RESEARCH ARTICLE



Effects of habitat deterioration on the population genetics and conservation of the jaguar

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Received: 2 September 2014/Accepted: 31 July 2015/Published online: 4 September 2015 © Springer Science+Business Media Dordrecht 2015

Abstract Over the past century, human activities and their side effects have significantly threatened both ecosystems and resident species. Nevertheless, the genetic patterns of large felids that depend heavily on large and well-conserved continuous habitat remain poorly studied. Using the largest-ever contemporary genetic survey of wild jaguars (*Panthera onca*), we evaluated their genetic diversity and population structure in natural (Brazilian Amazon) and highly modified habitats (e.g. Cerrado, Caatinga) including those close to the northern

Electronic supplementary material The online version of this article (doi:10.1007/s10592-015-0766-5) contains supplementary material, which is available to authorized users.

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(Yucatan, Mexico) and southern (Pantanal) edge of the species' distribution range. Data from our set of microsatellites revealed a pronounced genetic structure, with four genetically differentiated geographic areas. Geographic distance was not the only factor influencing genetic differentiation through the jaguar range. Instead, we found evidence of the effects of habitat deterioration on genetic patterns: while the levels of genetic diversity in the Amazon forest, the largest continuum habitat for the species, are high and consistent with panmixia across large distances, genetic diversity near the edge of the species distribution has been reduced through population contractions. Mexican jaguar populations were highly differentiated from those in Brazil and genetically depauperated. An isolated population from the Caatinga showed the genetic effects of a recent demographic decline (within the last 20-30 years), which may reflect recent habitat degradation in the region. Our results demonstrate that the jaguar is highly sensitive to habitat fragmentation especially in human-dominated landscapes, and that in Brazil, the existing but limited genetic connectivity in the central protected areas should be maintained. These conclusions have important implications for the management of wide-ranging species with high dispersal and low population density. The restoration of ecological connectivity between populations over relatively large scales should be one of the main priorities for species conservation.

Keywords Felid · Elusive · Habitat deterioration · Connectivity · Conservation

Introduction

Human impacts on ecosystems have increased dramatically throughout the world over the last century. Anthropogenic modifications of habitat (i.e., loss and fragmentation) that



Table 1 Sampling sites (n = 14) in the different biomes of the jaguar distribution in Mexico and Brazil, number of field-collected faeces after DNA extraction (N faeces) and other material (N other), species identification (N species ID), number of jaguar faeces (N jaguar),

number of jaguar individuals (N ind): in bold, total number of jaguars after the assignment strategy for both faeces and high quality DNA sources, and geographical coordinates. na: not applicable

Biome	Code biome	Sampling areas	Code area	N faeces	N other	N Species ID	N jaguar	N ind	Coordinates	
AMAZON	AMZN							12		
		Adolfo Ducke Reserve	DUCK	104	0	56	21	6	02°55′S	59°59′W
		Uatumã Biological Reserve	UATM	29	0	19	6	3	1°46′S	-59°16′W
		Maracá Ecological Station	MARA	19	0	13	2	1	3°24′26′'N	61°29′13′′W
		Viruá National Park	VIRU	46	0	33	8	2	1°29′9″ N	61°2′10′′ W
CAATINGA	CAPV	Capivara National Park	CAPV	82	0	57	53	18	8° 26′S	42° 19′W
CERRADO	CENT							14		
		Araguaia	ARAG	na	1 skin	na	na	1	3 25′13″	53 26'26"
			ARAG	na	11 liver	na	na	3	to 18 15'40"S	to 47 53'07"W
		Parque Estadual do Cantão	CANT	na	4 blood	na	na	4		
		Das Emas National Park	PNEM	61	0	49	14	3	18°19′S	52°45′W
			PNEM		3 blood	na	na	3		
PANTANAL	PANT	Refúgio Ecológico Caiman	PANT	98	0	79	37	34	19°57′S	56°18′W
			PANT	na	22 blood	na	na	22		
MEXICO	MEXC							24	Latitudes	Longitudes
		Ecological reserve El Zapotal	ZAPT	68	0	60	40	5	21°20′25″N	87°36′20″W
		Ecological reserve El Eden	EDEN	64	0	44	25	3	21° 13′ N	87°11 W
		Ejido20Noviembre	EJNV	4	0	3	0	0		
		Calakmul	CALK	18	0	16	5	3	18°11′05″N	89°44′ 49″W
		Petcacab	PETC	21	0	17	10	4	19°17′ 15″N	88°13′32.7″W
		Ejido Caobas	CAOB	34	0	27	14	9	18°14′N	89°03′W
			CAOB	na	6 skin	na	na	6		
			CAOB	na	1 blood	na	na	0		
TOTAL				209	50	167	94	102		

impact population size and connectivity can result in genetic erosion, which may seriously compromise the fitness of populations and increase extinction risk (Saccheri et al. 1998; Ceballos 2002; Frankham 2003, 2005; Reed et al. 2003; Palomares et al. 2012). In addition, the spatial distribution of populations and their dynamics may also be important in shaping the patterns of genetic diversity throughout a species' range. Models have suggested the vulnerability of natural populations would be determined in part by their spatial distribution (peripheral vs. core populations) because it directly influences the genetic variability and abundance of a species (Gyllenberg and Hanski 1992). In this context, one can predict that at the scale of species spatial distribution, the most vulnerable populations would be in areas impacted by both the demographic effects (i.e., location at the edge of the species' range) and environmental deterioration (i.e., habitat fragmentation and loss). Genetic analyses may provide early warning signals for the demographic consequences of these processes and provide specific recommendations for the design of effective conservation strategies.

