a male in the Front Range that
271	assigned mostly to the Western Slope, and an admixed male in the Western Slope that assigned mostly to
272	the Front Range (Figure 2b). We also analyzed both regions separately for population substructure (Figure
273	S1), and there was no signature of population differentiation within the Western Slope or Front Range,
274	further supporting two populations.
275	
276	Landscape Genomics
277	The Front Range has more urban development than the Western Slope, with more impervious
278	surface cover and a higher density of roads (Figure 1, Table 3, Table S2). The Front Range also has more
279	tree canopy cover, higher vegetation density, and higher annual precipitation than the Western Slope
280	(Table 3), likely due to the high desert habitats (i.e., the Colorado Plateau ecoregion) in the Western Slope
281	being drier than the grassland and shrub habitats found at lower elevations of the Front Range (i.e., the

282 Great Plains ecoregion; McMahon *et al.* 2001).

283 Prior to running Circuitscape, landscape raster surfaces were largely uncorrelated (i.e., Pearson's r 284 < 0.7), with the exception of elevation, which was positively correlated with annual precipitation and 285 negatively correlated with minimum temperature of the coldest month in both regions, and vegetation 286 density, which was negatively correlated with annual precipitation in the Front Range (Table S3). After 287 Circuitscape analyses, environmental resistance variables showed more collinear relationships than raw 288 raster surfaces (Table S4), likely due to Circuitscape resistances being higher for individuals separated by 289 larger geographic distances (McRae 2006). Therefore, we removed landscape variables from both regions 290 that were strongly correlated with many other variables, until all VIF scores were less than 10 (Row et al.

291 2017). Variables retained were geographic distance, river and stream riparian corridors, roads, impervious 292 surface cover, tree canopy cover, vegetation density, and minimum temperature of the coldest month. 293 However, vegetation density was still correlated with geographic distance in both regions, and impervious 294 surface was correlated with geographic distance and tree canopy cover in the Western Slope (Table S4). 295 We removed these variables as well, resulting in Pearson's r correlations less than 0.7 and VIF scores less 296 than or equal to 4.1 and 3.5 in the Western Slope and Front Range, respectively, for all explanatory 297 variables. Thus final MRDM and MLPE models for the Western Slope included geographic distance, tree 298 canopy cover, stream and river riparian corridors, roads, and minimum temperature of the coldest month; 299 and for the Front Range included the same landscape variables plus impervious surface cover. 300 Landscape genomic patterns of pumas were different in the rural Western Slope compared to the 301 more urbanized Front Range, with the exception of geographic distance being supported in both regions 302 (Tables 4 and 5). In the Western Slope, tree canopy cover was consistently positively correlated with gene 303 flow in MRDM and MLPE models, and low minimum temperatures of the coldest month (i.e., those found 304 in high elevation, alpine tundra habitats) were negatively correlated gene flow in one MLPE model 305 (Tables 4 and 5). In contrast, in the Front Range, tree canopy cover and percent impervious surface cover 306 were negatively associated with gene flow in the top MLPE models (Table 5). Since the relationship 307 between tree cover and gene flow was the opposite of what we hypothesized in the Front Range, we also 308 inverted the tree cover resistance surface (i.e., making higher tree cover = higher resistance), reran 309 Circuitscape and MLPE analyses, and higher tree cover still showed significant negative correlations with 310 gene flow in this region. 311 312 Discussion

The apex predator puma (*Puma concolor*) persists in many urbanized regions throughout its range, yet the localized effects of recent urban sprawl remain unclear. Here, we compared patterns of genomic landscape connectivity and diversity of pumas across two regions that span an urban-rural divide in

