Population Dynamics and Genetics of Gerbillus nigeriae in Central Sahel: Implications for Rodent Pest Control

Hima Karmadine1, *, Granjon Laurent2, Gauthier Philippe2, Ndiaye Arame3, Brouat Carine2, Dobigny Gauthier2

1Department of Biology, Abdou Moumouni University (UAM), Niamey, Niger
2Center of Biology for Population Management (CBGP), Campus International de Baillarguet, Montferrier-sur-Lez, France
3Laboratory for Sudano-Sahelian Animal Populations, BIOPASS (UCAD-IRD-CIRAD-ISRA), Dakar, Sénégal

Email address: karmadine@gmail.com (H. Karmadine)
*Corresponding author

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Abstract: Gerbillus nigeriae is a sand-dwelling and semi-arid adapted rodent species restricted to the West African Sahel where it causes extensive damages to cereal crops such as millet and sorghum. It also displays one of the most extensive floating chromosomal polymorphisms currently known in mammals, showing a non-random spatial distribution of diploid numbers (2N). We combined population dynamics and genetics to determine dispersal and mobility parameters of G. nigeriae in the species distribution range characterized by low 2N. To do so, we performed a three-year long population survey at Gangara, in the central east Niger. We used both time-dependent monitoring through capture-mark-recapture (CMR) methods and genetic analyses performed on the 134 monitored individuals. CMR results showed low to very low population densities (maximum 27.5 individuals/ha) throughout the study. Abundance cycle was single-phased and strongly dependent on rainfall patterns. Mobility parameters showed very low individual mobility, with means of distance between successive (re) captures (DRS) and maximal distance between (re) captures (DMR) of 7.8 and 14.4 meters, respectively. Genetic analyses revealed significant isolation by distance as well as spatial structuration, thus confirming poor dispersal capacity. Our results are discussed in terms of rodent pest control in arid areas of Niger where cereal crops production is crucial for human food security.

Keywords: Pest Rodent, Gerbillus nigeriae, Abundance, Mobility, Dispersal, Rodent Control, Niger

1. Introduction

Gerbillus nigeriae (Muridae, Gerbillinae) is a sand-dwelling rodent species that lives within a narrow Sahelian strip from Chad to the Mauritanian coast [1, 2]. This semi-arid adapted species is one of the most abundant and common rodent pests of cereal crops in the West African Sahel. Its diet consists mainly in seeds and, it causes major damage to seedlings and young plants in the field, as well as household and village food stocks of millet, sorghum, cowpea and peanut [3-5]. As an illustration, the destruction of up to 60% of millet seedlings was recorded in the Tanout department in the central-east part of Niger, and local but complete destruction is routinely reported in villages throughout Niger [6].

Methods used by local farmers to manage this rodent pest are often ineffective. In particular, they usually act too late, namely when rodents have already reached too high density to be efficiently controlled. Therefore, it appears essential to design preventive control strategies for G. nigeriae. In order to do so we require a better understanding of local demographic and spatial parameters (mobility, abundance, dispersal, structure) [7]. However, despite its importance for food security, there is a paucity of research on G. nigeriae in Sahelian countries [3, 4, 8].

A population dynamics survey was carried out in Kollo in
the South Western part of Niger, which, allowed the authors
to determine various demographic parameters of this
population as well as some of the factors which may be
responsible for population abundance variations. So far, no
data are available for *G. nigeriae* population dynamics in
eastern and central part of Niger which is mostly inhabited by
rural populations that are almost fully dependent on tropical
rain-fed cereal crops that are susceptible to severe damage by
*G. nigeriae*. It is unknown whether the results obtained in
Kollo can be generalized to all the country for two main
reasons. First, rainfall features as well as climate change
impacts in south-west Niger are not the same that those
observed in central and eastern parts of the country [9].
Second, *G. nigeriae* displays one of the most extensive
chromosomal polymorphism (i.e. due to centric fusions [10])
known in mammals since its diploid number (2N) ranges
from 2N = 60-74 [11, 12]. These karyotypic variations show
a non-random spatial distribution with major biogeographic
groups being defined according to their respective 2N values
ranges (i.e., “high 2N values”, 2N=70-74; “intermediate 2N
values”, 2N=65-69; “low 2N values”, 2N=60-64 [2]). In
Niger, two groups of very different chromosomal variants of
*G. nigeriae* are found (Hima et al., 2011): high 2N cytotypes
(i.e. 2N=70-74) characterize the western part of the country,
while low 2N cytotypes (i.e. 2N=60-64) inhabit the central
and eastern regions of Niger. Such genomic differences may
be associated with divergent eco-evolutionary paths and
potential differences in physiology, behaviour and social
structure [13], hence population dynamics.