Large felids have extensive home ranges and usually depend on well-conserved continuous habitat for reproduction and dispersal. They are thus particularly vulnerable to habitat degradation (Crooks 2002). During the last century, most of these charismatic species have experienced declines in population size worldwide, and the accelerated human-mediated habitat degradation (i.e., loss and fragmentation) and synergic effects of direct persecution such as hunting may be severely threatening their long-term survival (Nowell and Jackson 1996; Perez 2001; IUCN 2010). While population surveys of elusive carnivorous felid species are a challenge (Williams et al. 2002, Thompson 2004), genetic studies are even more limited by the difficulty of obtaining an adequate number of samples. As a result, the genetic patterns of many large felids and their responses to landscape scale habitat disturbance, including fragmentation and degradation, remain poorly studied. Improvements to non-invasive genetic testing through sampling of faeces can promote broader scale surveys in the near future (Janecka et al. 2008; Roques et al. 2011, 2014). However, to date, genetic studies on



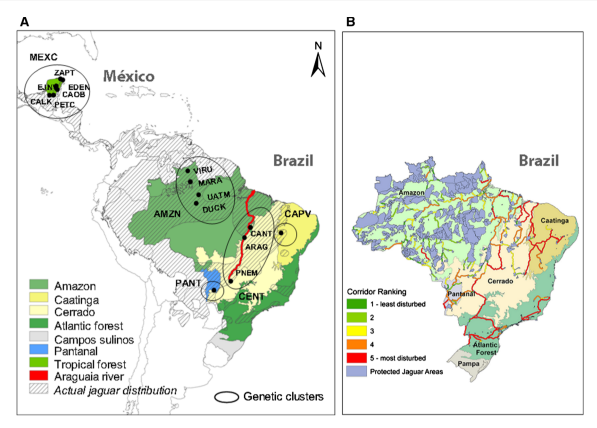


Fig. 1 a Map of the actual jaguar's geographic range (*Panthera onca*), sampling sites (*black points*), genetic clusters and principal ecosystems in Brazil and Mexico (see details and codes in Table 1). The map is based on information from the IUCN Red List of

Threatened Species (IUCN 2013). **b** Map of the potential corridors connecting protected jaguar populations in Brazil and degree of disturbance from Silveira et al. (2014)

declining populations of large carnivores are limited primarily to medium and small spatial scales, such as Amur tiger (*Panthera tigris altaica*, Henry et al. 2009; Alasaad et al. 2011), jaguar (*Panthera onca*, Eizirik et al. 2001; Moreno et al. 2006; Haag et al. 2010), leopard (*Panthera pardus*, Dutta et al. 2013) and tiger (*Panthera tigris*, Reddy et al. 2011; Joshi et al.2013; Sharma et al. 2013).

The jaguar is the largest felid in the American continent and the third-largest cat worldwide. Historically, its range encompassed a large area extending from the southwestern USA through the Amazon basin to the Rio Negro in Argentina, but today it occupies only about 50 % of this range (Mittermeier et al. 1998; Zeller 2007; Sanderson et al. 2002; Fig. 1). Years of poaching and livestock conflicts during the last century associated with massive rates of deforestation have reduced and severely fragmented the species' habitat and distribution (Zeller 2007). As a result, the IUCN classifies the jaguar as Near Threatened with declining population trends (IUCN 2010). Most of the loss of range has occurred at the edges in northern Mexico and Southwestern United States, and northern Argentina (Sanderson et al. 2002). In Brazil, which constitutes approximately 50 % of the current jaguar range (Zeller 2007), the Amazon rainforest and the Pantanal floodplains are thought to harbor the two largest continuous jaguar populations worldwide (Sanderson et al. 2002). However, there is extensive deforestation and development in Brazil, especially in the highly impacted southern Cerrado and Caatinga biomes, at the eastern limit of the jaguar distribution.

The first large-scale phylogeographic study of the jaguar was based on the analyses of mitochondrial DNA (mtDNA) control region sequences and 29 nuclear microsatellite loci of 44 individuals sampled from Mexico to southern Brazil (Eizirik et al. 2001). It revealed a low level of genetic differentiation in the species throughout its geographic range. This pattern of genetic homogeneity was interpreted as the result of a rather recent population expansion, about 300,000 years ago, followed by a history of demographic connectivity on a continental scale. The only partition observed between the northern and southern areas of the range was attributed to a reduced historical gene flow across the Amazon River, although such a reduced connectivity was not supported by a more recent study (Moreno et al. 2006).

The continued destruction and fragmentation of its habitat suggest that many jaguar populations likely became



demographically isolated and genetically depauperated in recent years. It appears that past and recent large-scale habitat loss and fragmentation have been sufficiently strong to promote genetic differentiation of jaguars in the Atlantic forest regions (Haag et al. 2010). Therefore, it is critical to gain a better understanding of genetic patterns and recent demographic processes at both local and large scales and to compare core and peripheral populations within the distribution range of the species.

In this study, we report on populations from Mexico and Brazil where jaguars are still found at high densities and in areas representing both highly modified, peripheral, as well as preserved core habitats. The results represent one of the most extensive genetic analyses of contemporary samples of jaguars to date. We assessed the genetic structure and diversity of jaguar populations from diverse areas, tested whether jaguars are still genetically connected throughout the entire distribution range, and evaluated the potential genetic consequences of habitat fragmentation on populations. Finally, we discuss the importance of potential corridors within Brazil and the Yucatan Peninsula in Mexico and the implications for conservation priorities.

Materials and methods

Study areas, samples and genotyping

Non-invasive genetic samples of jaguars were obtained by collecting faeces in several areas of Mexico and Brazil (Fig. 1a; Table 1, Supplementary Material S1, S2). We collected in six different areas in the Yucatan Peninsula, which is close to the northern limit of the jaguar's distribution and includes the largest remaining tract of tropical forest in Mexico. In Brazil, we sampled areas with relatively high densities of jaguars and large extensions of natural or semi-natural habitat, both in Pantanal and Amazon forests, and populations in the Cerrado and Caatinga biomes where the areas are highly modified, have a high human population density, and are less suitable for jaguars. Faeces were collected in four different areas in the Brazilian Amazon, which represents the largest area of relatively continuous jaguar habitat (Sanderson et al. 2002). Pantanal is used primarily for extensive cattle ranching, and is less affected by habitat fragmentation than areas with intensive agriculture. Sampling was carried out at the caiman ecological reserve (PANT), a cattle ranch and ecotourism center located in the southern Pantanal (Mato Grosso do Sul State). The Cerrado biome, originally covered by extensive areas of neotropical savannas and dry forest, has been severely fragmented by the agricultural activities of the last 50 years. Samples were obtained around three areas located within the Cerrado, and along the Araguaia river: the Emas National Park (ENP), one of Brazil's largest reserves located in the transition area with the Amazon biome; in Tocantins State, the Araguaia national Park (ANP) and the Cantão State Park (CSP), the only large conservation unit where jaguars are protected. The Caatinga of eastern Brazil represents the eastern limit of jaguar distribution in South America (Sanderson et al. 2002) and one of the most fragmented habitat remnants of the species in Brazil. Unique to Brazil, the Caatinga is a large and one of the most diverse regions of dry forests and arid scrubland of the world, but the high human population density has completely or partially transformed over 50 % of its area (Casteleti et al. 2005).

