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Small mammal community demography and reproductive seasonality under protection, pastoralism and agriculture in the Serengeti Ecosystem, Tanzania

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This work examined population demography and reproductive seasonality among small mammals under protection, pastoralism and agriculture in the Serengeti Ecosystem in 2017 and 2018. Small mammals were trapped to quantify variation in the proportions of actively breeding individuals, juvenile recruitment and adult sex ratio across land use types and seasons. Breeding peaked in the wet season, reflecting a corresponding underlying peak in resource availability and quality. The mean community proportion of reproductively active individuals was greater for females (60%) than males (40%) and under pastoralism than protection or agriculture. Juvenile recruitment was the highest in the pastoral land in the dry season. Adult sex ratio was biased towards males across all land use types and seasons but towards females in the pastoral than the agricultural land. This concurs with the Trivers-Willard hypothesis that nutritionally stressed females should produce more female offspring. Two generalist and pest species (*Mastomys natalensis* and *Arvicanthis niloticus*) dominated the pastoral areas indicating human disturbance. Anthropogenic activities degrade habitats, altering population demographics and promoting habitat generalists and pest species. Protection promotes habitat intactness and reduces opportunistic pests typically abundant in disturbed landscapes. Consequently, enhancing conservation around protected areas can improve diversity and control the abundance of opportunistic pest species.

Key words: Small mammals, seasonal breeding, juvenile recruitment, adult sex ratio, Serengeti ecosystem, pastoralism, agriculture, protection, rainfall, land use.

INTRODUCTION

Seasonal breeding and population demography are important components of animal population dynamics but can be altered by anthropogenic land use changes that

cause habitat degradation and loss (Blaum et al., 2007; Cao et al., 2016). Human activities often replace natural vegetation with managed systems with altered and often

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simplified structure, posing threats to biodiversity (Rickart et al., 2013; Schmidt et al., 2005). Such activities therefore threaten biodiversity conservation by reducing the geographic ranges and population sizes of many species (Newbold et al., 2015; Tittensor et al., 2014). Human activities can also adversely affect individual fitness and populations of many species, including small mammals (Bond et al., 2005; Boren et al., 1999; Galbraith, 1988; Hansen et al., 2019; Li et al., 2016; Peles and Barrett, 1996; Schieltz and Rubenstein, 2016; Schmidt et al., 2005). However, the precise consequences of human activities on wildlife vary among species, reflecting their contrasting adaptations to particular habitats and life history strategies, such as the degree of feeding specialization (generalists versus non-generalists) and timing of breeding.

Small mammals are some of the most important mammals contributing to ecosystem species diversity. Their community structure and species richness are related to variables such as habitat structure, rainfall and productivity (Avenant, 2003). Because land use often alters the availability and quality of essential resources, it can modify the timing and prolificacy of breeding in animals. By shaping food availability, nesting sites and other resources for small mammals, land use can alter their relative competitive abilities and reproductive seasonality (Newsome, 1969a, b). Alterations to natural habitats by land use can modify the constraints imposed by environmental structure and seasonality on the timing, synchrony and fecundity of breeding by small mammals. Even so, rainfall seasonality is a leading meteorological predictor of reproductive seasonality in small mammals in tropical savannas where photoperiod is absent, through its controlling influence on food and surface water availability and quality (Leirs et al., 2010; Lima et al., 2003; Madsen and Shine, 1999; Odhiambo et al., 2005). Rainfall, by modulating food resources, shapes the population abundance and reproductive phenology of small mammals (Bai et al., 2015; Lima et al., 2003; Makundi et al., 2007). As a result, small mammals typically breed seasonally in African savannas with strong rainfall seasonality. For many small mammal species, breeding occurs year-round but with peaks in the transition period between two seasons such that breeding peaks during the early wet season and reduces during the dry season (Francisco et al., 1995; Lima et al., 2003; Wube, 2005).

Seasonal reproduction leads to seasonal recruitment of new-borns and peaks in the number of juveniles at the end of the breeding season (Bantihun and Bekele, 2015). Thus, although juveniles can be found in the population throughout the year, their numbers often peak at the end of the wet season and during the dry season (Gebresilassie et al., 2006; Olenev and Grigorkina, 2011; Oli and Dobson, 1999). Abundant fresh vegetation during the wet season provides small mammals with adequate high-quality food, cover from predators and water for

lactating females (Linn, 1991; Massawe et al., 2005).