In this context, we conducted a three year-long monitoring
survey of *Gerbillus nigeriae* in central east Niger through a
combination of Capture-Mark-Recapture (CMR) and
microsatellite DNA-based genetics analyses. The aim of our
study was to identify relevant periods and spatial scales for a
better management of this major Sahelian rodent pest
species.

### 2. Material and Methods

#### 2.1. Monitoring Framework and Sampling

A Capture-Mark-Recapture experiment was undertaken in
a sandy soil millet field close to the village of Gangara
(14°36′N, 08°30′E) in the central-east part of Niger from
February 2008 to June 2011. It consisted of a 150×150 m
parcel (i.e. 2.25 ha) where 256 locally made wire mesh traps
were arranged in 16, 10 meter-spaced, lines. Two consecutive
traps were separated by ten meters, thus making a grid of 225
squares of 100m² each. Monitoring was conducted during
three and a half years with three sampling periods per year:
in the middle of the dry season (February), at the beginning
and the end of the rainy season (June and October,
respectively). Each trapping session consisted in five
consecutive nights with peanut butter-baited traps set in the
afternoon and checked early in the morning. Rodents were
captured alive, weighted, measured, and recorded for their
reproductive status (i.e. male vs. female, juvenile vs. adults,
internal vs. external testicle, pregnant and suckling females).
They were marked by phalange amputation and ear notch
[14] then released at their exact capture point after which we
systematically waited for the released individuals to enter
safely into a burrow. Phalanges and ears pieces from
individual marking procedures were collected and preserved
in ethanol for further genetics analyses.

#### 2.2. Population Dynamics

Population sizes at each sampling period were estimated
using the “Minimum Number Alive” method (MNA) [15].
MNA is defined as the number of individuals caught in a
capture session, plus those that were not caught at that time
but were caught both previously and subsequently within the
device. It is a widely used index to assess population size
with capture-mark-recapture data [16]. Population density
was obtained by dividing each MNA value by the area of the
grid increased with an “attraction area” which corresponds to
a surface that surpasses the grid on each side by the
equivalent of half the distance between successive recapture
(DRS; see below) [17]. We then tested correlations between
population density fluctuations and monthly rainfalls that
were locally recorded using a rain gauge located in the
village.

Individual mobility parameters were estimated by
calculating (i) the mean distance between successive (re)
captures (DRS), and (ii) the mean maximal distance between
(re) captures (DMR), of all recaptured individuals during one
given sampling temporal session. These distances (DRS and
DMR) were calculated for all individuals as well as for sex
and age categories (male vs. female; young vs. adult
individual). Finally, instantaneous (i.e. within one trapping
sessions) and cumulative (i.e. over the entire trapping history
of any individual) mean home range sizes were estimated
according to the “Minimum Convex Polygon” [18, 19].

#### 2.3. Population Genetics

Genomic DNA was extracted from phalanx or ear tissue
samples collected during individual marking procedures
using the Qiagen DNeasy Blood and Tissue Kit. It was then
used as a template for individual multiplex genotyping at 12
*G. nigeriae* species-specific microsatellite loci, (GN01,
GN11, GN19, GN21, GN24, GN27, GN29, GN37, GN48,
GN51, GN62 and GN78) that were previously developed
[20]. Microsatellite repeats were then genotyped on an
Applied Biosystems ABI 3130xl Genetic Analyzer (Life
Technologies) and analysed using GeneMapper v.4.0.