Sampling was carried out in one of the most important protected areas of the Caatinga, the Serra da Capivara National Park (CAPV). Sampling of faeces in all areas was conducted mostly during the dry season between 2007 and 2009 with the exception in the Adolfo Ducke Reserve (DUCK), where samples were also collected in 2004 and 2005. In all sites, faeces were collected by inspecting roads and trails frequently used by humans or animals, except in Parque Estadual do Cantão (CANT), Araguaia (ARAG), PANT, and PNEM where scat detector dogs were used to find samples (Vynne et al. 2011b). Faeces were collected in sterilized plastic vials with approximately 30 ml of absolute alcohol, subsequently transferred to 100-ml plastic jars containing silica pellets (Roeder et al. 2004), and stored at room temperature until DNA extraction. Most samples collected in the Amazon were put directly in silica gel without the first step involving an alcohol solution.

We also obtained blood samples from captured individuals (Table 1). Skin samples collected in 2007 were also obtained from ARAG, Brazil and from Ejido Caobas (CAOB) in Mexico. DNA isolation from blood, liver and skin samples followed a standard phenol-chloroform extraction protocol (Sambrook et al. 1989). DNA was extracted from faecal samples using protocols based on the GuSCN/silica method (Boom et al. 1990) as previously described in Roques et al. (2014). All scat samples collected in the wild were first screened for species identification using species-specific primers (Roques et al. 2011). Those samples belonging to jaguars were genotyped at a set of 11 microsatellite loci as described in Roques et al. (2014). Briefly, after scoring the alleles with GENE-MAPPER version 4.0 (Applied Biosystems), a unique consensus genotype was assigned to samples given a consensus criterion derived from that proposed by Taberlet et al. (1996) and based on the results of the four PCR replicates. The four genotype replicates were compared to the consensus genotype and the quality index value (QI) was calculated as described by Miquel et al. (2006). Full details on error rates, allelic dropout and false alleles are



available in a previous paper (see Supplementary Material 1 in Roques et al. 2014).

Population structure, size and gene flow

To explore the genetic evidence for subdivision among jaguars, we first used the program STRUCTURE over the 14 locations and to identify populations within Brazil (BRAZ) or within Mexico (MEXC). Simulations were conducted by varying the number of genetic clusters (k = 1-12; alternatively, k = 1-7 for within BRAZ and MEXC) with 30,000 steps of the Markov chain Monte Carlo (MCMC), following a burn-in period of 300,000 iterations, with and without a priori 'population' information. Twenty independent runs for each k were performed under an admixture model with correlated gene frequencies to determine the number of genetic clusters. The most likely number of k was calculated based on Δk as described in Evanno et al. (2005) and on visual inspection of the plot of lnP (D) as a function of k, using STRUCTURE HARVESTER (Earl and vonHoldt 2011). Once the number of k was estimated, two replicates of a longer run with 300,000 steps of burn-in followed by 1,000,000 steps were performed to assign individuals to clusters. The partition of the total genetic variation into different genetic clusters was further assessed based on a factorial component analysis (FCA) in GENETIX v.4.03 (Belkhir et al. 2004). The extent of genetic differentiation among the populations defined based on clustering approaches (see above) was estimated with FST statistics (Weir and Cockerham 1984) using Genetix (5000 permutations). Further, we tested whether patterns of neutral genetic structure were the product of isolation by distance. We calculated population-level pairwise genetic differentiation as FST/ (1 - FST) (Slatkin 1995) using Fst values calculated in Genetix (Belkhir et al. 2004). Geographic distance was calculated as the closest linear distance between pairs of sampling areas using Google Earth (http://earth.google.com). We tested whether genetic distance was related to geographic distance using Mantel tests, implemented in the program IBD (Isolation by Distance, Bohonak 2002).

Detection of migrants

STRUCTURE 2.3.2 and GENECLASS 2.0 were also used to identify first-generation migrants and individuals with mixed ancestry among the pre-defined populations. In STRUCTURE, prior population information was used in the USEPOPINFO option into determine the individuals that were not residents of their sampled population. MIGPRIOR was set to 0.05. GENECLASS 2.0 specifically identifies first generation migrants, i.e. individuals born in a population different to the one it was sampled (Paetkau et al. 2004; Piry et al. 2004). The Bayesian criterion of Rannala and Mountain

in combination with the resampling method of Paetkau and an alpha level of 0.01 were used to determine critical values.

Genetic diversity

Diversity parameters were first calculated for the pre-defined populations. Departures from linkage disequilibrium and the Hardy–Weinberg equilibrium (HWE) were tested using exact tests as implemented in GENEPOP on the web (Rousset 2008). Genetic diversity was assessed through the observed and expected heterozygosity (HO and HE) estimated using GENETIX. Further, allelic richness (i.e., the number of alleles per locus independent of sample size) and percentage of shared and private alleles were calculated using the program HPrare (Kalinowski 2005). Differences of indices among populations were tested with Wilcoxon signed-rank tests.

Population size reductions

We used two different approaches to test for a genetic bottleneck signature. Because violations of the panmixia assumption might bias these tests, genetic homogeneity within the pre-defined population units was confirmed based on both FST statistical significance (see Supplementary Material S1) and Structure approaches (see above). For the first approach, the mutation-drift equilibrium test which is implemented in BOTTLENECK 1.2.02 (Cornuet and Luikart 1996, Piry et al. 1999), tests whether the number of loci with heterozygosity excess is significantly higher than that expected by chance at mutation-drift equilibrium. In populations that have experienced a relatively recent (within the last ~ 0.2 -4Ne generations) reduction in effective size, the number of alleles is reduced faster than gene diversity, leading to a transient excess of heterozygosity (Luikart and Cornuet 1998). The program was initially run under either the 100 % infinite alleles model (IAM) or stepwise mutation model (SMM) of microsatellites evolution. In order to test the sensitivity of the analysis to the mutation model chosen, we ran the program under a two-phase mutation model (TPM model) because the microsatellites in this study are dinucleotide repeats, which better fit the IAM (Cornuet and Luikart 1996). We ran the program with proportions of either 5 or 30 % of SMM. Significance was assessed from 10,000 iterations using a Wilcoxon signed-rank test which gives the highest statistical power when population sample size is small (30 or fewer) (Cornuet and Luikart 1996). For the second approach, we used the M-ratio (Garza and Williamson 2001) which corresponds to the mean ratio of the number of alleles to the allele size range across all loci, and the value is expected to decrease following a population reduction. The M-ratio test is more sensitive than the other two tests and would detect a bottleneck signal longer after it occurred, and thus gives insights into population contractions occurring at a larger