Reproductive seasonality in mammals is often associated with changes in population demographic characteristics, such as sex ratio. Sex ratio is an important demographic parameter underpinning population fluctuations in mammals (Oli and Dobson, 1999) and is balanced for some populations or population segments but unbalanced for others (Bantihun and Bekele, 2015; Mulungu et al., 2013). At birth, mammals produce approximately equal numbers of males and females resulting in a balanced sex ratio (Rosenfeld and Roberts, 2004), anthropogenic activities can, however, strongly bias sex ratios with significant implications for population dynamics (Marealle et al., 2010; Ndbalema, 2009; Setsaas et al., 2007). Human activities can also indirectly skew sex ratios in mammals by influencing maternal body condition, type and nutritional sufficiency of the diet, competitive ability and stress levels (Hoffmann et al., 2003; Hunnink et al., 2018; Li et al., 2016; Rosenfeld and Roberts, 2004; Trivers and Willard, 1973). Biased sex ratio can have manifold population-level consequences, which can be sex specific and include heightened competition for mates (Fritzsche et al., 2016; Kvarnemo and Ahnesjo, 1996; Schärer et al., 2012), elevated risk of population decline and local extirpation (Le Galliard et al., 2005; Rankin et al., 2011) and breeding depression (Brook et al., 2002; Stephens and Sutherland, 1999).

Although small mammals have been widely studied in many ecosystems, including the Serengeti (Magige and Senzota, 2006; Makundi et al., 2007; Stanley et al., 2011), the variation in their breeding and population demography (abundance, age and sex ratios and juvenile recruitment) across seasons and land use types has attracted relatively little attention. Here, we investigate the influence of seasonality and land use on the breeding activity and population demography of small mammals in the Serengeti Ecosystem. Specifically, we examine seasonal variation in 1) relative abundance of actively breeding females and males, 2) juvenile recruitment among three contrasting land use types and 3) adult sex ratio.

We test predictions of the following seven hypotheses: (H1) The mean breeding probability for males and females should be higher for the wet than the dry season because the wet season receives higher rainfall and thus has higher resource availability and quality for small mammals (Kelly et al., 2013; Sarli et al., 2016; Taylor and Green, 1976). (H2) Further, the mean breeding probability should be higher for males than females in both seasons and in all the three land use types because females spend much more time on reproduction (gestation and rearing young) than males (Johnson et al., 2001). (H3) Because human activities such as cultivation and livestock grazing disturb small mammals, breeding probability should be higher in the protected area than in either the agricultural or the pastoral land. (H4) For the

same reason, juvenile recruitment is expected to be higher in the protected area than the agricultural or pastoral land. (H5) Juvenile recruitment should also be higher in the dry than the wet season if breeding peaks in the wet season. (H6) Adult sex ratio should be skewed in favour of females in the wet season but in favour of males in the dry season because female small mammals aggressively defend territories in the wet season and the majority of young females are philopatric whereas adult and juvenile males disperse from their natal home ranges to search for mates and food elsewhere (Barrett and Feldhamer, 2008). Furthermore, the local resource competition theory predicts that females should preferentially produce female offspring when resources are plentiful but male offspring when resources are limiting to minimize competition (Clark, 1978; Twining et al., 2017). (H7) Thus, males should be more abundant in the pastoral land due to relatively lower food availability caused by intense livestock grazing but females should be more abundant in the agricultural land and protected area, with potentially higher food availabilities.

MATERIALS AND METHODS

Study area

Data on small mammals were collected on the North-eastern part of the Serengeti ecosystem, encompassing the Serengeti National Park (2° 20' S, 34° 50' E) and the adjoining Serengeti (2°15' S, 34°68' E) and Ngorongoro (3°24' S, 35° 48' E) Districts. Serengeti National Park protects about 15,000 km² of tropical savanna ecosystem (Sinclair, 2008). The park encompasses a wide variety of vegetation types, including woodlands, treed grasslands, open grasslands and other more restricted habitat types (Byrom et al., 2014; Reed et al., 2009). Farming and livestock pastoralism are practiced in the vicinity of the protected area. Rainfall is bimodal and peaks in the short (November-January) and long (March-May) rainy seasons (Norton-Griffiths et al., 1975). There is a strong gradient of increasing rainfall from the dry south-eastern plains (700 mm/year) to the wet north-western woodlands (1,050 mm/year) inside the Serengeti National Park. The total monthly rainfall during the study period (2017-2018) averaged 100 mm for the wet and 62 mm for the dry season. During the same period, the daily temperature averaged 26°C in the wet and 24°C in the dry season.

