Genetic diversity and sex-bias dispersal of plateau pika in Tibetan plateau

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Abstract
Dispersal is an important aspect in organism’s life history which could influence the rate and outcome of evolution of organism. Plateau pika is the keystone species in community of grasslands in Tibetan Plateau. In this study, we combine genetic and field data to characterize the population genetic pattern and dispersal dynamics in plateau pika (Ochotona curzoniae). Totally, 1,352 individual samples were collected, and 10 microsatellite loci were analyzed. Results revealed that plateau pika possessed high genetic diversity and inbreeding coefficient in a fine-scale population. Dispersal distance is short and restricted in about 20 m. An effective sex-biased dispersal strategy is employed by plateau pika: males disperse in breeding period for mating while females do it after reproduction for offspring and resource. Inbreeding avoiding was shown as the common driving force of dispersal, together with the other two factors, environment and resource. In addition, natal dispersal is female biased. More detailed genetic analyzes are needed to confirm the role of inbreeding avoidance and resource competition as ultimate cause of dispersal patterns in plateau pika.

KEYWORDS
genetic diversity, microsatellite, plateau pika, population structure, sex-bias dispersal

1 | INTRODUCTION

Dispersal of individual is an important aspect in organism’s life history which has profound significance for genetic structure, population biology and gene flow (Garant, Forde, & Hendry, 2007; Slatkin, 1987) that eventually could influence the rate and outcome of evolution of organism (Wright, 1931). The potential driving forces of dispersal mainly focused on several aspects including inbreeding, kin competition, resource competition, environment, and so on. Among them, inbreeding avoidance is recognized as the main cause of dispersal (Bowler & Benton, 2005).

In animals, sexual differences of dispersal are found and documented mainly on rates and distance. Dispersal rate is male biased in most of mammal species (Greenwood, 1980) and males also disperse in longer distances than females (Waser & Strobeck, 1998). Recently, some empirical studies have demonstrated the differences in dispersed distance between sexes (Blundell, Ben-David, Groves, Bowyer, & Geffen, 2002; Fontanillas, Petit, & Perrin, 2004; Ji, Sarre, Aitken, Hankin, & Clout, 2001). The reasons of long- or short-distance dispersal are likely to be very different (Ronce, Olivieri, Clobert, & Danchin, 2001). It has been evidenced that short-distance dispersal is sufficient for avoiding inbreeding or kin competition, whereas long-distance dispersal might function to colonize a new territory or escape (Perrin & Goudet, 2001). Dispersed distance is very informative for explicating the evolutionary cause of dispersal (Murrell, Travis, & Dytham, 2002). Furthermore, dispersed time might also be able to be another cause of dispersal. The timing of dispersal can be quite variable which...
depends on the species’ life history (Schülke, 2003; Smale, Nunes, & Holekamp, 1997). Natal dispersal is thought to function as an effective means of spatially separating opposite-sexed kin, alleviating the risk of close inbreeding (Dobson, 1982; Hazlitt, Eldridge, & Goldizen, 2004); while many species undergo secondary dispersal later in life, which is motivated for several purposes: such as aggressive eviction, and mating competition (Packer, 1979). In addition, dispersal increases with the social complexity (Perrin & Lehmann, 2001).

Plateau pika (Ochotona curzoniae) is small, diurnal, and nonhibernating rodent, belonging to lagomorph family, Ochotonidae. They mainly distribute within and around the Tibetan Plateau. They are highly social burrow dwellers with seasonally breeding (Dobson, Smith, & Wang, 1998; Lai & Smith, 2003; Smith & Foggin, 1999; Smith & Wang, 1991). They play pivotal role in the community dynamics of high-altitude grasslands and are considered a key species in Tibetan Plateau.

With the development of statistical and genetic methods, many studies have performed spatial genetic structure analysis using genetic markers (Hazlitt et al., 2004; Peakall, Ruibal, & Lindenmayer, 2003). To date, most conclusions about genetic structure and dispersal of plateau pika are based on observational studies without full consideration of the spatial genetic structure of population (Dobson et al., 1998; Smith & Wang, 1991). In this study, we used a set of 10 microsatellite genetic markers to investigate the spatial genetic structure of plateau pika in a fine-scale population, together with the observation data of field. The goals of this study are as follows: (i) dynamic of genetic diversity and population structure in plateau pika; (ii) sex-biased dispersal pattern of plateau pika.