timescale. M-ratios were calculated using AGARST (Harley 2002) and the critical M-ratio (Mcrit) for each sample location was determined using the critical_M.exe software (Garza and Williamson 2001). We set the mean number of non-one-stepwise mutations (ps) to 0.12 and the mean size of larger mutation (thetaS) as 2.8 as conservative parameters (i.e., lower critical value), as suggested by the authors. Pre-bottleneck values were calculated using $\alpha = 5 \times 10^{-4}$ (Garza and Williamson 2001) and Ne values estimated in this study for the jaguar, as well as several Ne values (i.e., 20, 50, 150, 300). Two loci with odd-sized alleles (those that did not represent multiples of the recognized repeat unit) were omitted from these analyses (FC115 and FC566).

To estimate the effective size (Ne) in our populations, we first applied the linkage disequilibrium method using the program LDNE (Waples and Do 2008), assuming random mating and excluding all alleles with frequencies lower than 0.02. We also used an approximate Bayesian computation (ABC) approach as implemented in the program ONESAMP (Tallmon et al. 2008), which is considered more robust and less biased by substructure and overlapping generations than LDNE (Luikart et al. 2010).

In order to test the genetic effects of recent habitat degradation in the southeastern Brazilian areas and especially the probable recent isolation of the Caatinga population, we used a coalescent-based MCMC simulation implemented in 2MOD (Ciofi et al. 1999). This method tests whether the observed population structure would better fit a gene flowdrift equilibrium model or a pure drift model; the first model assumes a balance between gene flow and drift (i.e., populations at equilibrium) while the second model assumes that an ancestral panmictic population has evolved into several different units diverging by drift in the absence of gene flow. The MCMC search was carried out twice for 30×10^5 iterations with the first 3×10^4 discarded as burn-in. The posterior distribution of F (probability of co-ancestry of any two genes in the putative population) was estimated for each population. Simulations were run with 600,000 steps with a burn-in of 100,000 in three independent runs. We used Tracer v 1.40 (http://beast.bio.ed.ac.uk/) to evaluate the stationarity of model parameters, verify adequate sample sizes, determine an appropriate amount of burn-in, and verify the consistency between runs. Under the drift model, we estimated the time since isolation among the three areas relative to the population size, (T/N) as $-\log(1 - F)$, following Ciofi et al. (1999).

Results

Non-invasive genetics

We successfully determined the species for 73% (N = 473) of 651 faecal samples collected and processed



Table 2 F_{st} (left) indices of genetic differentiation among defined jaguar populations for Mexico (MEXC); Caatinga (CAPV); Amazon (AMZN); Pantanal (PANT); and Central areas (CENTR)

	MEXC	CAPV	AMZN	PANT	CENTR
MEXC	_				
CAPV	0.190	_			
AMZN	0.135	0.115	-		
PANT	0.162	0.168	0.087	_	
CENTR	0.107	0.067	0.026*	0.067	

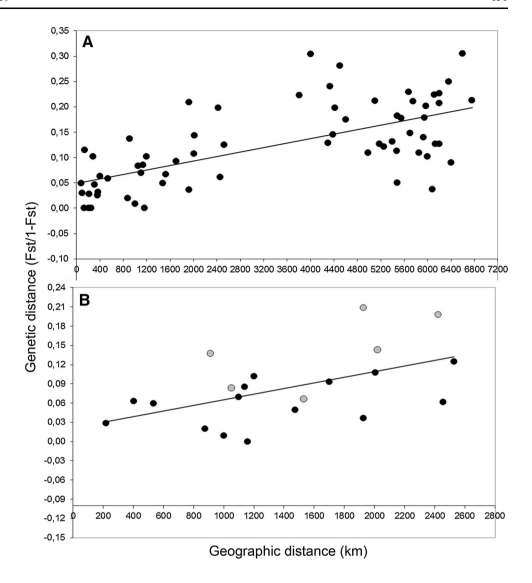
All values are highly significant ($P \le 0.01$) except * ($P \ge 0.05$)

(Table 1). Most of the faecal samples were from jaguars (49.7 %) and pumas (41.6 %), and to a lesser extent, smaller felids (ocelot/margay; 8.7 %). Among the 234 jaguar faecal samples, a high proportion (91 %) have \geq 50 %, quality (based on the Quality Index; QI; Miquel et al. 2006) and 71 % of genotypes have even higher quality (QI \geq 75 %). Consensus multilocus genotypes for each sample were grouped into 62 different genotypes representing distinct individuals following the assignment strategy described by Roques et al. (2014). Including the genotypes obtained from high quality DNA sources (blood: n = 31; liver: n = 13, and skin: n = 7) we gathered 102 distinct genotypes from 14 study areas across the current distribution range of the jaguars (Table 1 and Supplementary Material S3).

Genetic differentiation and connectivity

The overall genetic differentiation was high and jaguar populations were genetically structured throughout the species' range. Within Brazil, Fst values were low and not significantly different from zero among the four Amazonian localities (DUCK, UATM, VIRU, MARA) and among all central areas along the Araguaia river (CANT, ARAG, PNEM), but they were high and significant among the other populations studied (see Supplementary Material S1). Based on these results, we defined four differentiated genetic entities within Brazil (Table 1): AMZN (Amazon-DUCK, UATM, MARC, and VIRU); PANT (Pantanal); CAPV (Caatinga); and an intermediate area in the central region, namely CENTR (ARAG, CANT, and PNEM). Within the Yucatan Peninsula, estimates of genetic differentiation (Fst) were low and not significant for any pairwise comparison, thus corroborating genetic homogeneity at this scale. Differentiation among the inferred genetic units was very high and significant for comparisons between Brazil (PANT, CAPV, AMZN, CENT) and MEXC (Table 2; $P \le 0.01$), indicating high divergence in allele frequencies between these geographically distant areas. Within Brazil, the highest value

Fig. 2 Isolation by distance across jaguar populations. Pairwise genetic differentiation as FST/(1 – FST) at **a** multiregional scale including Mexico (n = 15 populations) and **b** regional scale; Brazil (n = 9 populations). In *grey*, genetic comparisons involving CAPV, the easternmost Brazilian sampling site



occurred with comparisons involving CAPV and the other sampling areas, while differentiation between CENT and the rest of the populations was lower and the differentiation between AMZN and CENT was not significant (Table 2).