316 Colorado, USA. Landscape genomic connectivity patterns differed between regions, such that genetic 317 distances were higher and urbanization (i.e., percent impervious surface cover) restricted gene flow in the 318 more urbanized Front Range, whereas forest cover was most important for enhancing gene flow on the 319 rural Western Slope. Despite finding reductions in gene flow associated with urbanization on the Front 320 Range, population-level genetic diversity and inbreeding measures were similar to those on the rural 321 Western Slope. This suggests that recent urban sprawl in the Colorado Front Range has not yet had a 322 substantial impact on the genetic diversity of pumas. This is in contrast to more isolated puma populations 323 in other highly urbanized landscapes such as southern California and Florida, which exhibit reduced 324 genetic diversity and strong evidence of inbreeding compared to Colorado pumas (Ernest et al. 2003, 325 2014; Johnson *et al.* 2010). However, a smaller effective population size, higher among-individual genetic 326 distances, and higher population substructure in the recently urbanized Front Range suggest habitat 327 fragmentation has already impacted this population and could cause further reductions of genetic diversity 328 as urbanization continues to expand in Colorado (Theobald 2005; U.S. Census Bureau 2017). If puma 329 populations decline, this could have important cascading effects into lower trophic levels, such as 330 overgrazing of vegetation by ungulate herbivores (Markovchik-Nicholls et al. 2008).

331

#### 332 *Population genomics and structure*

333 The Western Slope and Front Range were resolved as two genetically distinct groups (i.e., K=2; 334 Figures 1 and 2). Minimum temperature of the coldest month was also negatively associated with gene 335 flow in one of the top landscape genomic models on the Western Slope (Table 5), suggesting there may be 336 restricted gene flow through high elevation, alpine tundra habitats (McMahon et al. 2001). However, 337 potential immigrants and admixed individuals were identified moving in both directions (Figure 2) and 338 overall genetic differentiation between the two populations was low (pairwise  $F_{ST} = 0.02$ ; Table 2). Since 339 our sample archive consisted of opportunistically collected samples, our analyses were restricted to 340 populations in two distinct regions, whereas pumas occur throughout the southern Rocky Mountains in

341 Colorado. Therefore, potential immigrants and admixed individuals are not necessarily moving between 342 our specific Western Slope and Front Range study areas, but may originate from other unsampled 343 populations that share genetic ancestry with our two study regions. Nevertheless, results from our study 344 suggest pumas may be somewhat limited in dispersing across the high elevation peaks of the Continental 345 Divide, and future studies should attempt to sample more intensively across the entire region to further 346 investigate this trend.

347 We identified similar levels of genetic diversity and inbreeding between the rural Western Slope 348 and more urbanized Front Range (Table 2), suggesting urbanization is not yet having a large impact on the 349 genetic diversity of pumas in Colorado. One potential explanation is that urbanization in the Front Range 350 is primarily occurring on the eastern edge of the region, possibly creating a relatively impermeable urban 351 boundary on the eastern border, but not isolating pumas in fragments or limiting their connectivity to 352 wildland habitat to the west (Figure 1; Lewis et al. 2015; Blecha et al. 2018). Another possibility is that 353 many of the SNPs we sampled may not have high enough mutation rates to show a strong genomic 354 signature of the relatively recent effects of rapid urbanization occurring in the Front Range (Haasl and 355 Payseur 2011; Allendorf et al. 2013). As the human population continues to expand, future urbanization 356 could result in more fragmented populations and reductions in genetic diversity, as has been detected in 357 other more urbanized landscapes like southern California and Florida (Ernest et al. 2003, 2014; Johnson et 358 al. 2010).

Despite similar geographic extents and levels of genetic diversity in the Western Slope and Front Range, mean genetic distances among individuals were higher in the urban Front Range (Table 2), suggesting that fragmentation due to urbanization may be limiting puma dispersal and gene flow. In addition, a larger effective population size ( $N_e$ ) of pumas was detected on the rural Western Slope ( $N_e$ =69.3) compared to the urban Front Range ( $N_e$ =40.2; Table 2), with the caveat that some assumptions of this estimator are violated in both regions (e.g., closed populations with no immigration, nonoverlapping generations). The effect of non-overlapping generations on  $N_e$  is difficult to predict (Waples *et* 