2 | MATERIALS AND METHODS

2.1 | Field description

Plateau pika inhabits at the sample collection site, Maqin County (34°24′N, 100°21′E), Tibetan Autonomous Prefecture of Golog, Qinghai Province, People’s Republic of China, where the average altitude is about 3,846 m (Figure 1). It is dry and cold, and the rainfall is mainly in summer (from June to August). The predominant vegetation of this area is kobresia humilis meadow. In the habitat, plateau pika is the most abundant rodent (Jiapeng et al., 2016).

2.2 | Sample collection

Pikas were trapped using catch-mark-release method within a plot of 200 × 200 m region (the core region with 150 × 150, and
50 m around in distance as the relief area, Figure 1). We used toe clipping to mark pika and obtain tissues. Trapping is performed in May–August from 2005 to 2009 (Table S1). Body mass, age, time of recruitment, reproductive condition, and the relative coordinate of trapped sites were recorded. During breeding season, two groups could be recognized: (i) mature one: overwintered adults who attend breeding; (ii) immature one: born in mating season, sexual immaturity who do not attend breeding (Table S1). All samples are carried out in strict accordance with the ethical guidelines approved by the Animal Care Commission of Plateau Biology, Chinese Academy of Sciences.

A total of 1,352 samples were trapped. Toe tissues are immediately preserved in 90% ethanol and then stored in 20°C.

2.3 DNA extraction and microsatellite analysis

Genomic DNA was extracted by standard phenol–chloroform extraction protocol and quantified by spectrophotometry and agarose gel electrophoresis. Characteristics of microsatellite markers are summarized in Table S2. All 10 microsatellite loci are amplified and genotyped basing on Kexin, Jianing, Jin, Yanming, and Songnian (2009).

2.4 Detection of genetic outliers

GeneClass2 was used to identify genetic outliers which might have been misidentified in the field (Piry et al., 2004). Using the frequencies, we calculated the log-likelihood that an individual belongs to a population from a given year, and then we assessed the probability of an individual to be a resident using the Monte Carlo resampling algorithm (Peakall, Smouse, & Huff, 1995).

2.5 Genetic variability and genetic differentiation over time

Genetic variables in the five consecutive years were analyzed by CERVUS 3.0.3. Locus-specific information was obtained, including number of alleles (Ne), observed heterozygosity (Ho), expected heterozygosity (He), allelic richness (AR), gene diversity (GD), polymorphism information content (PIC), and Hardy–Weinberg equilibrium (Marshall, Slate, Kruuk, & Pemberton, 1998).

2.6 The genetic structure of fine-scale population

Population genetic structures were analyzed by STRUCTURE 2.3.4 (Evanno, Regnaut, & Goudet, 2005; Pritchard, Stephens, & Donnelly, 2000). We assumed an ancestry model: incorporating admixture, a model of correlated allele frequencies, without prior information corresponding to the origin of samples. Ten independent runs of K = 1–30 were performed at 100,000 Markov Chain Monte Carlo (MCMC) repetitions and a 100,000 burn-in period. The most likely K was identified by web-based tool STRUCTURE HARVESTER v0.6.8 (Earl & Von-Holdt, 2011).

2.7 Assignment tests and relatedness

Assignment tests were executed to evaluate the sex-bias dispersal of population via the program FSTATAT version 2.9.3.2 (Goudet, 2002). Genetic variance among populations (FST), mean/variance of assignment index (mAlc/vAlc), and the average relatedness of individuals within a population relative to the whole sample (r = 2FST/1+FST) were analyzed for both sexes using 10,000 permutations in the menu of biased dispersal.