A significant positive correlation between genetic and geographic distance was observed among the jaguar populations at both large (Fig. 2a; Mantel test, r=0.655, P<0.001) and regional (Fig. 2b; Mantel test, r=0.5232, P<0.019) scales. The result of this test showed that a considerable part of the genetic variation was explained by geographic distance. Within Brazil, these results supported the Factorial Component Analysis (Fig. 3b) since all geographically-close populations resembled each other. Also, we found that almost all comparisons involving CAPV (Fig. 2b, grey circles) stand above the line, corroborating that this area presents more differentiation with the other areas than expected by distance only.

The STRUCTURE analysis including all samples suggested K = 4 as the most likely number of genetic clusters (Fig. 3a and Supplementary Material S4 for Evanno's output table for all K values). The geographical samples with predominant membership in the four clusters were grouped into Mexico (MEXC: ZAPT, EDEN, CAOB, CALAK, mean Q = 0.66); Amazon (AMZN: MARA, VIRU, DUCK, UATM, mean Q = 0.84), Caatinga (CAPV mean Q = 0.71) and Pantanal (PANT mean Q = 0.72). When the Mexican areas were analyzed separately, a single and panmictic population (MEXC, K = 1) (results not shown) was the most likely scenario. Within Brazil, K = 3was the most likely number of genetic clusters. These three clusters correspond to the three distinct geographical areas of PANT, AMZN and CAPV. The individuals from the central localities CENT, namely CANT, ARAG, PNEM, cluster with individuals from AMZN, but show some ancestry in the other two populations (Fig. 3a).



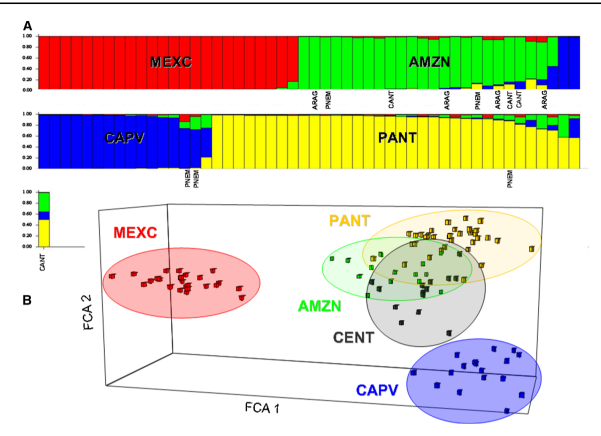


Fig. 3 a The genetic structure of the Brazilian populations identified by the STRUCTURE analysis assuming four genetic clusters (K=4; MEXC, AMAZ, PANT and CAPV) in the overall population. Individuals are represented as bars partitioned into segments corresponding to their membership in genetic clusters indicated by the colors. Individuals from the Central areas (CENT: ARAG, PNEM, CANT) show from 50 to 100 % ancestry in AMAZ, and the

remainder corresponding to the other two clusters **b** Three-dimensional Factorial Component Analysis graph. Names are referred to sampling sites (see Table 1). Jaguars from the central Brazilian areas (CENT) are intermediate between three differentiated groups (PANT, CAPV and AMZN). MEXC are genetically highly differentiated from the remaining samples

The representation of all individuals in the Factorial Correspondence Analysis was also highly congruent with the above clustering, clearly depicting the divergence of Mexican areas and the existence of three genetic entities in Brazil (CAPV, PANT, AMZN) and with CENT individuals occupying intermediate positions between these (Fig. 3b). The analyses clearly illustrated that CAPV is highly differentiated from the rest of populations and that jaguars from the central admixed area are genetically intermediate between those from AMZN and those from southern (PANT) and eastern (CAPV) populations.

Identification of migrants and admixed individuals within Brazil

STRUCTURE identifies a total of 17 migrants in Brazil (Table 3), most of them (n = 14) sampled in central areas (CENT), while two in PANT and one in AMZN. Assignments in GENECLASS were significant for nine migrants. Both programs were concordant in detecting four first-

generation migrants (i.e. not born in the sampled area), all from CENTR (n=2 in ARAG and two in PNEM). For the four migrants only detected by GENECLASS, the LOG(L_home)/(L_Max) values were relatively low and may be taken with caution. STRUCTURE also identified two individuals (CANT_H3-28 and PANT_SGH27) that were neither readily classified as migrants nor as residents (Q-values <0.60) suggesting that they might be of admixed ancestry (PANT/AMZN; Table 3).

Genetic diversity and population demography

None of the populations showed significant HWE disequilibrium after Bonferroni correction ($P \le 0.001$). Also, only two out of 55 tests for Linkage disequilibrium LD were statistically significant after applying the Bonferroni correction. Those tests involved different pairs of loci and occurred in different populations, suggesting that the assayed loci assorted independently. Mean expected and observed heterozygosities across loci and samples were



Table 3 Identification of migrant performed with STRUCTURE and GENECLASS

Sample name	Sampling location	STRUCT	URE Q $(k = 1)$	GENECLASS		
		No prior i	information, k			
		PANT CAPV		AMZN	LOG	ORIGIN
PANSGM11	PANT	0.224	0.006	0.770		
PANSGH27	PANT	0.420	0.010	0.570		
CANTH1-5	CENT	0.013	0.036	0.950		
CANTH2-6	CENT	0.100	0.047	0.853		
CANTM1-13	CENT	0.007	0.202	0.791		
ARAM1	CENT	0.009	0.063	0.928		
ARAM2	CENT	0.188	0.031	0.782		
PNEM2	CENT	0.110	0.010	0.880		
PNEHSG18	CENT	0.018	0.018	0.964		
CANTH3-28	CENT	0.446	0.145	0.409		
PNEM1	CENT	0.012	0.013	0.974		
PNEM3	CENT	0.029	0.659	0.312		
PNEHSG29 ^a	CENT	0.028	0.681	0.291	1.433	CAPV
PNEMSG15 ^a	CENT	0.830	0.013	0.158	2.772	PANT
ARAH3 ^a	CENT	0.012	0.009	0.979	0.789	AMZN
ARAH-M4 ^a	CENT	0.135	0.172	0.693	1.124	PANT
DUCM2	AMZN	0.018	0.079	0.903	1.865	CENT
DUCM4	AMZN				0.545	CENT
UATH1	AMZN				0.248	CENT
UATM1	AMZN				0.106	CENT
CAPM4	CAPV				0.205	CENT