366 al. 2016), and this assumption is expected to be violated similarly in both the Western Slope and Front 367 Range populations. Immigration, however, is expected to downwardly bias  $N_e$  by creating linkage 368 disequilibrium through a multi-locus Wahlund effect (Wahlund 1928; Waples and England 2011). Thus, it 369 is possible that the Front Range may be showing a lower  $N_e$  due to having more immigrants from outside 370 populations than the Western Slope. This is possible, and perhaps likely, given the higher overall 371 population substructure in the Front Range (Figure 2), which could indicate more potential immigrants 372 into this region. On the other hand, if immigration rates are similar for both regions, the relatively smaller 373 Front Range  $N_e$  may be due to (1) urbanization and fragmentation impacting and limiting population size, 374 and/or (2) species range limit theory (Abundant Center Hypothesis) predicting that smaller population 375 sizes are likely to occur at the edge of the geographic range relative to core areas (Brown 1984; Sagarin 376 and Gaines 2002). These potential underlying factors are not mutually exclusive and may both be acting 377 together. However, the lack of difference in most genetic diversity measures, in addition to slightly lower 378 allelic richness in the Front Range, which is the most sensitive metric to recent bottlenecks (Allendorf et 379 al. 2013), suggests lower effective population size on the Front Range may be more consistent with recent 380 urbanization impacts than historical range boundary effects.

381

382 Landscape genomics

With regard to general landscape genomics methodology, we found MRDM to be a much more conservative approach that adds fewer explanatory variables to the models than MLPE (Tables 4 and 5). Conversely, MLPE results in more complex models with more explanatory variables and higher r<sup>2</sup> values (genetic variation explained) than MRDM (Tables 4 and 5). The different genetic distance measures we used (D<sub>PS</sub> and r) showed largely consistent relationships with landscape variables, but still provided a few different insights, particularly using MLPE (Tables 4 and 5). Overall r<sup>2</sup> values were somewhat low (r<sup>2</sup> = 0.04 - 0.08 for MRDM, r<sup>2</sup> = 0.11 - 0.17 for MLPE), but this is expected for a large carnivore with extreme

long distance dispersal abilities (e.g., Short Bull *et al.* 2011, Balkenhol *et al.* 2016). Isolation by distance
was important across models for both regions (Tables 4 and 5).

392 On the rural Western Slope, tree canopy cover was most important for enhancing gene flow, 393 suggesting pumas prefer to disperse through forests rather than more open shrub and grassland habitats in 394 this landscape (Table 5). Forests provide more cover for concealment and ambush predation (Logan and 395 Sweanor 2001; Hornocker and Negri 2009; Warren et al. 2016). Use of open areas may also increase 396 susceptibility to mortality by hunters and ranchers (Newby et al. 2013), which are both more prevalent in 397 the rural Western Slope than the more urbanized Front Range. In addition, non-forested areas on the 398 Western Slope are dry, high elevation desert habitats (i.e., the Colorado Plateau ecoregion; McMahon et 399 al. 2001), which may provide less prey and water resources, and thus be poorer habitats for hunting and 400 dispersal (Sweanor et al. 2000; McRae et al. 2005; Dickson et al. 2013).

401 In the more urbanized Front Range, impervious surface cover restricted gene flow (Table 5). This 402 suggests urbanization is limiting gene flow, despite high levels of genetic diversity (Table 2). Similarly, 403 Lewis et al. (2015) found pumas were less likely to be detected in habitats with residential development, 404 even low-density exurban developments, which are increasingly encroaching into the foothills of the Front 405 Range region. Genetic studies on pumas from more urbanized and fragmented populations in southern 406 California and Florida have detected strong inbreeding and isolation associated with urbanization (Ernest 407 et al. 2003, 2014; Johnson et al. 2010; Riley et al. 2014). Our study detected more subtle impacts of 408 urbanization in a less fragmented landscape, within mountainous wildland habitats adjacent to a major 409 metropolitan center, which experiences high levels of human outdoor recreation activities such as hiking 410 and skiing (Figure 1). In addition, in contrast with the rural Western Slope and contrary to our initial 411 hypotheses, forest cover was negatively associated with gene flow on the Front Range (Table 5). This 412 pattern suggests pumas are more willing to disperse through open shrub and grassland habitats in this 413 region. The reasons for this are unclear, but pumas living in the more developed Front Range may be more 414 acclimated to human activities and thus more willing to travel outside of forested habitats, demonstrating

415	that pumas have a range of adaptable behaviors and will use and move through different types of habitat
416	(Dickson et al. 2005; Blecha et al. 2018). Pumas may also be hunting more urban mesopredators,
417	domestic, and agricultural animals in these open habitats on the more developed Front Range, which was
418	shown in a prior study using stable isotope analysis of Front Range puma diets (Moss et al. 2016b). There
419	is also less hunting of pumas in the Front Range compared to the rural Western Slope, so pumas may be
420	less wary of open areas, although this effect would be expected to be counteracted in part by higher traffic
421	mortality in the more urbanized region (Beier 1995; Crooks 2002).
422	

