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Tools and Technology

A Noninvasive Method to Detect Mexican Wolves and Estimate Abundance

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ABSTRACT Monitoring wolf abundance is important for recovery efforts of Mexican wolves (*Canis lupus baileyi*) in the Blue Range Wolf Recovery Area in Arizona and New Mexico, USA. Although radiotelemetry has been a reliable method, collaring and tracking wolves in an expanding population will be prohibitively expensive and alternative methods to estimate abundance will become necessary. We applied 10 canid microsatellite loci to 235 Mexican wolf samples, 48 coyote (*C. latrans*) samples, and 14 domestic dog (*C. lupus familiaris*) samples to identify alleles that provide reliable separation of these species. We then evaluated an approach for prescreening, noninvasively collected DNA obtained from fecal samples to identify Mexican wolves. We generated complete genotypes for only those samples identified as probable Mexican wolves. We used these genotypes to estimate mark–recapture population estimates of Mexican wolves and compared these to known numbers of wolves in the study area. We collected fecal samples during 3 sampling periods in 2007–2008 and used Huggins-type mark–recapture models to estimate Mexican wolf abundance. We were able to generate abundance estimates with 95% confidence for 2 of 3 sampling periods. We estimated abundance to be 10 (95% CI = 6–34) during one sampling period when the known abundance was 10 and we estimated abundance to be 9 (95% CI = 6–30) during the other sampling period when the known abundance was 10. The application of this noninvasive method to estimate Mexican wolf abundance provides an alternative monitoring tool that could be useful for long-term monitoring of this and other recovering populations. Published 2016. This article is a U.S. Government work and is in the public domain in the USA.

KEY WORDS alleles, Arizona, Blue Range Wolf Recovery Area, *Canis lupus baileyi*, fecal noninvasive DNA, mark–recapture, Mexican wolf, microsatellites.

The Mexican gray wolf (*Canis lupus baileyi*) is the most genetically distinct (Wayne et al. 1992, Garcia-Moreno et al. 1996, Chambers et al. 2012), smallest (Bogan and Mehlhop 1983), and most endangered (Ginsberg and Macdonald 1990) subspecies of gray wolf in North America. Historically, its range extended across much of Mexico; and southern New Mexico, the Trans-Pecos of

Texas, and southeastern and central Arizona in the United States (Young and Goldman 1944, Brown 1983). Sustained efforts to eradicate wolves and habitat alterations led to its extirpation from the wild in the United States and Mexico by the 1980s (Brown 1983, United States Department of the Interior, Fish and Wildlife Service [USFWS] 1996, Hedrick and Fredrickson 2010).

In 1976, under provisions of the Endangered Species Act, the Mexican wolf was listed as endangered (Parsons 1998). Subsequently, in 1998, the USFWS initiated releases of captive-bred Mexican wolves into the Blue Range Wolf Recovery Area (BRWRA) that included all of the Apache National Forest in southeastern Arizona and the Gila National Forest in southwestern New Mexico (Parsons 1998,

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USFWS 1998). Since then, individuals and family groups of Mexican wolves have been intermittently released or translocated into this portion of their historical range and many have begun to breed in the wild (Mexican Wolf Blue Range Adaptive Management Oversight Committee and Interagency Field Team [AMOC] 2005, USFWS 2013).

Monitoring the population status of Mexican wolves in the United States is a critical component of the recovery process and will likely remain an important priority for the foreseeable future (U.S. Department of the Interior, Fish and Wildlife Service 2013). A variety of techniques are used to monitor wolf populations, including howling and track surveys (Harrington and Mech 1982, Crête and Messier 1987, Fuller and Sampson 1988, Ballard et al. 1995), radiotelemetry (Ballard et al. 1987), and visual observations; each of these has their unique benefits and challenges. Radiotelemetry, observation, and track and howling surveys have been the primary tools used for Mexican wolf monitoring (USFWS 2013). These methods have been satisfactory for monitoring the Mexican wolf population in the BRWRA to date. However, as the population grows, the time-consuming nature of radiotelemetry, invasiveness of capture and handling required for collaring, and the prohibitive expense of tracking more wolves will limit the usefulness of these approaches. Therefore, alternative methods to monitor Mexican wolf populations are needed.

Contemporary molecular approaches have made it possible for biologists to identify individuals by obtaining and amplifying DNA from noninvasively collected samples

(Taberlet et al. 1999, Schwartz et al. 2007) including feces, hair, or saliva (Woods et al. 1999, Lucchini et al. 2002, Waits and Paetkau 2005, Bohling and Waits 2011, Nichols et al. 2012). Noninvasive genotyping of individuals from fecal DNA has been used successfully to generate mark–recapture population estimates of various North American wild canids, including gray wolves, in recovery areas (Creel et al. 2003, Marucco et al. 2012, Stansbury et al. 2014), endangered red wolves (*Canis rufus*; Knapp et al. 2009), and coyotes (*C. latrans*; Kohn et al. 1999, Prugh et al. 2005). A potential limitation of effectively using noninvasive DNA data in a mark–recapture statistical framework for the endangered Mexican wolf is their known low genetic diversity; only 7 captive-bred individuals were used as founders of the current wild population (Hedrick and Fredrickson 2010). Therefore, finding suitable markers to reliably distinguish individuals in the BRWRA population and distinguishing Mexican wolves from dogs and coyotes is critical for successful noninvasive, mark–recapture, population estimation. Thus, our study had 2 objectives: 1) to compare samples of known Mexican wolves, coyotes, and dogs to identify canid microsatellites with unique alleles for Mexican wolves and sufficient variability to distinguish closely related individuals and 2) to evaluate the feasibility of generating population estimates of endangered Mexican wolves using noninvasive DNA (fecal samples) combined with mark–recapture methods.