 $LOG = LOG(L_home)/(L_Max)$ significant for probability below 0.01

ORIGIN means the group to which migrants are reclassified

Jaguars marked with awere identified as migrants from the same area with both methods

0.800 and 0.730, respectively. Both heterozygosity and allele number were higher in Brazil (mean He = 0.812, mean A = 9.45) than in Mexico (mean He = 0.634, A = 4.45) (Table 4). Expected heterozygosity, He, calculated for the genetic clusters identified above, ranged from 0.654 to 0.805, with values significantly higher in AMZN (Wilcoxon sign-rank test, $P \le 0.03$) and lower in MEXC $(P \le 0.03)$ than in the other areas. However, the difference between MEXC and CAPV was not significant (P = 0.22). Allelic richness was also highest for AMZN (P < 0.02) and lowest for MEXC and CAPV (Table 4). The allelic richness in PANT was moderate and not significantly different from the values found in MEXC (P = 0.09) and CAPV (P = 0.22). The jaguar population at CAPV had the lowest proportion of private alleles (4 %) in Brazil, less than half of that found for AMZN, and the population at CENT shared the highest proportion of alleles with the other studied populations (74, 69 and 63 % for AMZN, PANT, and CAPV, respectively). While the highest effective population size was estimated for AMZN (>250), the effective population sizes were much lower for the remaining populations (between 13 and 30) (Table 4).

When we applied BOTTLENECK, we observed clear signatures of recent bottlenecks for both MEX and CAPV under IAM (P < 0.05) and TPM with either SMM = 5 or 70 % (see Table 4). However, all tests were non-significant under SMM. Among all populations sampled, the M-ratio ranged from 0.670 (CI = 0.057) to 0.888 (CI = 0.041), with the lowest values found in CAPV and PANT (Table 4). However, only the value for CAPV was lower than almost the whole range of simulated critical values (Mcrit20 = 0.662, Mcrit50 = 0.650, Mcrit150 = 0.629,and Mcrit300 = 0.600), suggesting a stronger reduction in size of this population than in the other populations. In contrast, the M-ratio of MEXC was high (0.888) and contrasts with the highly significant P value when BOT-TLENECK was applied; these values suggest a more recent population contraction event in this region (Cornuet and Luikart 1996).

Using the 2Mod program, we evaluated alternative hypotheses of whether the isolation of the Caatinga population was the result of a recent isolation (i.e., the pure drift model) or if this reflected an equilibrium situation of an historically small and weakly connected population (i.e., the gene flow-drift



Table 4 Summary of genetic indices of defined populations for Mexico (MEXC); Caatinga (CAPV); A mazon (AMZN); Pantanal (PANT); and central areas (CENT)

Genetic indices	Parameters/methods	MEXC	AMZN	PANT	CAPV	CENT
	N	24	12	34	18	14
Diversity	HE	0.654 + 0.147	0.805 + 0.084	0.726 + 0.097	0.709 + 0.133	0.837 + 0.0490
	НО	0.684 + 0.135	0.848 + 0.099	0.734 + 0.161	0.779 + 0.148	0.758 + 0.1692
	AR	5.10	6.73	5.61	5.20	7.26
Effective pop. size (Ne)	Onesamp	30 (22–38)	298 (na)	14 (10–17)	14 (12–16)	na
	LDNe	25 (14–45)	na (21-inf)	17 (10–28)	13 (7–28)	na
Bottleneck	Wilcoxon test					
	P (SMM 5 %)	0.0005^{S}	0.0615^{NS}	0.0508^{NS}	0.0268 ^S	na
	P (SMM 70 %)	0.0100^{8}	0.1302^{NS}	0.4410^{NS}	0.0500^{S}	na
	AF distribution	L -shaped NS	L -shaped NS	L -shaped NS	L -shaped NS	na
	M ratio	$0.888 + 0.041^{NS}$	$0.752 + 0.029^{NS}$	$0.717 + 0.041^{NS}$	$0.670 + 0.057^{NS}$	na

Values are provided for number of jaguars (N), expected (HE) and observed (HO) heterozygosities, and allelic richness (AR), P values are noted as statistically significant ($P \le 0.001$) (S) and non significant, NS na signifies no applicable. Details of the methods are provided in the "Materials and methods" section

equilibrium model). The results of 2Mod overwhelmingly supported a pure-drift rather than a migration-drift equilibrium scenario [P (drift model) = 0.9] for the CAPV, AMZN, CENT populations. Under the drift model, we calculated F values (FCAPV = 0.1481, 95 % CI 0.1361–0.1494; FAMZN = 0.0741, 95 % CI 0.0.0737–0.0746; FCENT = 0.0531, 95 % CI 0.0.0536–0.0541) and the T/N was estimated to be 0.1602 (2Ne = 28) for CAPV; 0.0544 (2Ne = 400) for CENT; and 0.0768 (2Ne = 596) for AMZN. Based on a generation time of 5 years and the effective population size estimates (reported here), these values suggest the population in CAPV has been isolated for approximately 20 years.

Discussion

Genetic effects of habitat deterioration and biogeography

Our study examined genetic diversity and connectivity of jaguars on a large spatial scale in Mexican and Brazilian ecosystems. The results indicate that despite prior evidence for historical connectivity and panmixia (Eizirik et al. 2001), the jaguar is genetically structured throughout its range. While genetic differentiation of areas of the jaguar distribution range is primarily driven by isolation resulting from distance (Fig. 2a) and putative barriers to gene flow (e.g., Amazon River, Darien Straits; Eizirik et al. 2001), the recent habitat deterioration (i.e., habitat fragmentation and loss) may have caused a disruption of gene flow and an intensification of genetic drift in part of its range. The population of Capivara in the eastern edge of the species

distribution is separated by a large area of unsuitable habitat, suggesting that such barrier may further contribute to genetic divergence and to the pronounced genetic isolation found in this area.