423 Conclusions

424 Our findings are consistent with prior comparative landscape genetic studies that have revealed 425 varying effects of landscape factors on movement and gene flow across different portions of a species' 426 geographic range (e.g., Vandergast et al. 2007; Short Bull et al. 2011; Trumbo et al. 2013). We found that 427 in the rural Western Slope with high hunting pressure, forests with high tree canopy cover are most 428 important for conserving puma genetic connectivity. In contrast, in the more urbanized Front Range, non-429 forested habitats such as shrublands and grasslands habitats are utilized for dispersal and gene flow, 430 effective population sizes are smaller, genetic distances among individuals are higher, and gene flow is 431 being restricted by urbanization (Tables 2, 4, and 5). Next generation sequencing techniques can provide 432 dense, genome-scale SNP datasets of thousands of putatively neutral markers, which gives researchers 433 increased power to detect the often subtle effects of landscape factors, such as urbanization, on gene flow 434 (Luikart et al. 2003; Lowe and Allendorf 2010; Allendorf et al. 2013). This is particularly important for 435 wide-ranging species with broad geographic distributions, since landscape effects on gene flow occur at 436 broader geographic scales and may be weaker and more difficult to detect compared to more dispersal-437 limited species with smaller home ranges (Holderegger et al. 2006; Epps et al. 2007; Balkenhol et al. 438 2016). Indeed prior work on pumas using 16 microsatellites found no population structure across the 439 southern Rocky Mountains of Colorado and northern New Mexico (McRae et al. 2005). Our results

440 demonstrate that large SNP datasets can allow researchers to identify impacts of urbanization on gene

- flow, effective population sizes, and patterns of population genetic structure of wide-ranging species, even
- before fragmentation is extensive enough to greatly reduce genetic diversity. Maintaining genetic
- 443 connectivity in these "umbrella" species can have outsized benefits towards conserving biodiversity, since
- 444 preserving broad swaths of contiguous habitats that are necessary for their persistence also benefits many
- 445 other species with smaller home ranges and narrower habitat requirements (Sergio *et al.* 2006, 2008;
- 446 Thorne *et al.* 2006).
- 447

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- 452

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- 624

## 625 Author Contributions

626

D.R.T. performed laboratory work, analyzed landscape and population genomic data, and wrote the
manuscript; P.S., R.B.G., C.P.K, S.K., and N.F.J. performed laboratory work and analyzed landscape and
population genomic data; K.L. and M.A. directed fieldwork and collected field data; M.E.C., S.C., H.B.E.,
K.C., S.V., and W.C.F. conceived of study questions and directed research; and all authors contributed
input to draft and final versions of the manuscript.

## 633 Data Accessibility

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635 ddRADseq data used in genomic analyses will be uploaded to Dryad (datadryad.org) upon acceptance of636 the manuscript for publication.

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## 638 Supporting Information639

**Table S1**: Library replicate analysis of error rates from different Stacks parameter settings for minimum number of identical raw reads required to create a stack (-m), number of mismatches allowed between loci when processing a single individual (-M), number of mismatches allowed between loci when building the catalog (-n), and maximum number of stacks at a single de novo locus (-max\_locus\_stacks). Locus error rate was the number of loci present in only one of the samples of a replicate pair divided by the total number of loci, allele error rate was the number of allele mismatches between replicate pairs divided by the number of loci, and SNP error rate was the proportion of SNP mismatches between replicate pairs.

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**Table S2**: Landscape resistance transformations for the Western Slope and Front Range.

Table S3: Correlations (Pearson's r) between environmental raster surfaces used in landscape genomic
analyses for the Western Slope and Front Range regions of Colorado. Pearson's correlations > 0.7 are in
bold.

Table S4: Correlations (Pearson's r) between Circuitscape environmental resistances used in landscape
 genomic analyses for the Western Slope and Front Range regions of Colorado. Pearson's correlations >
 0.7 are in bold.