Study design

The study was carried out along the Mto Wa Mbu-Musoma road transect (Serengeti northern road), which traverses protected, agricultural and pastoral land use types. The transect was selected because it crosses agricultural lands (north-west), pastoral lands with limited agriculture (north-east) and the protected Serengeti National Park situated between these two land use types (Figure 1). Each of the three land use types was subdivided into four plots, for a total of 12 plots ("habitats"). Out of the 12 habitats (plots), 10 were selected for trapping small mammals. The selected 10 habitats were categorized into five habitat types and included four habitat types (riverine forest, shrubland, wooded grassland and grassland) in the Serengeti National Park, four habitat types (shrubland, cropland, grassland and wooded grassland) in the agricultural lands and two habitat types (cropland and shrubland) in the pastoral

lands. A noteworthy difference in farming systems between the pastoral and agricultural lands is that the fallow period is relatively longer in the pastoral than the agricultural land. In this study, maize and bean croplands represented cropland habitats.

Trapping procedures

Traps were set within a 100 × 100 m sub-plot in each of the 10 plots (habitats). A total of 141 traps (100 Sherman, 30 wire mesh and 11 bucket pitfall traps) were set in each sub-plot for five consecutive nights and then transferred to the next sub-plot. The wire mesh traps are funnel-shaped, multi-capture traps made of thin wire and are widely used in Tanzania by local hunters. They have one door which allows an animal to enter but not to leave the trap because of inward projecting wires that form an effective barrier to exit.

Each sub-plot was assigned one pitfall line consisting of 11 buckets, spaced 5 m apart, and buried into the ground such that the top of the bucket was at the ground level. Each of the 11 buckets per plot was buried 26 cm deep and had a 15 L capacity, upper and lower diameters of 26 and 24 cm, respectively. Each pitfall line had 50 cm high black plastic drift fence running over the center of each bucket. These non-baited traps captured animals moving on the habitat floor that encountered the drift fence after following the trail and falling into a bucket. A similar procedure has been used with considerable success in other small mammal surveys (Byrom et al., 2014; Nkwabi et al., 2018; Stanley et al., 2011). For the Sherman traps (23 × 9.5 × 8 cm), 10 lines spaced 10 m apart were set up in the grid. Sherman traps were arranged along the lines at 10 m intervals for a total of 100 traps per sub-plot. To maximize capture of a wide variety of species; 30 wire mesh traps ('*mgono*') were placed in-between and after every Sherman trap line such that in every Sherman line five wire mesh traps were placed 20 m apart. Freshly fried coconut coated with peanut butter and mixed with Lake Victoria sardine (*Rastrineobola argentea*) was used as a bait and traps were rebaited twice daily, in the morning (6.30-8.30 am) and late afternoon (17:00-19:00 pm). All traps were checked twice daily, early in the morning (6.30-8.30 am) and early evening (17:00-19:00 pm). Both in 2017 and 2018, traps were laid out in April-May in the wet season and in August-September in the dry season. Traps stayed in one plot for five consecutive days. Using Kingdon (2015) and Kirsten et al. (2010) as guides, we identified trapped animals to the genus or species and recorded their morphometric (external shape and dimensions) measurements. Moreover, individually distinctive features, including sex, size, reproductive status and presence of scars were recorded and used to aid individual identification (Graham and Lambin, 2002; Kirsten et al., 2010).

Captured individuals were aged using body size, fur colour and texture. The animals were sub-divided into juvenile and adult age classes. Juveniles and subadults have smaller body size, greyer, softer and down-like baby fur, and typically have not undergone the changes associated with adulthood such as reproductive development (Searle, 1985). By contrast, adults have larger body size, different fur colour patterns and texture and fully developed reproductive organs (Kingdon, 2015; Searle, 1985). All adults and sub-adults were amalgamated together as "adults" whereas all young individuals and juveniles were lumped together as "juveniles" to minimize the likelihood of misclassification. Individuals were sexed using external genitalia and secondary sexual characteristics such as the presence of testes, status of nipples, number of urogenital openings (three in shrews) and distance between the anus and urogenital opening, which is relatively shorter for female than male rodents (Carraway, 2009; Kunz et al., 1996). To ascertain whether an animal was actively breeding, we noted the breeding status of every individual by examining the position of the testes; scrotal or abdominal in males and perforated or imperforated vagina in females. During the breeding season, males have descended testes caused by a temporary bulge in the perineal