STUDY AREA

Within the roughly 17,750 km² BRWRA, we demarcated 2,500 km² in the Apache–Sitgreaves National Forest in

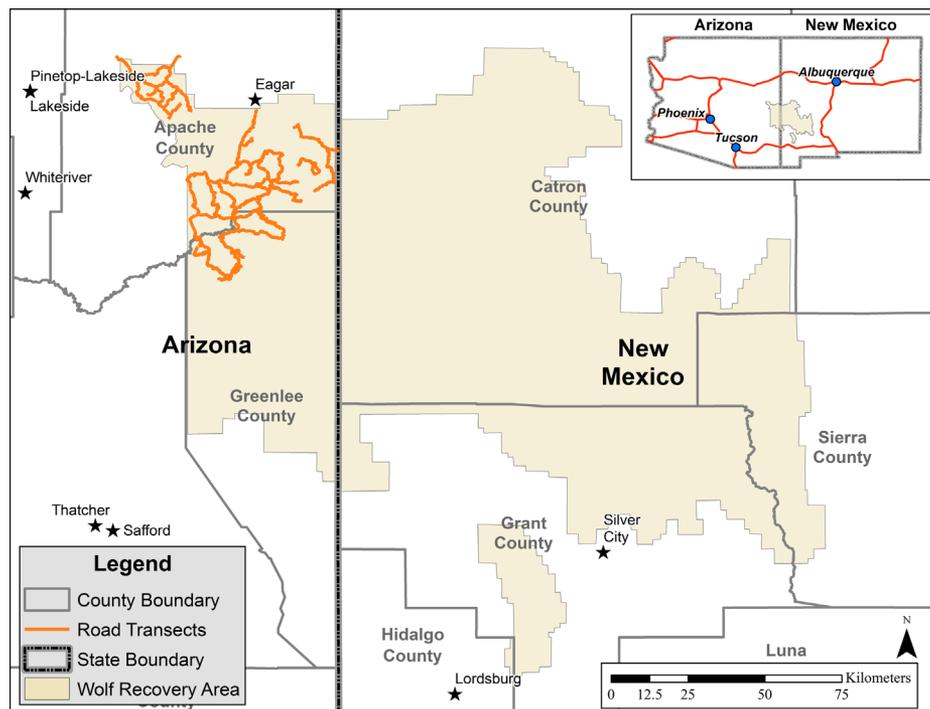


Figure 1. Boundaries of the Blue Range Wolf Recovery Area with the road transects in Arizona, USA, used for collecting canid fecal samples (e.g., Mexican wolf) during 2007 and 2008. The study area of approximately 2,500 km² was bounded by the Fort Apache Indian Reservation on the west; the Mogollon Rim on the south; U.S. Highway 191 on the east; and State Highway 261, County Road 118, and U.S. Forest Service Road 65 on the north.

Arizona (Fig. 1) to conduct the study. The Fort Apache Indian Reservation formed one border of the study area to the west; U.S. Highway 191 to the east; State Highway 261, County Road 118, and U.S. Forest Service Road 65, to the north; and the Mogollon Rim was the southern border. An extensive network of U.S. Forest Service roads existed within the study area (Fig. 1). Four Mexican wolf packs (Paradise, Hawk's Nest, Bluestem, and Rim) occupied the area during the study.

Within the study area, the major types of vegetation included ponderosa pine (*Pinus ponderosa*), aspen (*Populus tremuloides*), fir (*Abies* spp.) juniper (*Juniperus* spp.), piñon pine (*Pinus cembroides*), mesquite (*Prosopis* spp.), oaks (*Quercus* spp.), and several varieties of grasses and forbs (Reed 2003). Potential prey for Mexican wolves in the BRWRA included elk (*Cervus canadensis*), white-tailed deer (*Odocoileus virginianus*), and mule deer (*O. hemionus*). To a lesser extent, pronghorn (*Antilocapra americana*), bighorn sheep (*Ovis canadensis*), and collared peccary (*Pecari tajacu*) were also present (USFWS 1996). Other large predators in the BRWRA included coyotes, mountain lions (*Puma concolor*), and American black bears (*Ursus americanus*; USFWS 1996). Domestic dogs (*Canis lupus familiaris*), including feral individuals, also were present.

METHODS

We identified Mexican wolf fecal samples by detecting the presence or absence of a suite of alleles we characterized as unique to Mexican wolves. To characterize Mexican wolf allelic diversity, the USFWS National Forensic Laboratory (NFL) generated genotypes from tissue and blood samples from 235 Mexican wolves (including 12 confirmed by aerial radiotelemetry by USFWS to have been in our study area and 2 not collared but identified in our study area). To compare this allelic diversity to other canids whose fecal samples might be confused with Mexican wolves, the NFL also genotyped 48 coyotes from BRWRA, and 14 domestic dogs. The NFL genotyped all samples using 10 canid-specific microsatellite loci developed and characterized from the dog genome (Ostrander et al. 1993, Francisco et al. 1996). The NFL used 5 panels, using the PCR conditions described by Fain et al. (2010) and the following primer concentrations for the loci used in this study: Panel 1: CXX.204 (0.83 μ M), CXX.140 (0.04 μ M); Panel 2: CXX.123 (0.33 μ M), CXX.225 (0.83 μ M), FH2096 (0.063 μ M); Panel 3: CXX.172 (0.1 μ M), CXX.250 (0.05 μ M); Panel 4: FH2001 (0.17 μ M), FH2137 (0.083 μ M); Panel 5: FH2010 (0.04 μ M). The NFL binned, evaluated, and scored alleles using genemapper v. 4.0 (Applied Biosystems [ABI], Foster City, CA, USA). The NFL assessed allele frequencies using the program GenAlEx (see below). Loci with allelic frequencies found to be unique to Mexican wolves (e.g., not found in sampled dogs or coyotes) were identified. We included these loci in a panel used as a prescreen protocol for all fecal samples (see Panel A below) to identify and eliminate nontarget fecal samples.