Our results are similar to those reported by Eizirik et al. (2001) for the same area and show that the genetic diversity values in Mexico are some of the lowest reported for the species (Table 5). The low diversity and high differentiation for this particular region may be attributable to the recent colonization of jaguar populations in the northern areas and to a global pattern of isolation by distance (Eizirik et al. 2001). However, the significant signs of recent bottlenecks found in this region suggest that individuals from the Mexican population might be exhibiting the genetic signals of recent anthropogenic perturbations and isolation. This area is situated close to the northern limit of the species' range and is probably more vulnerable to stochastic demographic effects (Vucetich and Waite 2003; Chávez et al. 2005). Additionally, the Yucatan Peninsula population is connected northward to areas with groups of individuals that occur at the lowest densities reported for jaguars, including the relict populations of Sinaloa and Baja California (Navarro-Serment et al. 2005; Rosas-Rosas and Bender 2012) (see Fig. 1a). Jaguars have been extirpated to the south of the Yucatan, in parts of Nicaragua and Honduras, and this loss may have disrupted the gene flow with individuals from further south (Sanderson et al. 2002).

Genetic evidences for the effects of recent isolation were compelling for the Caatinga (CAPV) population. All population structure analyses indicated increased genetic drift and reduced gene flow between CAPV and the other



Table 5 Genetic surveys based on microsatellites markers that estimate the diversity of jaguar populations at different geographic scales

•		,	3 0	1 1			0 0 1
Study sites *	Geographic scale	N ^S	N^L	N ^{SH}	N ^A	HE	References
Mexico (Yucatan peninsula)	Regional	24	11	_	5.10	0.654	This study
Central America (Mexico, Guatemala, Panama, Costa Rica, Nicaragua)	Multi regional	16	29	5	5.20	0.622	Eizirik et al. (2001)
North -South America (Mexico-CA-Venezuela, French Guyana)	MultiRegional	25	29	5	6.80	0.695	Ruiz-García et al. (2001)
Guatemala-Paraguay	MultiRegional	107	12	4	11.00	0.846	Ruiz-Garcia (2007)
Colombia	Regional	62	12	4	10.00	0.835	Ruiz-Garcia (2006)
PERU	Regional	na	12	4	7.00	0.860	Ruiz-Garcia (2007)
BOLIVIA	Regional	na	12	4	7.00	0.860	Ruiz-Garcia (2007)
Brazil		59	11				
Amazon	Regional	18	11	_	6.90	0.805	This study
Cerrado	Regional	12	11	_	7.45	0.802	This study
Pantanal	Regional	34	11	_	7.00	0.726	This study
Caatinga	Regional	17	11	_	5.55	0.709	This study
North Argentina/South Brazil Atlantic Forest (Upper Parana)	Regional	13	13	0	6.00	0.737	Haag et al. (2010)
South -South America (Brazil, Bolivia, Paraguay)	MultiRegional	17	29	5	6.70	0.724	Eizirik et al. (2001)
Mexico-Brazil	Distribution range	42	29	5	8.30	0.739	Eizirik et al. (2001)
	Distribution range	102	11	_	10.55	0.800	This study

Study sites are ordened from north to south of the jaguar distribution range (See also Fig. 1a) Number of samples (N^S), loci (N^L), shared loci with this study (N^{SH}), alleles (N^A), and expected (HE) heterozygosity. na indicates not applicable. See Supplementary Material S2 for additional information on studied areas (codes, biomes, country, distances between sites, etc.)

regions. A significant reduction of diversity is reflected in low values of allelic richness (Table 4), whereas both estimates of heterozygosity were close to those estimated previously for the species (He = 0.732 in Haag et al. 2010 and He = 0.724 in Eizirik et al. 2001), but lower than those in the Amazonian strongholds (Table 5). This difference may be a reflection of the generally faster response of allelic richness to population contractions than heterozygosity (Cornuet and Luikart 1996; Srikwan and Woodruff 2000), with the former being thus a more sensitive signal of recent genetic erosion in isolated populations. The preponderance of genetic drift and the increased isolation of the CAPV population in recent times are also supported by the selection of a pure-drift model by the coalescent-based simulations. The Bayesian approach suggests a very recent (about 20 years) genetic isolation of the CAPV population, while jaguars from the Amazon and Cerrado regions probably were well connected until 100 years ago. This observation, along with the low proportion of private alleles in CAPV and the fact that it shares a major proportion of its alleles with the central areas, corroborates historical evidence that CAPV was once part of a much larger population that included the Cerrado.

The detection of only two migrants from PNEM (assigned to CAPV) is thus consistent with restricted connectivity and disturbed potential corridors recently described in this area (Silveira et al. 2014 and Fig. 1b). The Cerrado biome, which

marks the transition between the Amazon and the southern populations, has been intensively modified since the 1950s through extensive cattle farming and agricultural monocultures (rice, corn, soybean), and today up to 80 % of this region is considered degraded (Cavalcanti and Joly 2002). The isolation of the jaguar population in the Caatinga may have been driven in the last few decades by the lack of suitable habitat for connectivity with surrounding populations. The relatively low estimate of effective population size calculated for CAPV is supported by results of recent field studies in the region. While the Capivara National Park is considered to have an important jaguar population (Silveira et al. 2010), substantial contractions as the result of habitat changes, scarcity of prey and persecution have been reported recently in the Brazilian Caatinga (Sollmann et al. 2008). The semiarid climate and poor soil limit large scale agriculture and cattle ranching, and about 60 % of this area still maintains the native vegetation cover, although as fragmented blocks (Casteleti et al. 2005). The low estimated effective population size suggests that further genetic erosion will occur until the population size or the gene flow from other regions increases (Frankham et al. 1999; England et al. 2010; Palomares et al. 2012).

Jaguar populations in other Brazilian areas (AMZN, CENT, PANT) were generally more diverse than the ones at the northern and eastern limits of the species range (MEXC, CAPV). The Amazon was the most genetically



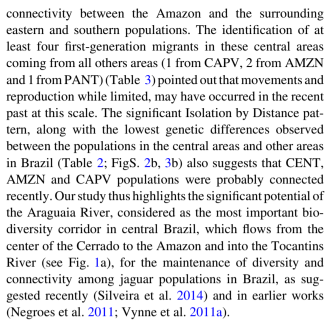
diverse region and had the highest proportion of private alleles, and variability indices were comparable to values found in other tracts of forest in Colombia, Bolivia, and Peru (Table 5). Many areas in the Amazon are still connected, forming enormous blocks of evergreen forest that support large effective populations (de Oliveira et al. 2012) and panmictic breeding, and our estimate of a moderate to large effective population size agrees with that reported in this biome (Sollmann et al. 2008).