Figure S1: Principle Components Analyses (PCAs) and Admixture plots of (a) 78 Western Slope pumasand (b) 56 Front Range pumas, analyzed separately within each region.

## 661 662 Tables

#### 663 Table 1: Environmental variables used for landscape genomic analyses, data sources, spatial resolution,

#### 664 and ecological justification.

Category	Landscape Variable	Code	Description	Data Source, Spatial Resolution	Calculation	Ecological Justification
Distance	Isolation by geographic distance	Geo. dist.	Euclidean, straight- line distance between individuals	No environmental data; model assumes only distance affects gene flow, 30 meter	ArcGIS Reclassify tool, Circuitscape	Model of isolation by straight-line distance (Wright 1942).
Land cover	Land cover: forested, open- natural, and developed	Land cover	Multiple land cover categories collapsed into 3 costs of movement: forested (lowest), open natural areas (medium), and developed (highest)	National Land Cover Database (mrlc.gov/nlcd2011.php; Homer <i>et al.</i> 2011), 30 meter	ArcGIS Spatial Analyst	Forested habitats provide the most cover for hunting and dispersal, open natural areas are intermediate, and developed areas are the least suitable habitat for dispersal (Crooks 2002; Lewis <i>et al.</i> 2015).
	Percent impervious surface cover	Imperv.	Percentage of impervious surface	National Land Cover Database (mrlc.gov/nlcd2011.php; Homer <i>et al</i> . 2011), 30 meter	ArcGIS Spatial Analyst	Human development results in increased noise, lights, and hunter access, limiting dispersal (Riley <i>et al.</i> 2006; Ernest <i>et al.</i> 2014; Maletzke <i>et al.</i> 2017).
	Road corridors	Roads	Roads, with 50 meter buffers on each side	Colorado Department of Transportation (dtdapps.coloradodot.info /otis), 30 meter	ArcGIS Analysis Tools, Spatial Analyst	Roads increase mortality, noise, lights, and hunter access, limiting dispersal (Riley <i>et al.</i> 2006; Newby <i>et al.</i> 2013; Maletzke <i>et al.</i> 2017).
	River and stream riparian corridors	Riparian	River and stream riparian corridors, with 50 meter buffers on each side	National Hydrography Dataset (nhd.usgs.gov), 30 meter	ArcGIS Analysis Tools, Spatial Analyst	River and stream riparian corridors provide vegetative and topographical cover for dispersal, as well as water sources attracting prey species (Naiman <i>et al.</i> 1993; Hilty and Merenlender 2004; Dickson <i>et al.</i> 2005).
Vegetation	Percent tree canopy cover	Tree cover	Percentage of tree canopy cover	National Land Cover Database (mrlc.gov/nlcd2011.php; Homer et al. 2011), 30 meter	ArcGIS Spatial Analyst	Low tree canopy limits cover for ambush predation and concealment, and restricts dispersal (Sweanor <i>et al.</i> 2000; Logan and Sweanor 2001; Warren <i>et al.</i> 2016; Blecha <i>et al.</i> 2018).
	Enhanced vegetation index	Veg. density	Density of vegetation calculated from chlorophyll reflectance in visual and near- infrared spectra	Moderate Resolution Imaging Spectroradiometer (modis.gsfc.nasa.gov), 250 meter	ArcGIS Spatial Analyst	Low vegetation density limits cover for ambush predation and concealment, and restricts dispersal (Sweanor <i>et al.</i> 2000; Hilty and Merenlender 2004; Warren <i>et al.</i> 2016; Blecha <i>et al.</i> 2018).
Climate	Minimum temperature of the coldest month	Min. temp.	Mean annual minimum temperature of the coldest month (°C) calculated from	Global Climate Data (worldclim.org/bioclim; Hijmans <i>et al.</i> 2005), 1 kilometer	ArcGIS Spatial Analyst	Low minimum temperatures and high snowfall, found at high elevation mountain ridgelines (e.g., alpine tundra habitats) restrict