To collect Mexican wolf fecal samples, we established 8 60-km transects on U.S. Forest Service roads because wolves travel along existing roads, trails, and waterways and often

deposit feces along these pathways (Mech 1970). Transects were navigable by 4-wheel-drive, high-clearance vehicles. After initially clearing transects of fecal material, teams of 2 volunteers in vehicles driven at speeds not exceeding 20 km/hr surveyed and collected all potential canid fecal samples from each transect during 3 sampling periods (Sep 2007, Nov 2007, and May 2008). During each sampling period, we surveyed transects 5 times (every 5 days) and recorded Global Positioning System coordinates for all sample collection points. During September 2007, we stored 2-g pieces of fecal samples in vacuum-sealed 25-mL vials with 4 g of silica/g of feces (Wasser et al. 1997). Filter paper barriers separated fecal samples from silica dust. Within 2 days of receiving the samples, we performed fecal scraping (Kohn et al. 1999, Prugh et al. 2005) to procure epithelial cells of the predator species rather than any prey DNA. We adapted the storage protocol for fecal samples for the remaining 2 sampling periods (Nov 2007 and May 2008) because of poor results from storing fecal material in silica beads (see Results). During these collection periods, we stored 2-g samples of fecal scrapings (performed in the field) in 2.0-mL microcentrifuge tubes containing a DET buffer in a 1:4 volume-sample-to-solution ratio (Seutin et al. 1991, Prugh et al. 2005). Field personnel stored all samples in freezers until they shipped them to the National Wildlife Research Center (NWRC) laboratory and DNA was extracted promptly after samples arrived.

We extracted DNA from fecal samples following the protocol for DNA isolation from human stool using QIAamp DNA Stool Minikits (Qiagen, Valencia, CA, USA). We conducted all DNA extractions in a room that was free of canid DNA, polymerase chain reaction (PCR) products, and separate from the laboratory where PCRs and post-PCR processing was conducted. We used positive and negative controls with all extractions and PCRs and a single individual (MAN) performed all genotypic scoring to ensure consistency.

We developed and tested our laboratory techniques for fecal species identification and scoring at the NWRC by obtaining fecal and blood samples from 8 captive Mexican wolves at the Sevilleta National Wildlife Refuge, La Joya, New Mexico. Fecal samples were collected from the refuge in vacuum-sealed 25-mL vials with 4 g of silica/g of feces (Wasser et al. 1997) to act as a desiccant. We used filter paper barriers to prevent silica dust from embedding itself on the surface of the fecal samples. We extracted DNA from blood samples using a Qiagen DNeasy Blood and Tissue Kit (Qiagen). Further, for validation of consistency of results and alleles binned at the NFL and NWRC, we tested 16 Mexican wolf DNA extractions from tissue samples in both laboratories. All fecal genotyping was conducted at the NWRC. The NWRC amplified 10 loci using 3 panels in 11.0–12.0- μ L reactions. Panel A contained the markers CXX.172 and CXX.204, which have private alleles unique to Mexican wolves (see Results). Locus FH2010 (Ostrander et al. 1993, Francisco et al. 1996) was also in Panel A, and does not have alleles unique to Mexican wolves but amplified well in fecal samples with good-quality DNA and therefore,

served to test fecal sample quality. Panel A was used as a prescreen, where all samples were tested along with a negative control and 2 positive controls (DNA from Mexican wolf blood and Mexican wolf fecal samples from Sevilleta National Wildlife Refuge). If the positive controls amplified well and the negative control was negative, then any sample that failed at all 3 markers was considered low-quantity or low-quality DNA and discarded. If at least one locus amplified for a sample and it was judged to be a Mexican wolf or inconclusive then Panel B was run. If at least one locus amplified in Panel B and those alleles or a combination of alleles suggested Mexican wolf (see Results) or was inconclusive, then Panel A was run 2 more times, Panel B run 2 more times, and Panel C run 3 times (see Supporting Information for detailed protocol) following the multiple tubes approach (Taberlet et al. 1996) to identify and account for alleles missed through allelic dropout or added as false alleles. The prescreen approach was used to minimize the cost of processing hundreds of fecal samples where only a small proportion were likely to be Mexican wolf.