Genotypic linkage disequilibrium and deviation from
Hardy-Weinberg equilibrium (HWE) were investigated for
each pair of loci and for each locus, respectively, using
Genepop v.4.7 [21]. We corrected for multiple testing by the
false discovery rate approach [22] implemented in the
QVALUE R-package [23]. Genetic diversity was estimated
over all loci by calculating unbiased [24] expected (H_e)
and observed (H_o) heterozygosities as well as the allelic richness.
Deviation from HWE was quantified by computing FIS using
Fstat program [25, 26].

We tested individuals’ relatedness within annual samples using SPAGEDI v.1.4 [27]. To do so, we computed the kinship coefficient of Loiselle ($\rho$) [28] between all pairs of individuals within annual samples, using the whole sample (i.e. all individuals sampled throughout the three year-long monitoring period) as the dataset of reference for allelic frequencies. The corresponding histograms were plotted in R v.3.5.1 [29].

Genetic differentiation was evaluated by comparing $F_{ST}$ values between pairs of sub-samples in Genepop v.4.7 [25, 21]. To assess isolation by distance (IBD) pattern, we tested for the correlation between pairwise geographical (log-transformed) distances and genetic differentiation between individuals, using a Mantel test (10,000 permutations). Genetic differentiation between individuals was estimated using $\varepsilon$ statistics [30], which present a lower but slightly biased variance in contrast to $\hat{a}$ statistics [31]. Confidence intervals of IBD regression slopes were obtained using the ABC bootstrap method [32, 30] while plots were obtained under R v.3.5.1. [29]. IBD was investigated over the whole sample. The mean effective dispersal distance was estimated using the slope of IBD regression, which is inversely proportional to the product of effective density ($D_e$, estimated as the average density of the population) and the mean squared parent–offspring dispersal distance ($\sigma^2$) (IBD slope = $1/4\pi D_e \sigma^2$; [21]). We tested for sex biased dispersal using Fstat program [26], and by comparing spatial autocorrelation patterns for males and females, using SPAGEDI and the Kinship coefficient of Loiselle [27, 28].

3. Results

3.1. Annual Variations of Population Density and Cycle of Abundance

Trapping effort was constant at a rate of 256 trap-nights per day, thus making 1,280 trap-nights per temporal session and a total effort of 14,080 trap-nights for the 11 sessions conducted from 2008 to 2011. In total, 146 individuals were captured (Table 1), thus representing an overall trapping success of 1.13%. Though globally low, important annual (between sampling periods from the same year) and inter-annual (between sampling periods of different years) variations in trapping success were noted: density peaks were observed in June during years 2008 and 2011, and in February during years 2009 and 2010. The minimum of population density was always observed in October.

| Captures | Recaptures | Trapping success (%) | Number of individuals | Estimated numbers of individuals (MNA) |
|----------|------------|----------------------|-----------------------|---------------------------------------|
| February 2008 | 5 | 1 | 0,4 | 4 | 4 |
| June 2008 | 34 | 10 | 2,65 | 24 | 24 |
| October 2008 | 2 | 0 | 0,15 | 2 | 4 |
| February 2009 | 61 | 7 | 4,76 | 54 | 68 |
| June 2009 | 40 | 14 | 3,12 | 26 | 26 |
| October 2009 | 0 | 0 | 0 | 0 | 1 |
| February 2010 | 8 | 2 | 0,62 | 6 | 6 |
| June 2010 | 5 | 2 | 0,4 | 3 | 3 |
| October 2010 | 1 | 0 | 0,08 | 1 | 1 |
| February 2011 | 4 | 1 | 0,31 | 3 | 3 |
| June 2011 | 43 | 6 | 2,89 | 37 | 37 |
The maximal number of individuals that were present within our CMR device was observed in February 2009 (N caught = 54; MNA = 68), whereas the absolute minimum (N = 1) was recorded in November 2009 (Figure 1). Along the study period, densities ranged between 0.44 (October 2009 and 2010) and 30 (February 2009) individuals/ha. Our results show a single-phase cycle, with maximum abundance reached during, or at the end of, the dry season, between February and June. From this period, the number of individuals tends to decrease until it reaches the minimum of the year towards the end of the rainy season (October-November).