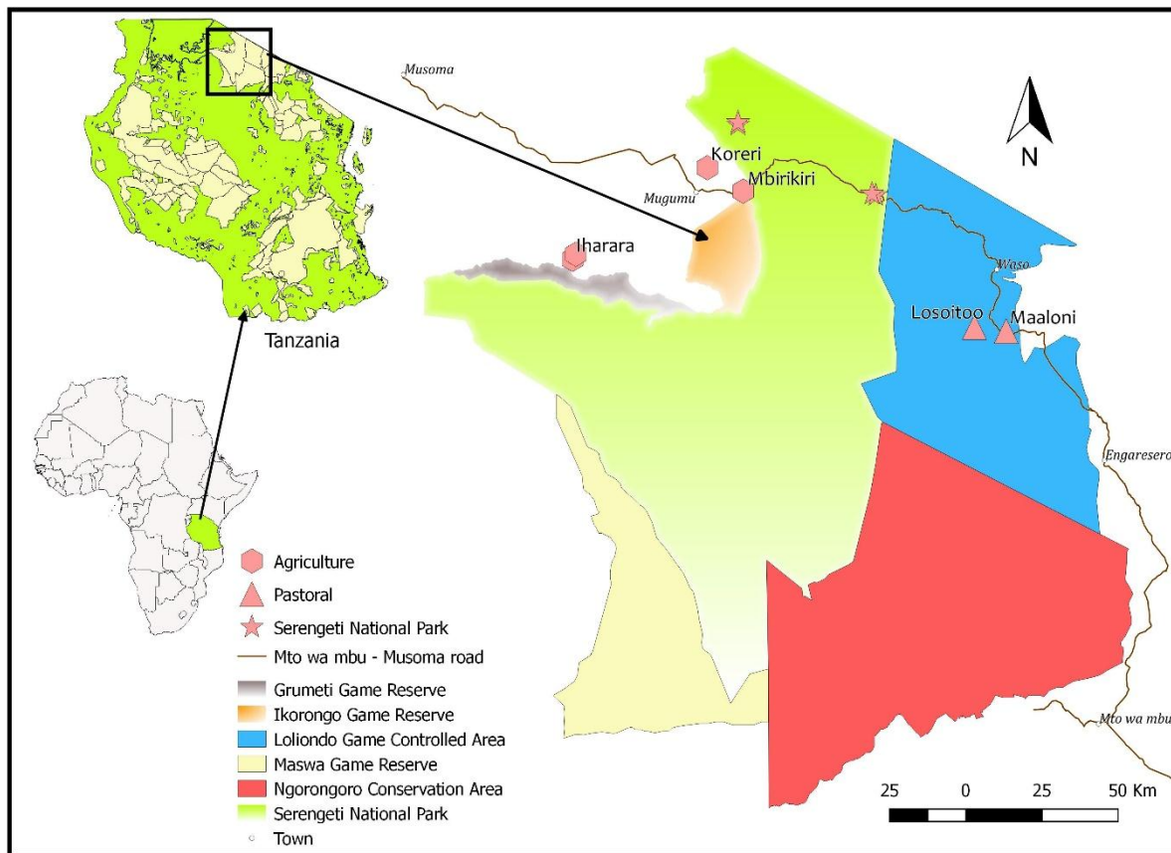


Figure 1. Map of the Serengeti Ecosystem showing the study area including the Mto Wa Mbu – Musoma road transect (study plots are in pink colour).

region (between the anus and the urinogenital opening) whereas females have a swollen vaginal tissue and gaping vaginal opening. Animals whose sex and breeding status could not be accurately determined were classed as “undefined” and excluded from the statistical analyses involving either trait. In particular, juveniles were hard to sex accurately and hence were omitted from all analyses involving sex ratios. As a result, only “adults” (adults and sub-adults), which were accurately identified to sex ($n = 552$ of 566 adults), were included in analyses involving information on the sex of captured individuals. Similar methods have been used previously to sex, age and ascertain the breeding status of small mammals (Makundi et al., 2007; Monadjem and Perrin, 2003).