2.8 Spatial genetic structure and sex-biased dispersal analysis

Program GenAlEx 6.5 was used to calculate the matrices of pairwise genetic distances among groups (Peakall & Smouse, 2006). Multilocus spatial autocorrelation analysis was proven powerful and flexible for detecting fine-scale genotypic structure in animals (Peakall et al., 2003; Smouse & Peakall, 1999; Smouse, Peakall, & Gonzales, 2008). This method does not need to make assumptions about the relationship between geographic and genetic distances. Banks and Peakall (2012) described a spatially explicit individual-based simulation framework to investigate the effects of sex-specific variation in dispersal on fine-scale genetic structure (Banks & Peakall, 2012). In this study, we used this method to analyze the dispersed pattern of pika. Briefly, there were three steps. First, we estimated the bootstrap 95% confidence intervals about r by drawing with replacement within the set of relevant pairwise comparisons for a given distance class. Second, the standard spatial autocorrelation analysis of sexes was performed in separate populations, and then the nonparametric heterogeneity tests were computed (Smouse et al., 2008). Finally, the two correlograms drawn on the p values for the T2 statistic at each distance class across both populations were compared, and the correlograms wide “Omega” (w) were computed. In our study, the distance class is 10 m (first distance class is 0–10 m). The first distance class was determined based on the family zone of plateau pika from previous study and our observation of field (Dobson et al., 1998). Data from each sampling period were explored at geographic distance intervals ranging from 10 to 200 m and 10 m intervals.

3 RESULTS

3.1 Population genetic diversity and inbreeding coefficient in five consecutive years

Genetic diversities in five consecutive years are listed in Table 1. We find high genetic variability in the population: AR varies between 8.21 and 10.87 (Table S4); observed heterozygosity (Ho) is about 0.70 in 5 years; the mean effective number of alleles (Ne) is among 13.7 to 14.9; GD is about 0.8 (Table S3). The PIC ranges from 0.77 to 0.82, which indicate high genetic diversity in populations (PIC value < 0.25, low polymorphism; 0.25 < PIC value < 0.5, intermediate polymorphism; and PIC value > 0.5, high polymorphism). Mean inbreeding coefficient (FIS) ranges from 0.11 to 0.21 among 5 years (Table S5), which implies that the nonrandom mating occurred in populations.
3.2 | Population structure in five consecutive years

We identified genetic clusters following the procedure described by Evanno et al. (2005). In independent simulations of structure analysis, a distinct apex value of DK is at K (Fig. S1). The most probable number of genetic clusters in different years is listed in Table 1. According to the rule of STRUCTURE HARVESTER, the most genetic clusters are 2 in 2005 (K = 2), but it is in contrary with the observation in the field. Considering the result of observation, the clusters are defined 22 (K = 22).

3.3 | Assignment tests and sex differences in relatedness

The assignment test is applied to analyze the sex-biased dispersal in population level. Result suggests that there is no significant difference between opposite sexes in estimates of population structure (FST and r), assignment index (mAlc and vAlc), or inbreeding index (FIS; Table 2). Except for that in 2008. The mAlc is negative for males and positive for females in all years (Table 2). FST, r and mAlc are expected to be higher in philopatric ones than that of dispersed ones. Immigrants tend to have lower AIc values than residents (Goudet, Perrin, & Waser, 2002). In our population, females have greater probability of being residents because the mAIC of females is negative for males and positive for females in all years (Table 2).
Spatial genetic structure correlograms for different pika population at study area from May to August. The sample size at each distance class is provided above each correlogram. The scale for the autocorrelation coefficients is not standardized because of the inherent skew in relatedness between the sexes. (a) Spatial genetic structure correlograms for mature female pika population; (b) Spatial genetic structure correlograms for mature male pika population; (c) Spatial genetic structure correlograms for immature female pika population; (d) Spatial genetic structure correlograms for immature male pika population.

**FIGURE 2** Spatial genetic structure correlograms for different pika population at study area from May to August. The sample size at each distance class is provided above each correlogram. The scale for the autocorrelation coefficients is not standardized because of the inherent skew in relatedness between the sexes. (a) Spatial genetic structure correlograms for mature female pika population; (b) Spatial genetic structure correlograms for mature male pika population; (c) Spatial genetic structure correlograms for immature female pika population; (d) Spatial genetic structure correlograms for immature male pika population.
FIGURE 2 (Continued)
positive while the male is negative. And in 2008, mAic and vAic are significantly higher in males than females (both \( p < .05 \), Table 2), it is consistent with male-biased dispersal. But the number of female is much larger than that of male in 2008, which maybe resulted into the difference of Aic.