Panel B contained the markers CXX.123, CXX.225, and CXX.250 (Ostrander et al. 1993). Panel C contained the markers FH2001, FH2096, FH2137, and CXX.140 (Ostrander et al. 1993, Francisco et al. 1996). Panels A and B were 12.0- μ L reactions and Panel C was an 11- μ L reaction. Panel A used 3.2- μ L 5 \times buffer C (Invitrogen, Carlsbad, CA, USA), 0.42 mM dNTP (Invitrogen), 0.17 μ g bovine serum albumin (BSA), primers (CXX.172 0.25 μ M, FH2010 0.67 μ M, CXX.204 0.16 μ M), 0.8 units of Taq (Promega, Madison, WI, USA), and ddH₂O to total volume of 10 μ L. Panel B used 3.0 μ L 5 \times buffer C (Invitrogen), 0.42 mM dNTP (Invitrogen), 0.17 μ g BSA, primers (CXX.123 0.25 μ M, CXX.225 0.25 μ M, CXX.250 0.16 μ M), 0.8 units of Taq (Promega), and ddH₂O to bring total volume to 10 μ L. Panel C used 1.0 μ L ABI 10 \times buffer, 0.84 mM dNTP (Invitrogen), 2.27 mM MgCl₂ (ABI), 0.17 μ g BSA, primers (FH2001 0.03 μ M, FH2096 0.02 μ M, FH2137 0.03 μ M, and CXX.140 0.02 μ M), 0.2 μ L AmpliTaq (ABI), and ddH₂O to bring total volume to 9 μ L. We added 2 μ L of DNA template to the mix for each panel. The PCR amplification profile for Panels A and B was an initial denaturation at 94 $^{\circ}$ C for 4 min, followed by 30 cycles of denaturing at 94 $^{\circ}$ C for 15 s, annealing at 58 $^{\circ}$ C for 15 s, and extension at 72 $^{\circ}$ C for 30 s, with a final extension period of 30 min at 60 $^{\circ}$ C. The PCR amplification profile for Panel C was initial denaturation at 95 $^{\circ}$ C for 10 min, followed by 52 cycles of denaturing at 94 $^{\circ}$ C for 30 s, annealing at 55 $^{\circ}$ C for 30 s, and extension at 72 $^{\circ}$ C for 45 s, with a final extension period of 7 min at 72 $^{\circ}$ C. We mixed PCR products with GeneScan 400HD ROX Size Standard (ABI) and visualized all fragments on an ABI 3130 genetic analyzer following standard protocols. Alleles per locus were binned and scored using genemapper v. 4.0 (ABI).

We used the program GenAlEx to 1) compare allelic frequencies between Mexican wolves, coyotes, and dogs; 2) estimate the probability of identity between individuals and siblings (P_{ID} and $P_{ID_{sib}}$; Peakall and Smouse 2006) from

Mexican wolf fecal genotypes with all 10 markers; 3) detect initial occurrence and reoccurrences of fecal genotypes; and 4) identify matches between genotypes from our fecal collection and of Mexican wolves that were known to exist within our study area during our study period. We calculated the probability of identity for all individuals (P_{ID}) and between potential siblings ($P_{ID_{sib}}$) to confirm whether or not we had enough power to distinguish individuals, including closely related ones with the markers we were using (Waits and Leberg 2000, Waits et al. 2001). We used only the final genotypes identified as Mexican wolf from our fecal samples for this analysis. Individual Mexican wolves had been captured, collared, tracked, sampled, and genotyped as part of the USFWS monitoring program. This program was ongoing during our study. We compared genotypes of Mexican wolves known to be on the ground in our study area during our study period to fecal genotypes we obtained to see how many known individuals we captured in our noninvasive sampling effort. We used the program Microchecker to check for null alleles and scoring errors in our fecal genotypes (Van Oosterhout et al. 2004).

We performed analyses of the data (our fecal genotypes, Mexican wolf genotypes from blood and tissue samples from individuals known to be in our study area during our study period, coyotes, and dogs) in STRUCTURE with a burn-in of 100,000 and 500,000 total iterations to assess genetic clusters. Our model was run assuming admixture and correlated allele frequencies because we considered this was a conservative approach that would maximize evidence of admixture if it existed. We tested for $K = 1-6$ and tested each K 5 times. This analysis was performed to assess the ability of our chosen markers to reliably distinguish Mexican wolves from coyotes in the study area from dogs.

We generated encounter histories for each genotype and used these histories in conjunction with 14 Huggins-type robust design mark-recapture models to estimate abundance (Pollock 1982, Huggins 1989). We estimated survival (S) between primary sampling periods for all models. We estimated probability of emigration (γ') for some models (indicated by G) and fixed the parameter at 0 for other models (indicated by $G = 0$). We estimated initial capture probabilities as constants (indicated by $p[.]$) and as varying across primary capture periods but not across secondary capture periods (indicated by $p[t]$). Some wolves were likely to have a greater capture probability than others because of the location of particular packs relative to our study area boundary. We attempted to model capture heterogeneity by recording the average distance to the edge of the study area from the location at which a genotype was found and used this individual covariate to explain the variation in capture rates between genotypes. Our choice of Huggins-type models to model our data was dictated by our attempt to use individual covariates to explain capture heterogeneity (indicated by $p[\text{Distance}]$). We estimated recapture probabilities in a similar manner (indicated by $c[.]$, $c[t]$, and $c[\text{Distance}]$). We derived the population estimate (\hat{N}) for each primary sampling period with Program MARK (White and Burnham 1999). We removed all models with model

weight (w_i) < 0.01 from the candidate set and used model averaging to generate point estimates and confidence intervals of Mexican wolf abundance. We incorporated model uncertainty for our model-averaged estimates as outlined by Buckland et al. (1997).

RESULTS

Using genotypes from 235 Mexican wolves, 48 coyotes, and 14 dogs in combination with our fecal genotypes, we identified unique Mexican wolf alleles or combinations of alleles (Table 1). Two loci (Panel A: CXX.172 and CXX.204) had diagnostic alleles for Mexican wolves when compared with coyotes from the area and domestic dogs. All other loci had alleles that were not unique to Mexican wolves, but some allele combinations across loci were diagnostic of a Mexican wolf (Table 1; Supporting Information).