			1970-2000 weather station data, interpolated between stations			hunting, breeding, and dispersal (Hornocker and Negri 2009).
	Mean annual precipitation	Ann. precip.	Mean annual precipitation accumulation (mm) calculated from 1970-2000 weather station data, interpolated between stations	Global Climate Data (worldclim.org/bioclim; Hijmans <i>et al.</i> 2005), 1 kilometer	ArcGIS Spatial Analyst	Dry habitats with low precipitation accumulation limit prey species for hunting and vegetative cover, restricting dispersal (Logan and Sweanor 2001; McRae <i>et al.</i> 2005).
Topography	Topographic roughness	Topo. rough.	Topographic complexity based on variance in elevation within a moving window	National Elevation Dataset (lta.cr.usgs.gov/ned) National Map Tool (viewer.nationalmap.gov), 30 meter	Geomorphometric and Gradient Metric Toolbox (Cushman et al. 2010), ArcGIS Spatial Analyst	Steep, topographically- complex canyons and mountain slopes provide cover for hunting and dispersal (Dickson <i>et al.</i> 2005; Hornocker and Negri 2009).
	Elevation	Elev.	Elevation calculated from digital elevation models.	National Elevation Dataset (lta.cr.usgs.gov/ned) National Map Tool (viewer.nationalmap.gov), 30 meter	ArcGIS Spatial Analyst	Low minimum temperatures and high snowfall, found at high elevation mountain ridgelines (e.g., alpine tundra habitats) restrict hunting, breeding, and dispersal (Hornocker and Negri 2009).

668	Table 2: Study areas (km <sup>2</sup> ), number of individuals genotyped (N <sub>gen</sub> ), and population genomic parameter
669	estimates from the Western Slope and Front Range of Colorado. Population genomic measures are

670 observed heterozygosity ( $H_{obs}$ ), expected heterozygosity ( $H_{exp}$ ), nucleotide diversity ( $\pi$ ), allelic richness

 $(A_r)$ , inbreeding coefficient (F<sub>IS</sub>), genetic differentiation among populations (pairwise F<sub>ST</sub>), mean genetic

distance among individuals corrected for geographic distance (D<sub>PS</sub> and r per km) with standard errors

673 (S.E.), and effective population size (N<sub>e</sub>) with 95% confidence intervals (C.I.) based on parametric

674 bootstrapping.

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Region	Area (km²)	$N_{gen}$	$H_{obs}$	$H_{exp}$	π	A <sub>r</sub>	$F_{IS}$	$F_{ST}$	D <sub>PS</sub> /km (S.E.)	r/km (S.E.)	N <sub>e</sub> (95% C.I.)
Western Slop	pe 11889	78 indiv.	0.240	0.272	0.0029	1.93	0.117	0 024	0.28 (0.05)	0.15 (0.03)	69.3 (66.2-72.4)
Front Range	11958	56 indiv.	0.242	0.263	0.0028	1.89	0.084	0.024	0.46 (0.16)	0.24 (0.09)	40.2 (38.7-41.7)

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# Table 3: Habitat differences between the Western Slope and Front Range of Colorado. Units are percent cover for impervious surface and tree canopy cover; resistance values for land cover, river and stream

680 riparian corridors, and roads; degrees Celsius for temperature; millimeters for precipitation; meters for

elevation; and unitless measurements based on chlorophyll reflectance and variance in elevation,

respectively, for enhanced vegetation index and topographic roughness.

683

Landscape Data	Western	Western Slope					Front Range			
	Min	Max	Median	Mean	Std Dev	Min	Max	Median	Mean	Std Dev
Elevation (m)	1453.5	4362.9	2354	2418.0	552.5	1474.5	4347.1	2365	2374.9	629.3
Tree canopy cover (%)	0	100	20	29.9	31.4	0	100	32	35.0	33.6
Impervious surface (%)	0	100	0	0.5	4.1	0	100	0	4.0	13.5
Minimum temp. coldest month (°C)	-20.2	-9.5	-13.3	-13.9	2.8	-19.9	-8.3	-12.7	-12.6	2.8
Annual precipitation (mm)	208	1137	458	483.3	171.5	359	1006	452	496.4	121.9
Enhanced vegetation index	-1806	8955	4634	4434.9	1858.3	-1969	9132	5416	4957.8	1800.9
Topographic roughness	0	27924.6	11	53.1	129.6	0	20067.0	25	56.2	100.8
Landcover	1	10	1	3.2	3.0	1	10	1	4.2	3.7
Roads	1	10	1	1.7	2.5	1	10	1	2.7	3.6
Riparian	1	10	10	9.4	2.3	1	10	10	9.4	2.3