3.4 Spatial genetic structure of plateau pika population

In the study, individuals are divided into two groups as described above. Individuals from each period were analyzed in mature group. Mature females are genetically similar to local ones rather than distant ones in May to July; pairs exhibit significant positive spatial genetic structure at 0–20 m, while they are more genetically similar to geographically distant ones in August (Figure 1a). But the mature males are in contrary: more genetically similar to geographically distant males in May to July; pairs exhibited almost no positive spatial genetic structure at 0–20 m, while they are more genetically similar to local distant ones in August (Figure 1b). The difference in spatial genetic structures between sexes is quite distinct at 0–20 m.

In immature group, females are genetically similar to local ones rather than distant ones in May and June; pairs exhibited significant positive spatial genetic structure at 0–50 m, while they are more genetically similar to geographically distant ones in July and August (Figure 1c). However, the males exhibit genetically similar to local ones in whole periods (Figure 1d).

3.5 Sex-biased dispersal based on the genetic spatial autocorrelation

Spatial analysis of multilocus is a powerful approach for detecting sex-biased dispersal in natural populations (Banks & Peakall, 2012). Three statistical tests were used: bootstrap confidence intervals about the autocorrelation \( r \) values and two recently developed heterogeneity tests at the distance class. And whole correlogram levels have been used to identify the sex-biased dispersal.

For maturity group, it showed that female pairs have more pronounced genetic structure than male pairs at the shortest distance class (especially in the first class), as indicated by significantly lower spatial autocorrelation coefficients of males in May to July, while the result is opposite in August. The result indicates that male dispersed while female was more philopatric in mating period from May to July, but the situation reversed in the end of breeding in August (Figures 2 and 4).

For immaturity group, both sexes displayed higher spatial autocorrelation coefficients in May and June, while male was with much higher than female in July and August (Figures 3 and 4). It indicates that the immature males were more philopatric in whole period, and immature females dispersed in July and August. The immature females disperse in July when they reach body maturity and the behavior will continue until August when all females finish the dispersal.
4 | DISCUSSION

4.1 | Genetic diversity and population structure

Genetic diversity is an important aspect of population dynamics which directly relates with evolutionary potential and inbreeding (Hughes, Inouye, Johnson, Underwood, & Vellend, 2008). In our research, plateau pika possesses high genetic diversity (Table 1 and Tables S3–S5). It was evidenced that decrease of observed heterozygosity (Ho) in population could induce the decrease in average fitness of individuals (Reed & Frankham, 2003; Szulkin, Bierne, & David, 2010), so high Ho may indicate great potential fitness of individual. Furthermore, AR is closely related to the adaptive ability of population for environmental changes, and allelic richness is considered as a strong indicator of the evolutionary potential of population (Allendorf, 1986; Allendorf, Luikart, & Aitken, 2012; Caballero & García-Dorado, 2013). Genetic diversity of plateau pika is much more abundant than others. American pika is characterized by low levels of allelic richness (2.77) and observed heterozygosity (0.39). It has reported that genetic diversity decreases as population density declines (Jiayan & Zhibin, 2006). American pika population decline obviously, which is considered as an endangered species (Gibson, Van Der Marel, & Starzomski, 2009; Kelsey, Clayton, & Michael, 2016). Collared pika, which lives primarily on talus, showed a medium genetic diversity (Jessie & David, 2012). We compared several species and found the genetic diversity

| TABLE 3 | Dispersed characteristic of plateau pika |
|---------|----------------------------------------|
| Age     | Dispersed sex | Dispersed time | Dispersed purpose | Dispersed name       |
| Maturity | Male          | May, June, July| Mating, inbreeding avoiding | Second dispersal     |
|         | Female        | August         | Inbreeding avoiding, resource |                   |
| Immaturity | Female   | July, August   | Inbreeding avoiding, resource | Natal dispersal      |
reduces with the loving of elevation (Table S6). It indicates that pika is more suitable for high elevation environment. The divergence in genetic diversity of species may be caused by ecological disturbances. Ecological disturbance is considered as a driver of genetic diversity. Disturbance may cause a loss of genetic diversity within population (in particular, allelic diversity or richness) when population sizes were reduced (Banks, Blyton, Blair, McBurney, & Lindenmayer, 2012).