We collected 103, 282, and 194 fecal samples during September 2007, November 2007, and May 2008, respectively. The Panel A prescreen was run on 579 samples, only 66 samples were tested for Panel B's first replicate, and 24 samples were tested for all remaining panel replicates (Supporting Information; Table 1). To utilize only the

most reliable genotypes, our criteria for final analysis were that a sample was genotyped at all 10 loci ($N=15$) and was categorized conclusively as a Mexican wolf ($N=13$). No fecal samples from September met our criteria because none produced complete genotypes. Therefore, for subsequent results we will not refer to this sampling period. Thus, November 2007 is the first sampling period and May 2008 will be the second sampling period. We identified 6 fecal samples from the November collection period and 7 fecal samples from May as Mexican wolf. Overall, there were 8 unique fecal genotypes identified throughout the study. One genotype was encountered 3 times—once during the first sampling period, and twice during the second one. Three genotypes were encountered twice—one was encountered in each period, one was encountered twice during the first sampling period, and one was encountered twice during the second sampling period. Four genotypes were only encountered once—2 during the first and 2 during the second primary session. Therefore, during each sampling period, we identified 5 unique genotypes (2 genotypes were found in both periods). As expected, allelic diversity was low among fecal genotypes with 1 fixed locus and 1–4 alleles/loci. Yet, the estimated P_{ID} ($1.4E-04$) and $P_{ID_{sib}}$ ($1.6E-02$) at 10 loci for the fecal Mexican wolf genotypes were lower than the generally accepted guidelines for genetic mark-recapture studies (Waits et al. 2001).

There were 12 wolves from 4 packs (Paradise, Hawk's Nest, Bluestem, and Rim) detected in the study area based on Interagency Field Team aerial radiotelemetry across the sampling periods. There were 2 more individuals that were not collared that were later determined to have been in the area during one or both study periods. When we examined matches of the fecal genotypes to genotypes from known individuals (total $N=14$), we found matches to 5 individuals (36% of wolves in area) from 3 packs (Hawk's Nest, Bluestem, and Rim). Of these 5 individuals, we detected 3 of them twice (see above) and 2 of them a single time both in the second sampling period. Further, we had 3 fecal genotypes that identified as Mexican wolves but did not match to any individuals detected on the ground through Interagency Field Team capture efforts. One of these fecal genotypes was detected 3 times (see above) across both sampling periods and the other 2 were detected only once; both in the first sampling period. The known abundance of Mexican wolves in our study area during our November sampling period was 10 and during May it was 10 (USFWS 2008) so for each sampling period we identified 50% of the wolves known to be in our study area. No null alleles were detected.

The analyses of genetic clusters using STRUCTURE found that $K=3$ had the same probability as $K=4$ (Fig. 2). However, $K=3$ had less variance; and when the clusters at $K=4$ were observed, each individual coyote was divided evenly between 2 clusters (Fig. 2). Therefore $K=3$ was the most biologically meaningful answer. When assignment to clusters for each individual across all 5 runs of $K=3$ was averaged, Mexican wolves (including our fecal genotypes and 14 Mexican wolves from BRWRA) assigned to a single

Table 1. Loci and alleles that were used to identify Mexican wolves from fecal genotypes. The diagnostic alleles, or combination of alleles, were identified from genotyping 235 Mexican wolves, 49 coyotes, and 14 dogs from canid scats collected via road transects conducted during 2007 and 2008 in the Blue Range Wolf Recovery Area in Arizona, USA. "Inconclusive" means that these alleles can be found in Mexican wolves but are also found in coyotes and dogs. Panels B and C inconclusive alleles serve as confirmatory alleles when Panel A alleles point to a potential Mexican wolf species identification.

Panel A	Locus CXX.172	Locus CXX.204	Locus FH2010	
Mexican wolf	158	215		
Inconclusive	157 ^a	217 209 211	232 236 240 244 248	
Panel B	Locus CXX.123	Locus CXX.225	Locus CXX.250	
Mexican wolf				
Inconclusive	152 158	167 169 ^b	140 144	
Panel C	Locus FH 2001	Locus CXX.140	Locus FH2096	Locus FH2137
Mexican wolf				
Inconclusive	150 154 158	137 145 151	104 108	166 168 174

^a Allele 157 from locus CXX.172 is very common in Mexican wolves (>90%) but is also reported in the literature in coyotes (<3%; allele 152; Fain et al. 2010), but it has not been seen in coyotes from the Blue Range Wolf Recovery Area.

^b Allele 169 from locus CXX.225 is the most common allele in Mexican wolves (>90%), so although it is found in other canid species (from our samples = 11% dogs and 45% coyotes), it can serve as a diagnostic when other alleles suggest a Mexican wolf identification.