The relationship between annual population densities and cumulative rainfall (Figure 1) suggests that the less it has rained during the previous year, the more precocious is the peak of abundance (218 mm cumulative rainfall in 2008 for 68 individuals observed in February 2009). In contrast, higher rainfall leads to a later peak of abundance (311 mm and 473 mm of rainfall in 2007 and 2010, for 24 and 27 individuals in June 2008 and 2011, respectively). It also appears that after two consecutive poor rainy seasons, population density may decrease significantly (a maximum of 6 individuals recorded in 2010 following the weak rainfalls of 2008 and 2009) but it may raise rapidly after one good rainy season. As an example, following the 473 mm rainfall in 2010, the number of individuals reached 37 individuals in 2011, while it has not exceeded 6 in 2010.

Table 2. Estimated individual mobility parameters: average values of DRS and DMR. Numbers of individuals from which DRS and DMR were calculated are shown in parentheses.

|                | DRS (m) | DMR (m) |
|----------------|---------|---------|
| February 2008  | 10 (1)  | 10      |
| June 2008      | 7.7 (10)| 28.3    |
| February 2009  | 8.0 (7) | 10      |
| June 2009      | 6.0 (14)| 14.14   |
| June 2010      | 12.1 (2)| 14.14   |
| June 2011      | 3.3 (6) | 10      |
| Average        | 7.8     | 14.4    |

3.3. Population Genetic Structure

Genetic analyses do not include the subsample for the year 2010 due to its limited size (only 7 individuals over the three sessions of the year). Annual subsamples were constituted by grouping individuals caught within the same year (2008, 2009 and 2011). When an animal was captured at several temporal sessions, it was referred to the sessions of its first capture.

Of the 66 exact tests performed to test for linkage disequilibrium, only 5 (7.5%) were significant after correction for multiple testing. Significant values systematically involved different pairs of loci (data not shown), thus suggesting that the 12 loci can reasonably be considered as independent.

The Gangara population was not at Hardy-Weinberg equilibrium and exhibited significant heterozygote deficiencies, as indicated by the mean positive \( F_{IS} \) value of 0.24 (95% CI = [0.11 – 0.36]). The allelic richness per population ranged from 2.18 to 13.34 for annual subsamples, with an overall average of 7.34 (Table 3). Median values of Loiselle’s kinship coefficients were globally low for each annual subsample (0.0062 for 2008, 0.0026 for 2009, and 0.0011 for 2011), and their distribution is largely unimodal indicating that there is no pairs of individuals which are more related than others (data not shown).

Table 3. Genetic diversity parameters estimated on the whole sample and the annual subsamples 2008, 2009 and 2011. N = Number of individuals; \( A_r \) = Number of alleles; \( H_e \) and \( H_o \) respectively expected and observed heterozygosités; HWE (p) = probability of HWE deviation test.

| Sample | N  | \( A_r \) | \( H_o \) | \( H_e \) | \( F_{IS} \) |
|--------|----|-----------|-----------|-----------|------------|
| 2008   | 28 | 6.751     | 0.5062    | 0.6663    | 0.25       |
| 2009   | 64 | 6.996     | 0.5041    | 0.6634    | 0.25       |
| 2011   | 35 | 6.821     | 0.5392    | 0.6569    | 0.19       |
| All    | 134| 7.340     | 0.508     | 0.686     | 0.24       |

Estimated pairwise \( F_{ST} \) values between annual subsamples.
ranged from 0.0056 to 0.0191 (mean $F_{ST} = 0.042$, 95% CI = [0.11 – 0.36]). They indicated small, but significant variations in allelic frequencies between annual sub-samples. IBD was significant ($P < 0.001$; $R^2 = 0.038$; regression slope $b = 0.014$, 95% CI [0.0078-0.0255]; Figure 2).