Furthermore, the inbreeding coefficient of plateau pika is medium in three species (Table 2 and Table S6), but it is still harmful for the stabilization of population. Inbreeding depression could reduce population performance through increasing the recessive or deleterious traits of offspring (Banks et al., 2012). Inbreeding depression is closely related to the evolutionary of animal (Charlesworth & Willis, 2009). For plateau pika, the $F_{is}$ value showed that inbreeding indeed existed in population (Table 2). But this is contradicted with the high genetic diversity in some extent. Inbreeding coefficient is different in three pika species (Table S6). The result show that inbreeding is severe in American pika (Kelsey, Clayton, & Michael, 2016), and then in plateau pika. However, there is almost no inbreeding in collared pika population. This may be related to their life history and dispersal patterns (Lawson & Perrin, 2007). In addition, population size is another factor (Dlugosch & Parker, 2008). American pika reduced rapidly, which may be cause by serious inbreeding in ever-reduced population.

### 4.2 | Characteristic of dispersal in plateau pika population

Among animals, dispersal pattern is different between sexes in terms of distances or rates. It reported that plateau pika was philopatric, and dispersal distance was extremely restricted. Furthermore, they dispersed often among families (Dobson et al., 1998). Our results showed that dispersal distance of plateau pika was very limited, within 20 m, which is consistent with the previous observation of 1.3 family ranges (average distance between centers of activity within families is 5.7 m and average distance separating the centers of activity of neighboring families is 23.8 m; Dobson et al., 1998). The restricted dispersal has also been demonstrated in American pika (Kelsey et al., 2016). It reported that short-distance dispersal was probably sufficient for avoiding inbreeding or kin competition (Perrin & Goudet, 2001).

In our study, the male-biased dispersal is not obvious in pika population. Since warm season is very short in Tibetan Plateau, resource (especially food) is considered as a great limiting factor for performance of alpine biota. So in this analysis, dispersal time has been in full consideration. The timing of dispersal could provide interesting clues to ultimate pattern. Natal dispersal was thought of a fundamental parameter in life history, and population biology (Garant et al., 2007; Slatkin, 1987). And natal dispersal was the most common mechanisms proposed to reduce the risk of inbreeding (Lawson & Perrin, 2007; Perrin & Mazalov, 1999). In this study, the natal dispersal of plateau pika is female biased: female juveniles dispersed while the male juveniles were philopatric, which is uncommon in small mammals. We also saw the second dispersal of pika in the mating period. The mature male dispersed in mating period while female dispersed when they finished reproduction (Figures 2 and 4; Table 3).

The extraordinary dispersal pattern of plateau pika can explain the contrary between genetic diversity and inbreeding coefficient. Dispersal strategies evolve a large variety of reasons. For mature male, the dispersal in mating period may drive by two factors: inbreeding avoidance, which was the common driving force in mammal (Charlesworth & Willis, 2009); and local mate competition, because the survival rate can be almost lower than 50% in small mammals, and the “mortality costs” require high reproductive capacity, so adults need more chance to mate (Johnson & Crossman, 1991). The short-distance dispersal is an effective way for male to get more mating chance with limited costs.

For female (including juvenile and maturity), the dispersal is also related to breeding avoiding, that is reducing the risk of father–daughter mating (Lawson & Perrin, 2007), and resource (Greenwood, 1980). In Tibetan plateau, the environment is harsh; resource is limited for survival, especially in winter. The cost of immigration must be in full of consideration. Female dispersal may be a better choice with less cost; they could be more readily accepted into a new group for social animals. Furthermore, postweaning territory is inheritable, where juveniles (especially for male) need to acquire resource such as a burrow to survive in severe environment (Price & Boutin, 1993). The main reasons of distinctive dispersal pattern for pika are inbreeding avoiding and resource.

The dispersal strategy of plateau is effective, but there are still some defects. Firstly, it is cannot completely avoid inbreeding (Table 2). This may be caused by restricted dispersal. Although parallel dispersal between sexes can reduce the inbreeding greatly, in the confined spaces the related individuals also can mate. Moreover, the short life span can also reduce the inbreeding in some extent. However, the strategy is still successful for plateau pika population.

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### CONFLICT OF INTEREST

None declared.

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**SUPPORTING INFORMATION**

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