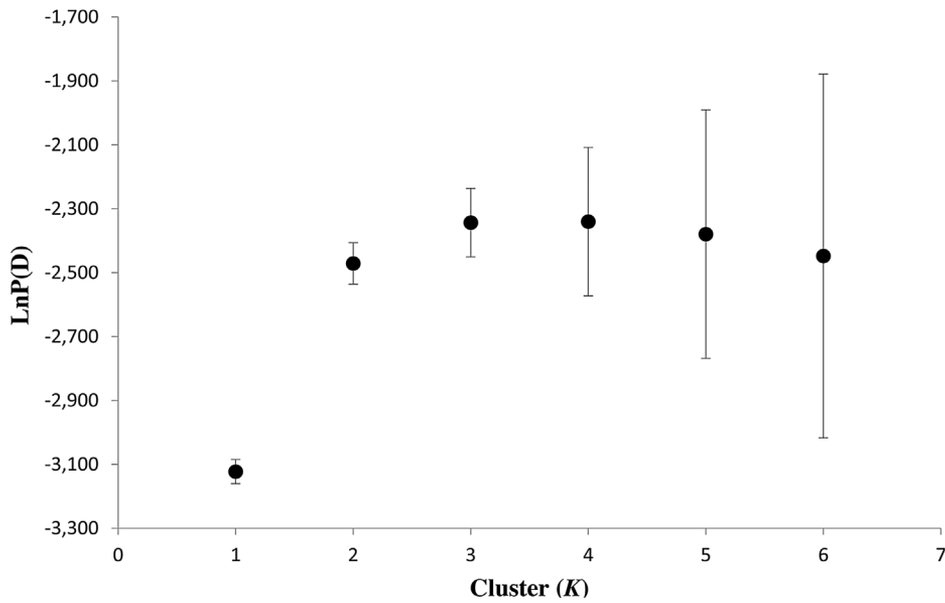


Figure 2. Graph of results from STRUCTURE ($K=1-6$). Log-likelihood values $\text{LnP}(D)$ and the number of clusters tested, including the variance of values across each of 5 independent runs. This analysis tested the genetic clusters formed by fecal genotypes presumed to be Mexican wolf, genotypes from tissue or blood of Mexican wolves known to be in the study area (Blue Range Wolf Recovery Area, AZ, USA) during the study, coyotes from the Blue Range Wolf Recovery Area, and domestic dogs (collected via road transects during 2007 and 2008). $K=3$ was the best result and all our fecal genotypes fell into a genetic cluster with known Mexican wolves, while coyotes formed a second cluster, and dogs formed the third cluster.

cluster (0.993) with a standard deviation (SD) of 0.009; coyotes on average assigned to a single cluster (0.971) with SD of 0.041; and dogs assigned to a single cluster (0.951) with SD of 0.061. Therefore, each species formed its own genetic cluster and our fecal genotypes all clustered with Mexican wolves. STRUCTURE results show that the private alleles detected in Mexican wolves drive much of the Mexican wolf cluster assignment.

Our robust design models contained only 2 primary collection periods where 8 unique Mexican wolf genotypes were encountered. Of the 14 original models (Table 2), we

retained the top 8, each with a $w_i > 0.01$, and discarded the remaining 6 from our candidate set. The estimates of abundance derived from each model varied between 5.1 and 13.1 for November and between 5.7 and 9.9 for the May collections (Table 3). Our model-averaged abundance estimate was 10 (95% CI = 6–34) during November. Our model-averaged abundance estimate was 9 (95% CI = 6–30) during May. All final estimates were rounded up to the next greatest integer. Distance from the site of collection of a genotype to the edge of the study area did not improve the modeling of initial or recapture probabilities even though we

Table 2. Model support for Huggins-type robust mark–recapture models used to estimate Mexican wolf abundance in the Blue Range Wolf Recovery Area in Arizona, USA. Data are fecal genotypes generated from samples collected within the study area during 2 sampling periods in 2007 and 2008. We used Akaike Information Criterion corrected for small sample size (AIC_c) to rank models and present associated model weights (w_i).

Model ^a	Parameters estimated	AIC_c	ΔAIC_c	w_i
{ $S, G=0, p[.] = c[.]$ }	2	61.5	0.0	0.49
{ $S, G=0, p[.], c[.]$ }	3	63.3	1.9	0.20
{ $S, G=0, p[t] = c[t]$ }	3	64.6	3.1	0.10
{ $S, G, p[.] = c[.]$ }	3	64.9	3.5	0.09
{ $S, G=0, p[t], c[.]$ }	4	66.6	5.2	0.04
{ $S, G=0, p[.], c[t]$ }	4	66.9	5.4	0.03
{ $S, G, p[.], c[.]$ }	4	67.7	6.1	0.02
{ $S, G, p[t] = c[t]$ }	4	68.9	7.5	0.01
{ $S, G=0, p[\text{Distance}] = c[\text{Distance}]$ }	2	70.1	8.6	0.01
{ $S, G=0, p[t], c[t]$ }	5	71.4	10.0	0.00
{ $S, G, p[t], c[.]$ }	5	72.2	10.7	<0.01
{ $S, G, p[.], c[t]$ }	5	72.5	11.0	<0.01
{ $S, G, p[\text{Distance}] = c[\text{Distance}]$ }	3	73.6	12.1	<0.01
{ $S, G, p[t], c[t]$ }	6	78.9	17.4	<0.01

^a S indicates that the survival parameter was estimated; $G=0$ indicates that the γ'' (probability of emigration) parameter was fixed at 0, and G indicates that the γ'' parameter was estimated; $P[.]$ or $c[.]$ indicates that the initial capture or recapture probabilities were modeled as constants, $p[t]$ or $c[t]$ indicates that they varied across primary capture periods but not across secondary capture periods, and $p[\text{Distance}]$ or $c[\text{Distance}]$ indicates that they were modeled as a function of the average distance from the site of collection of a genotype to the edge of the study area.