Results for the Pantanal region indicate that even though population bottlenecks were not statistically detectable, this area may be showing early signs of genetic erosion and isolation. Allelic richness and heterozygosity in the population from the Caiman Ecological Reserve were medium to low (Table 4) and close to those found in the nearby area of the Upper Parana (Haag et al. 2010; Table 5). These results were striking for several reasons: as the largest seasonally flooded landlocked area in the world, the Brazilian Pantanal still is covered by native vegetation over most of its territory and relatively well-connected; the extensive cattle ranching on native pastures (Harris et al. 2005) has maintained some level of habitat quality for jaguars and has provided them with additional sources of prey (Swartz 2000), what may explain the reported high jaguar density (Soisalo and Cavalcanti 2006), even in non-protected areas. However, in some areas of this biome, the genetic patterns we detected in our research support the observations made in earlier work (Altrichter et al. 2006), namely a decrease in the size of some populations and increased isolation. These results are not unexpected because some intensive cattle ranching practices have resulted in a major loss of native habitat and increased direct persecution (i.e., hunting) of jaguars resulting from the increased conflict with cattle ranchers (Crawshaw and Quigley 2002). Additionally, populations in the southern Pantanal are connected southwards with the Atlantic forest region, a heavily human-impacted biome where jaguar populations also show clear signs of genetic isolation and loss of genetic diversity (Haag et al. 2010). The results of our work can serve as a starting place for discussion and evaluation of the role of the Pantanal as a secure refuge for jaguars.

The importance of connectivity for jaguar conservation

The population structure observed at this scale intimate that connectivity with the extreme eastern (i.e., Caatinga) and southern areas (i.e., Pantanal) is limited (Table 2) and that much of the existing connectivity may be at risk because of continued habitat erosion, and might be enhanced through habitat restoration or genetic exchange among them.

Interestingly, our research suggests that the central areas of Brazil within the Cerrado region (PNEM, ARA and CANT) (Fig. 1a), may act as "stepping stones" to maintain



The restoration of ecological connectivity between populations over relatively large scales should be one of the main priorities for the conservation of the jaguar and for other wide-ranging species with high dispersal, low population density and that are particularly vulnerable to anthropogenic impacts. We stress the importance of ambitious programs to conserve a continuous north to south habitat corridor through the range of the species (Rabinowitz and Zeller 2010; Fig. 1b) and to evaluate the potential for large scale jaguar corridors in Brazil (Silveira et al. 2014).

Implications for species viability, conservation and management

Our work showed that genetic patterns differed among jaguar populations and biomes but were highly consistent with the known status of the populations as well as with the degree of habitat deterioration and connectivity with neighboring populations. Large continuous forested areas, such as the Amazon, still maintain genetically healthy jaguar populations. In contrast, the geographic and genetic isolation of the Caatinga population suggests that the jaguar may be at risk of extinction in those areas of its range not connected, and especially those near the edge, or those which may become isolated in the near future by the high rates of fragmentation. With the exception of the groups in the Amazon, estimates of effective population sizes were low (N = 13-30) and much below the number of 85 individuals proposed as the minimum threshold for long-term population viability (>200 years; Sollmann et al. 2008). These low population values reinforce other evidence showing a continued trend of declining jaguar populations. While large carnivores with widespread geographic ranges should be at lower risk from habitat



fragmentation, our research showed that jaguar connectivity may be limited by the difficulty of dispersing in modified habitats. In a changing landscape, protection and/ or establishment of reserves are one of the most important tools for habitat preservation as a buffer against anthropogenic impacts (Noss et al. 1996; Margules and Pressey 2000; Rylands and Brandon 2005; Shivik 2006). In Brazil, a system of connected protected areas extensive enough to hold long-term viable jaguar populations is currently implemented in the Amazon, but it is absent in other important jaguar areas such as the Caatinga biome. Long-term jaguar conservation may depend on alternative strategies integrating non-protected landscapes, as well as cultural and political mechanisms (Sollmann et al. 2008).

Acknowledgments This study was carried out with the support of the project BIOCON 05—100/06 of the Fundación BBVA, the project CGL2010-16902 of the Spanish Ministry of Research and Innovation, the project CGL2013-46026-P of MINECO, the excellence project RNM 2300 of the Junta de Andalucía, and projects UAM-PTC-333 and PROMEP/103.5/12/3823. Sampling in the Mexican areas under the license SGPA/DGVS/549 provided by Martín Vargas of the Dirección General de Vida Silvestre (Semarnat). Faecal samples were exported from Mexico to Spain under the export licences no MX33790 and MX42916 of the Secertaria de Medio Ambiente/ CITES. Sampling in Brazil was carried out in RAPELD sites installed or maintained by the Brazilian Program for Biodiversity Research (PPBio) and under licenses #131/2005 CGFAU/LIC, 13883-1 SISBIO and 15664-1 SISBIO of the Instituto Brasileiro do Meio Ambiente-IBAMA. Faecal samples were exported from Brazil to Spain for genetic analysis under IBAMA/CGEN Autorização de Acesso license #063/05 and IBAMA/CITES export licenses #0123242BR and 08BR002056/DF". We thank the management of the Edén Ecological Reserve (Marco Lazcano) and El Zapotal Ecological Reserve (Pronatura Península de Yucatán: Juan Carlos Faller and María Andrade) for their logistical support. We are grateful to J.S. Lópes and J. Tavares for the collection of most of the field samples in Brazil. Julia Martínez, Gloria Clemencia Amaya, Juan Carlos Faller, Meredic Calleja and Ana Alicia Morales helped with the fieldwork in Brazil and Mexico, as well as the local reserve staff of El Zapotal and El Edén (Mexico). L. Soriano and A. Piriz provided technical advice on multiple issues, and A. García, E. Marmesat, and B. Gutiérrez assisted in the analysis of samples. Logistical support was provided by Laboratorio de Ecología Molecular, Estación Biológica de Doñana, CSIC (LEM-EBD). The Spanish Ministry of Education and Sciences supported the visit of S. Roques in Mexico. We thank Manuela Gonzalez-Suarez and Philip Hedrick for an early revision of the manuscript.

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