Using the regression slope value, and a density of 0.0016 individual /m$^2$ corresponding to the mean density observed in the quadrat, we estimated the mean parent-offspring genetic distances to be 59.5 m.

Figure 2. Results of Isolation by Distance (IBD) using Mantel tests correlating log-transformed geographical distances and genetic distances.

$F_{ST}$ values were not significantly different ($p = 0.77$) between males (mean $F_{ST} = 0.0115$) and females (mean $F_{ST} = 0.0036$). Spatial autocorrelations also indicated similar dispersal patterns for females and males (Figure 3). Effective dispersal distance estimated were of 54 m (95% CI: [38 m-90 m]) and 60 m (95% CI: [39 m-85 m]), for females and males, respectively, and the overlapping confidence intervals suggested that there is no significant difference between sexes.

4. Discussion

4.1. Population Dynamics and Genetic Diversity of $G. nigeriae$

Our study was conducted at a spatial scale (2.25 ha) somewhat smaller than those at which population dynamics and/or genetic surveys are usually carried out in rodents [33-38]. Nevertheless, we believe that the grid here considered is well representative of $G. nigeriae$ habitat conditions in this area, where millet crops and open savanna on sandy soils dominate in the landscape. Its location has been chosen based on evidence of $G. nigeriae$ presence at the site, and further confirmed by trapping investigation prior to the installation of the trapping grid.

Throughout the monitoring period, the recorded numbers of individuals, and consequently population densities, were very low. Limited “trappability” associated with behavioural characteristics (trap avoidance/shyness, social learning…) or unattractiveness of our traps could be put forward as explicative factors for the relatively low number of capture events recorded over the entire study despite an significant trapping effort. Another explanation may reside in our choice of 10m as inter-trap interval that is slightly larger than the average DRS retrieved from this study (7.8 m). Indeed, in theory, trap interval should be less than or equal to the DRS, so that the trapping device allows each individual to encounter at least one trap when foraging in its home range [39]. Nevertheless, much higher densities of $G. nigeriae$ were successfully detected in Kollo, western Niger, using a similar protocol [3]. In addition, despite higher trapping success values, the DRS obtained in Kollo (7.7 m) and in this study were similar. The low trapping success in Gangara (< 2%) suggests that gerbils were globally rare on our experimental field. Search and excavations of apparently active burrows in the area surrounding our grid (at least 30 burrows excavated per trapping session) largely confirmed the scarcity of gerbils in the area (data not shown). Altogether, these data bring strong support that our protocol enables a reliable reflection of demographic patterns in $G. nigeriae$ from Gangara.
As noted earlier, densities of *Gerbillus nigeriae* are low in the study area. The maximal density value recorded throughout the three and a half year long monitoring period (27.5 gerbils per hectare) was much [less than what was observed in Kollo between 1997 and 2000 (peaks of 120 individuals/ha [4])]. Taking into account the very similar protocols used, such density differences between Kollo and Gangara populations can be attributed to geographical (south west vs. central east Niger) and/or historical differences (1997-2000 in Kollo vs. 2008-2011 in Gangara). Kollo is located within a typical Sahelian region with an annual rainfall of 500 mm, while Gangara is located further north, between the Sahara and Sahel ecoregions, and receives an average of 350 mm annually [40] (DMN, 2014). Moreover, Gangara has been more affected by the anthropic and climatic changes than the southernmost zones, with isohyets moving from north to south since the early 1970s and resulting in rarer and more erratic rainfall [9, 41]. Such a rapid aridification and subsequent degradation of local habitats have induced a loss of biodiversity in northern Sahel [42]. As an example, many plant species have gone extinct locally, probably because they had reached their drought tolerance threshold [43]. Like these plants, *G. nigeriae*, which is not adapted to Saharan conditions [44, 1, 2], may also have suffered from recent aridification processes at work along the northern edge of its geographic distribution, such as in the region of Gangara. This impact of climatic and environmental conditions on the evolution of *G. nigeriae* species range may further be exemplified by its occurrence in Southern/Sudanian localities of Niger [12] where it was not known by the inhabitants until recently (Hima, unpublished). Yet, the genetic diversity observed in *G. nigeriae* from Gangara was higher than that found in the expansion area of the species (e.g. in Senegal, Ndiaye et al. unpublished), and similar to that observed in large-scaled studies focusing on other African rodent species (e.g. in *Mastomys* [33, 45-47] or *Praomys* [48]). If low densities of *G. nigeriae* in Gangara are attributed to climatic and environmental changes, such processes are probably too recent to be reflected by local genetic diversity levels. Unfortunately, no genetic data are available for Kollo’s population, which would have helpful in terms of evaluating whether genetic diversity may reflect the contrasted density levels observed in Gangara and Kollo.