Table 3. True abundance and abundance estimates (\hat{N}) as well as estimates of survival (S), emigration (G), capture probability (p), and recapture probability (c) of the top 8 Huggins-type robust mark–recapture models used to estimate abundance of Mexican wolves from fecal genotypes collected in a study area within the Blue Range Wolf Recovery Area in Arizona, USA, during 2 sampling periods in 2007 and 2008.

Model ^a	Nov 2007						May 2008					
	(True $N = 10$)						(True $N = 10$)					
	\hat{N}	SE	S	G	p	c	\hat{N}	SE	S	G	p	c
{ $S, G = 0, p[.] = c[.]$ }	9.9	4.8	0.95	0.00	0.13	0.13	9.9	4.8	0.95	0.00	0.13	0.13
{ $S, G = 0, p[.], c[.]$ }	5.7	1.3	0.85	0.00	0.35	0.11	5.7	1.3	0.85	0.00	0.35	0.11
{ $S, G = 0, p[t] = c[t]$ }	13.1	10.9	0.92	0.00	0.09	0.09	8.3	4.1	0.92	0.00	0.17	0.17
{ $S, G, p[.] = c[.]$ }	9.9	4.8	0.99	0.19	0.13	0.13	9.9	4.8	0.99	0.19	0.13	0.13
{ $S, G = 0, p[t], c[.]$ }	5.1	0.48	0.91	0.00	0.52	0.11	7.8	7.5	0.91	0.00	0.18	0.11
{ $S, G = 0, p[.], c[t]$ }	5.7	1.26	0.85	0.00	0.35	0.06	5.7	1.3	0.85	0.00	0.35	0.17
{ $S, G, p[.], c[.]$ }	5.7	1.26	0.87	0.08	0.35	0.11	5.7	1.3	0.87	0.08	0.35	0.11
{ $S, G, p[t] = c[t]$ }	13.1	10.8	0.92	0.00	0.09	0.09	8.3	4.0	0.92	0.00	0.17	0.17

^a S indicates that the survival parameter was estimated; $G = 0$ indicates that the γ'' (probability of emigration) parameter was fixed at 0, and G indicates that the γ'' parameter was estimated; $p[.]$ or $c[.]$ indicates that the initial capture or recapture probabilities were modeled as constants, and $p[t]$ or $c[t]$ indicate that they varied across primary capture periods but not across secondary capture periods.

found the suggestion of a relationship between the number of times an individual was captured and distance to edge of the study area (Table 4). We conducted a Poisson regression of the data and the parameter estimates were not statistically significant, primarily because of small sample size (Table 4).

DISCUSSION

Our primary objective was to evaluate the utility of using noninvasive fecal DNA genotypes of Mexican wolves combined with mark–recapture methods to generate population estimates. Our results indicated that such techniques could be a viable alternative for monitoring Mexican wolf populations. Another objective was to compare samples of known Mexican wolves, coyotes, and dogs to identify canid microsatellites with unique alleles for Mexican wolves. Our genetic methodology proved effective for distinguishing Mexican wolf scat from dogs and coyotes in the BRWRA and during 2 sampling occasions we were able to identify 5 of 14 individual Mexican wolves known to be members of 3 different packs and 3 previously undetected wolves occurring in the study area during both our sampling periods.

Although the low numbers of wolves we detected and recaptured resulted in wide confidence intervals, our

estimates of abundance corresponded with the known number of wolves in the study area. Intensive capture and surveillance efforts of Mexican wolves did not detect 3 individuals identified from fecal genotypes. This is not unexpected because traditional mark–recapture methods can miss elusive animals including juveniles. We detected each of these individuals initially during the November 2007 sampling effort, so these could be young of the year not yet captured and collared. Alternatively, there could have been uncollared adult wolves in the area. We believe that our lack of success in generating complete genotypes from September samples was due to the inefficacy of the preservation technique we used during that collection period. We were able to generate complete genotypes for subsequent collections after we implemented a different preservation technique. If we had been able to successfully amplify samples from the September 2007 period at all 10 loci, our population estimates may have been more precise and accurate. Regardless, this study demonstrated that our approach could prove useful for monitoring Mexican wolves; however, we suggest a number of changes to the sampling design that could increase the number of Mexican wolf detections.

Our sampling protocol called for surveys of transects every 5 days, but studies (published after we conducted our study) have reported that wild canid fecal samples degrade significantly and provide less robust genotypes after 1–3 days (Santini et al. 2007, Panasci 2009). Ideally, before the implementation of another attempt at fecal collection, an initial study would be conducted to examine Mexican wolf fecal deposition and degradation rates, along with estimates of field and lab costs to allow quantitative evaluation of the ideal temporal sample collection that balances all these issues as has been done for coyotes and foxes in Utah, USA (Lonsinger et al. 2015). Our sampling protocol also called for the collection of all suspected canid feces detected along transects because field methods to distinguish between feces deposited by Mexican wolves and coyotes based on fecal size are unreliable (Reed et al. 2004). Consequently, we collected and prescreened numerous fecal samples that were deposited

Table 4. Evidence of a potential relationship between capture frequency of individual genotypes of Mexican wolves and the average distance from the site of collection of a fecal genotype to the edge of a study area within the Blue Range Wolf Recovery Area in Arizona, USA, generated from samples collected during 2 sampling periods in 2007 and 2008.

Genotype label	No. times encountered	Average distance to edge (km)
A	3	14.2
B	2	11.9
C	2	9.7
D	2	10.6
E	1	4.2
F	1	3.8
G	1	5.1
H	1	0.1

by coyotes and other canids. Financial resources that could have been devoted to surveying transects, either more frequently or more times during additional collection periods, were instead devoted to prescreen genotyping nontarget fecal samples. Improving field identification would help reduce this burden.