686 Table 4: Multiple regression on distance matrices (MRDM) landscape genomic results from the Western Slope and Front Range of Colorado. Response variables were individual-based genetic distances, i.e., 687 688 proportion of shared alleles (Dps) and relatedness (r). Explanatory variables, after removing correlated 689 variables, were the geographic (Euclidean) distance model (geo. dist.), percent impervious surface cover, 690 percent tree canopy cover, river and stream riparian corridors, roads, and minimum temperature of the 691 coldest month. Forward selection followed by backward elimination was performed, with 1,000 random 692 permutations of the dependent distance matrix per step, using Bonferroni-corrected p-to-enter and p-to-693 remove alpha values of 0.05. Standardized beta coefficients were used to assess the direction of effect of 694 each landscape variable on gene flow. Only univariate models were supported. 695

Region	Genetic distance	Landscape factors	Direction of effect	r <sup>2</sup>	р
Western Slope	Dps	tree cover	+	0.08	0.001
	r	geo. dist.	-	0.04	0.001
Front Range	Dps	geo. dist.	-	0.05	0.001
	r	geo. dist.	-	0.04	0.001

696 697

**Table 5**: Maximum likelihood of population effects (MLPE) landscape genomic results from the Western

699 Slope and Front Range of Colorado. Response variables were individual-based genetic distances, i.e.,

proportion of shared alleles (Dps) and relatedness (r). Pairwise comparisons of individuals were controlled

as a random effect. Fixed effects, after removing correlated variables, were the geographic (Euclidean)

distance model (geo. dist.), percent impervious surface cover, percent tree canopy cover, vegetationdensity, river and stream riparian corridors, roads, and minimum temperature of the coldest month.

704 Standardized beta coefficients were used to assess the direction of effect of each landscape variable on

705 gene flow. Models reported are within the top 5 BIC units. Landscape factors are in order of standardized

706 beta coefficients (largest to smallest).

707

Region	Genetic distance	Landscape factors	Direction of effect	r <sup>2</sup>	∆BIC
Western Slope	Dps	tree cover	+	0.15	0
		min. temperature	-		
		tree cover	+	0.15	0.6
		geo. dist.	-		
		tree cover	+	0.14	3.0
	r	geo. dist.	-	0.17	0
		tree cover	+		
		geo. dist.	-	0.17	3.8
Front Range	Dps	geo. dist.	-	0.12	0
		tree cover	-		
		impervious surface	-		
		geo. dist.	-	0.11	2.9
		tree cover	-		
	r	geo. dist.	-	0.13	0
		tree cover	-		
		impervious surface	-		
		geo. dist.	-	0.13	0.5
		tree cover	-		



747 Figure 1: Study area in the Western Slope and Front Range of the southern Rocky Mountains of 748 Colorado, USA. Landscape genomic analyses included 78 pumas from the Western Slope and 56 pumas 749 from the Front Range (white circles). Resistance surfaces, shown for the Western Slope, represent 750 alternative hypotheses of the effects of landscape variables on puma dispersal and gene flow (red=high 751 gene flow, blue=low gene flow) for: (a) percent impervious surface cover (negative effect on gene flow), 752 (b) land cover (forested, open-natural, and developed: positive, neutral, and negative effects on gene flow), 753 (c) percent tree canopy cover (positive effect), (d) vegetation density (positive effect), (e) river and stream 754 riparian corridors (positive effect), (f) roads (negative effect), (g) minimum temperature of the coldest 755 month (negative effect), (h) annual precipitation (positive effect), and (i) topographic roughness (positive 756 effect). We also tested isolation by geographic Euclidean distance. Land cover base maps show forests 757 (green), shrub and grasslands (tan), urban areas (red), agriculture and ranchlands (brown and yellow), and 758 alpine tundra (grey).



Figure 2: Population structure from (a) Principal Components Analysis (PCA), (b) Discriminant Analysis

787 of Principal Components (DAPC), and (c) Admixture analysis. Individuals assigned to the Western Slope 788 and Front Range are green and blue, respectively. K=2 was most supported in Admixture analysis using

789 cross validation error.