Our results indicate that the *G. nigeriae* population of Gangara presents a single-phase cycle characterized by important inter- and intra-annual variations. Density peaks appeared towards the end of the dry season, or sometimes even at the beginning of the rainy season. This cycle is closely related to duration and intensity of the rainy season with heavy rainfall on year N leading to earlier and more important reproduction in year N+1. Similar results were found in another gerbilline rodent species, namely *Taterillus pygargus*, in semi-arid regions of Senegal [49]. A population dynamics pattern with a unique peak of density at the beginning of the rainy season was also noted in *G. nigeriae* from Northern Burkina Faso [8]. In contrast, in Kollo...
(southwestern Niger), *G. nigeriae* abundance cycle seems biphasic and unbalanced, including two distinct periods of abundance with varying intensities and durations [3]. The first, less marked and shorter, corresponds to the end of the dry season / beginning of the rainy season, while the second one, characterized by higher densities over longer periods, is observed between November and February. Such a pattern showing two maxima and two minima is generally typical of rodents from tropical regions characterized by two rainy seasons [50, 51]. In our study, however, the time period that separated consecutive sessions (4 months) may have been too long to highlight a biphasic cycle. Indeed, when comparing the results obtained in Southwestern (Kollo) and central east (Gangara) Niger at the only periods when trapping occurred in Gangara (February, June and November), the abundance cycles of the Kollo and Gangara populations appear similar, suggesting that the cycle could also be biphasic in Gangara. Abundance cycles, if they exist, may contribute to explain the significant heterozygote deficiencies observed if they lead to genetically different cohorts through the year [52]. However, estimated pairwise $F_{ST}$ values between annual subsamples were very low, indicating weak differentiation between annual subpopulations, hence between generations.

The weak low abundance of *G. nigeriae* in eastern Niger during some periods may be due to a possible estivation of gerbils as already observed in northern Burkina Faso [53, 54]. No estivation was mentioned in *G. nigeriae* from Kollo [3], but contrasted aridity levels could lead to differences in life cycle between populations. In support of the estivation hypothesis in *G. nigeriae* from Gangara, important quantities of cultivated and wild grass seeds (e.g. *Pennisetum* spp., *Sorghum* spp., *Cenchrus biflorus*, *Cymbopogon gayanus*) were found stored in the numerous galleries of the excavated burrows. Food storage inside burrows may also partly explain our low trapping success, via a reduction of the mobility of gerbils rather feeding on stocked food than attracted by the bait.

### 4.2. Mobility and Dispersal Capacities

Dispersal can be estimated either directly by CMR, or indirectly using genetic markers for inferring differentiation between individuals in both space and time [55, 56]. Here, both approaches were used and found to be quite congruent in their conclusions. First, our CMR results show that gerbils from central-east Niger are not very mobile with DRS values ranging from 3.3 to 12.1 m (average 7.8m) and DMR values ranging from 10 to 28.3 m (average 14.4m). The individuals thus seems to live on and explore only very small home range. This is perfect agreement with what was observed in south-west Niger (average DRS of 7.7 m and average DMR of 24.8 m) despite higher densities and a slightly different annual cycle of abundance [3]. These small home ranges may result in spatial structure, and explain the heterozygotes deficiencies observed, as already shown for instance in desert rodents [57]. The significant IBD pattern detected in Gangara confirms the limited dispersal abilities of *G. nigeriae* at the scale of the trapping grid. Indeed, significant IBD is usually associated with low individual dispersion capacities, especially when individuals are genealogically and geographically close, i.e. on a reduced spatio-temporal scale [58, 59]. We showed here that dispersal is limited for both females and males, although the latter is usually the most dispersing sex in mammals [60]. Low dispersal abilities should theoretically result in high inbreeding. At the scale of the quadrat, the unimodal distribution of kinship coefficients did not enable to reconstruct different family groups. Further investigations would be required to better understand spatial structure and mate choice strategies of this species with limited dispersal abilities but relatively high genetic diversity levels.