Statistical analyses

We used a generalized linear model with a binomial error distribution and logit link function (logistic regression) to analyse the variation in the community proportion of all actively breeding small mammals (number of breeding adults of all species/number of all adults of all species) with season, land use, sex and all their interactions. We used the logarithm of the total trap nights per land use type as an offset to account for variation in total trap nights across the three land use types. We calculated adjusted mean breeding probabilities for all significant main effects and interactions. We also performed a partitioned analysis of the adjusted mean breeding probabilities for the significant interaction effects (analysis of simple effects) (Winer, 1971), carried out

pairwise comparisons of the adjusted mean probabilities and adjusted the family wise error rate for multiplicity by using simulation adjustment (S1 Data). We similarly analysed variation in juvenile recruitment (juveniles/(juveniles + adults, S2 Data) across seasons, land use and their interaction and variation in adult male: female ratio (Males+Females) across land use types and seasons (Becker et al., 2016). The logistic regression models were implemented in the SAS GLIMMIX procedure (SAS Institute 2019, SAS/STAT Version 15.1).

RESULTS

A total of 612 small mammals consisting of 18 rodent species ($n = 528$, 86%) and one genus of shrews ($n = 84$, 14%) were trapped and recorded during 28,200 trap nights of effort. Adults constituted 92.5% whereas juveniles 7.5% of all the 612 captured small mammals. Of the 566 adults, 65.7% were males, 31.8% were females; whereas 2.5% were not identified to sex. Four species dominated the captured small mammals, namely *Arvicanthis niloticus* ($n = 162$), *Mastomys natalensis* ($n = 101$), *Mus* sp ($n = 91$) and *Crocidura* sp ($n = 84$). The two most abundant species (*A. niloticus* and *M. natalensis*) were particularly abundant on the pastoral landscape

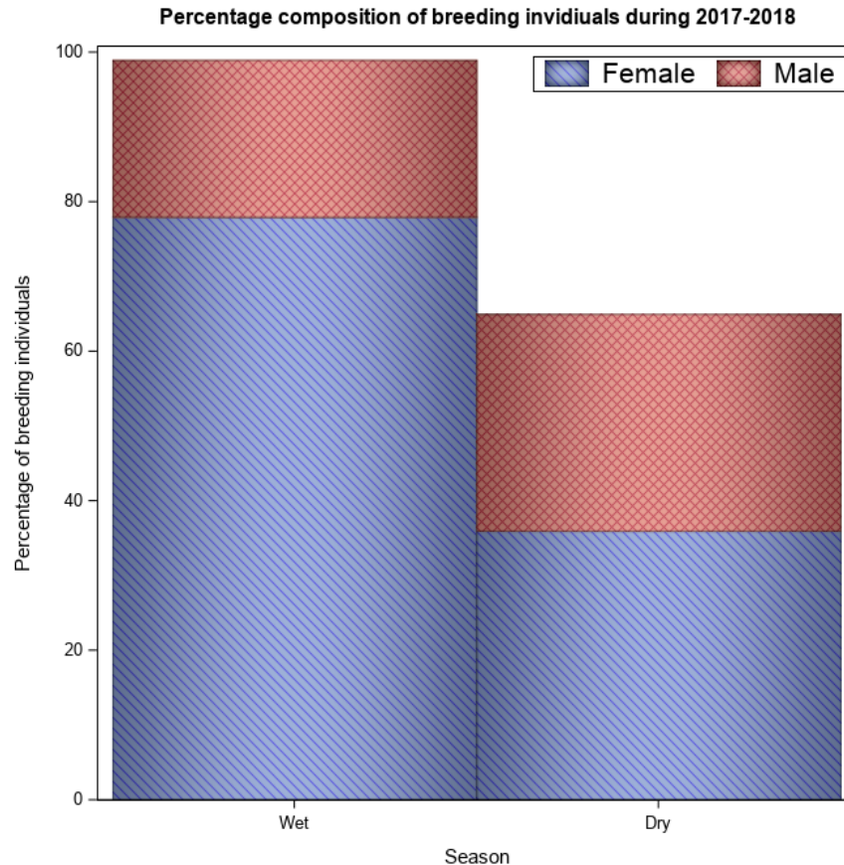


Figure 2. Percentage composition of the actively breeding female and male small mammals captured in the Serengeti ecosystem during the wet and dry seasons of 2017 and 2018.

where they contributed 73% of all the 96 actively breeding individuals captured.