Rinkevich (2012) used detection dogs to conduct a similar study to ours in another portion of the BRWRA in 2008 and 2009. Using dogs she collected 558 fecal samples, 265 of which amplified and produced DNA sequence data. Of these, 197 sequences were carnivore and 21 of those were Mexican wolf (her population size estimate was 19), while 111 were coyote. The study concluded that the dogs found a large number of nontarget scats and a low number of wolf scats in the study. We believe further investigation into whether detection dogs can be useful is warranted and should be done in conjunction with a cost–benefit analysis that compares our methodology with that using detection dogs and mark–recapture through radiotelemetry.

Our candidate models only contained the γ'' parameter (which is the probability of being off the study area and unavailable for capture during period i , given that the animal was not present during the previous period and given that it survived) and not γ' (which is the probability of being off the study area and unavailable for capture during period i , given that the animal was present during the previous period and given that it survived). This is because estimation of both these parameters requires ≥ 3 primary sampling sessions, whereas we only had 2 primary sampling sessions as a result of the unavailability of data from our September sampling effort.

We expected heterogeneity in capture rates between individual wolves and therefore used Huggins-type models (Huggins 1989) to account for this heterogeneity. Two of the 4 wolf packs in our study had territories that were completely contained within our study area, while the other 2 packs occupied territory outside the study area. Thus, we tried to model individual heterogeneity by recording the average distance to the edge of the study area from the location at which a sample was collected. However, our hypothesis that individual capture rates might vary depending on the location of pack territories was not sustained by model selection methods. It is likely that our model selection methods strongly penalized the use of additional parameters because of the dearth of usable data; with more data, this may be an effective way of modeling individual capture heterogeneity.

Erroneous addition or omission of new individuals occur when genotypes are incorrectly determined because of PCR replicate error, error from degraded DNA, human error in scoring, or lack of robust loci (Taberlet et al. 1999, Waits and Leberg 2000). Even miniscule errors can lead to significant positive bias in the estimation of population size (Taberlet et al. 1999, Waits and Leberg 2000, Lukacs and Burnham 2005, Waits and Paetkau 2005). Further, unique animals might appear as a single genotype when loci that are not robust enough to distinguish closely related individuals are applied (Taberlet et al. 1999; Waits and Leberg 2000;

McKelvey and Schwartz 2004*a, b*). This artificially increases the recapture rate and results in the underestimation of abundance (Mills et al. 2000). Thus, one of our objectives was to identify microsatellites that were robust (i.e., not error prone) and variable enough to allow discrimination of closely related individuals even though this population is inbred (lacks allelic diversity).

The estimated P_{ID} and P_{IDsib} for the genotypes we collected were sufficiently low for us to conclude that we effectively distinguished closely related individuals. We did not attempt to estimate genotyping error either statistically or within the framework of mark–recapture modeling because of the extremely limited number of Mexican wolf genotypes detected. However, it is our belief that we avoided significant genotyping errors by following suggestions of Paetkau (2003), who argued that genotyping errors are reduced to negligible levels or completely eliminated by using careful laboratory protocol, experienced observers, reliable microsatellite loci, and by rejecting samples of poor quality, which was accomplished through our prescreening protocol (Supporting Information).

We have demonstrated that noninvasive DNA collection paired with mark–recapture analysis is an effective tool for monitoring Mexican wolves, which is consistent with Stenglein et al. (2010, 2011) and Marucco et al. (2012) who conducted noninvasive fecal DNA studies of other wolf species. By genotyping captured Mexican wolves, we have identified a suite of alleles that differentiate Mexican wolves from coyotes and domestic dogs. Therefore, we are confident that the method presented herein is effective for identifying Mexican wolf species from fecal samples. This information will help future researchers perform species identification from noninvasive samples. Our technique provides managers a way to identify and monitor uncollared wolves that may be missed by traditional surveys. Our technique can also be used to estimate temporary emigration and survival. However, managers will have to sample more intensively than we did to tighten the confidence intervals on estimates of abundance or on other parameters that may be biologically pertinent. More intense sampling efforts should include increasing the number of 1) transects, 2) surveys during a collection period, and 3) collection periods during the year (Marucco et al. 2012). Optimization of sampling design could be achieved through another study assessing Mexican wolf fecal deposition and degradation rates (Lonsinger et al. 2015). Sampling can and should be conducted on an ongoing basis and in an adaptive manner until the desired tightness in confidence intervals is achieved. Such efforts will likely lead to the more efficient use of scarce resources and effective monitoring of this endangered species.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article at the publisher's web-site. Supporting information includes a detailed document on the laboratory work flow for identifying Mexican wolves from fecal samples and a Genalex output of allelic frequencies between Mexican wolf blood and tissue samples of wolves on the ground in our study area during the time of our sampling, our fecal samples, coyotes, and dogs. Supporting Information Table 1 shows the total number of PCR runs, failures, Mexican wolves, other canid species, and inconclusives that were identified at each step of the work flow.

Table S1. Number of PCRS performed for each panel, including the number of PCRS/samples that failed to amplify at each marker in the panel, the number identified as Mexican wolf and/or inconclusive. n/a, not applicable.

Supporting Word File. Work flow for Mexican wolf samples.

Supporting Excel File. Genalex output of allelic frequencies between Mexican wolf blood and tissue samples of wolves on the ground in our study area during the time of our sampling, our fecal samples, coyotes, and dogs.