In short, *G. nigeriae* population dynamics results are very consistent wherever they are obtained in Kollo [4] or Gangara (this study) which are characterized by two very distinct chromosomal variants [2]. They show that the species may display different demographic patterns in the two areas, but share the same low mobility. They highlight that, whatever the cytotype, *G. nigeriae* has one of the lowest mobility ever documented in sudano-sahelian rodent species (e.g. average DRS values of *G. gerbillus*, *G. nanus*, *Mastomys erythroleucus*, *Taterillus gracilis* and *Gerbilliscus guineae* are of 10.7, 16.9, 17.7, 19.5 and 20.5 m, respectively [61, 1]). In contrast with its low dispersal abilities in Niger, the expansion of *G. nigeriae* in Northern Senegal seems to be very rapid, probably as a result of the opening environment and increased aridity driven by anthropo-climatic changes ([62]; Granjon et al. unpublished). The reason for such a difference remains to be investigated.

### 4.3. Implications for *G. Nigeriae* Control as a Rodent Pest

In the central-east region of Niger, *G. nigeriae* may be at the limit of its distribution range. As aridity increases, the local persistence of the species may become difficult, and gradual extinction may occur. No evidence exist about a potential distribution shift from north to south in Niger, as seems to be the case between Mauritania and North Senegal [62]. However, if this was to be the case in Niger too, the impacts in terms of damage to crops would be probably significant since the southern regions are currently the main cereal production areas in the country.

In view of the low mobility and limited-dispersal capacities of *G. nigeriae* in Niger, locally-designed (i.e. at the field or village scales) control strategies against this rodent pest appear conceivable. Such rodent control campaigns could be carried out during low population’s densities periods which are now well characterized whatever the chromosomal pool (i.e. high vs. low diploid numbers) / geographical region (west vs. east of Niger) considered. Specifically, the control should be conducted between January and May when densities are low and when farming activities are at their lowest. Nevertheless, before launching any control activities, it would first be necessary to evaluate its cost-benefit ratio [63]. Indeed, it may happen that, below a certain threshold of pest density, the damage caused may be less costly than the control procedure itself.
Interestingly, traditional traps known as “Kornaka” traps (which are quite similar to pitfall traps) could be used for controlling rodent populations in Niger. Indeed, a preliminary test of these traps showed that they could significantly reduce gerbil densities (37 captures in 3 night-traps with 49 “Kornaka” traps versus 22 captures in 5 night-traps with 256 wire mesh traps; Hima et al. unpublished). As such, “Kornaka” traps may be an effective, cheap and accessible alternative for the control of pest gerbils by vulnerable Sahelian farmers.

5. Conclusion

Our results showed that in the Central Sahel, *G. nigeriae* populations are characterized by very low population densities with a single-phased abundance cycle which was strongly dependent on rainfall patterns. We also found that mobility parameters indicated very low individual mobility. In addition, genetic analyses revealed spatial structuration, thus confirming rather poor dispersal capacity. So, in terms of control, our results suggested that, locally-designed control strategies appear conceivable, in view of the low mobility and limited-dispersal capacities of *G. nigeriae* in Niger. Specifically, control measures should be conducted between January and May when densities are low and when farming activities are at their lowest. Interestingly, traditional traps known as “Kornaka” traps could be an effective, cheap and accessible alternative for this control by vulnerable Sahelian farmers.

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