Variation in breeding activity across sexes, seasons and land use types

Out of the 566 adults trapped, 167 (47%) were actively breeding and consisted of 67 (40%) males and 100 (59.8%) females. The mean breeding probability was higher for the wet than the dry season (Figure 2) because substantially more males and females were engaged in breeding activity in the wet than the dry season (Tables 1-3) as predicted (H1). As well, more females than males were engaged in breeding activity in both the wet and dry seasons (Tables 1 to 3 and Table S1), contrary to the prediction (H2) that the mean breeding probability should be higher for males than females regardless of land use or season.

In addition, the mean breeding probability differed across land use types but was surprisingly the highest for the pastoral, intermediate for the agricultural and lowest for the protected land (Figure 3 and Tables 1 and 2), contrary to the prediction that it should be the highest for

the protected area (H3). The mean breeding probability for each land use type was, moreover, similar for both males and females and for the wet and dry seasons (Table 1).

Variation in juvenile recruitment across seasons and land use types

The mean juvenile recruitment was higher for the dry (10.5%) than the wet season (3.2%, $n = 612$; Tables 4 and 5, Table S2) as predicted (H5). The mean juvenile recruitment was also higher for the pastoral land than for either the protected or the agricultural land, both of which had comparable mean recruitment probabilities (Tables 4 and 5, Table S2); a pattern inconsistent with the prediction that it should be the highest for the protected area (H4).

Variation in adult sex ratio across seasons and land use types

Sex ratio was biased in favour of males for both the wet

Table 1. Tests of fixed effects of land use, season, sex and their interactions on variation in breeding probability of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	NDF	DDF	F	Pr > F
Landuse	2	11	20.46	0.0002
Season	1	11	23.63	0.0005
Landuse×Season	2	11	0.94	0.4203
Sex	1	11	33.16	0.0001
Landuse×Sex	2	11	1.06	0.3794
Season×Sex	1	11	7.22	0.0211
Landuse×Season×Sex	2	11	0.06	0.9466

NDF and DDF are the numerator and denominator degrees of freedom, respectively.

Table 2. Adjusted mean breeding probability and the associated 95% confidence limits (CL) for small mammals trapped in the Serengeti Ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	Landuse	Season	Sex	Mean probability	Standard error	Lower 95% CL	Upper 95% CL
Landuse	Agricultural			0.3441	0.08543	0.1857	0.5469
Landuse	National park			0.2493	0.04076	0.1705	0.3491
Landuse	Pastoral			0.5375	0.05482	0.417	0.6538
Season×Sex		Dry	F	0.3049	0.0608	0.1891	0.4519
Season×Sex		Dry	M	0.1387	0.02709	0.0891	0.2097
Season×Sex		Wet	F	0.8372	0.06796	0.6318	0.9391
Season×Sex		Wet	M	0.2465	0.03859	0.1716	0.3408

F and M denote females and males, respectively.

Table 3. Tests of simple effect slices (decomposition or partitioned analysis) of the interaction between season and sex on variation in breeding probability of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	Slice	NDF	DDF	F	Pr > F
Season×Sex	Sex F	1	11	18.31	0.0013
Season×Sex	Sex M	1	11	5.31	0.0417
Season×Sex	Season Dry	1	11	7.51	0.0192
Season×Sex	Season Wet	1	11	26.01	0.0003

NDF and DDF are the numerator and denominator degrees of freedom, respectively.

and dry seasons, the protected area and agricultural land but was approximately balanced for the pastoral land (Figure 4, Table 6 and Table S3 and S4). These patterns contradict the prediction that females should be more abundant than males in the wet season but less abundant than males in the dry season (H6) as well as the prediction that males should be more abundant than females in the pastoral land (H7). Furthermore, pairwise comparisons showed that male sex ratio was higher for the agricultural than the pastoral land but comparable between the agricultural and the protected land. But female sex ratio was higher for the pastoral than the

agricultural land but comparable between the pastoral versus the protected land and the agricultural versus the protected land (Table S4).

DISCUSSION

We examined variation in breeding activity and community demography of small mammals with season and land use in the Serengeti Ecosystem during 2017 and 2018. This involved quantifying seasonal variation in the relative abundance of actively breeding females and males, adult sex ratio and juvenile recruitment among

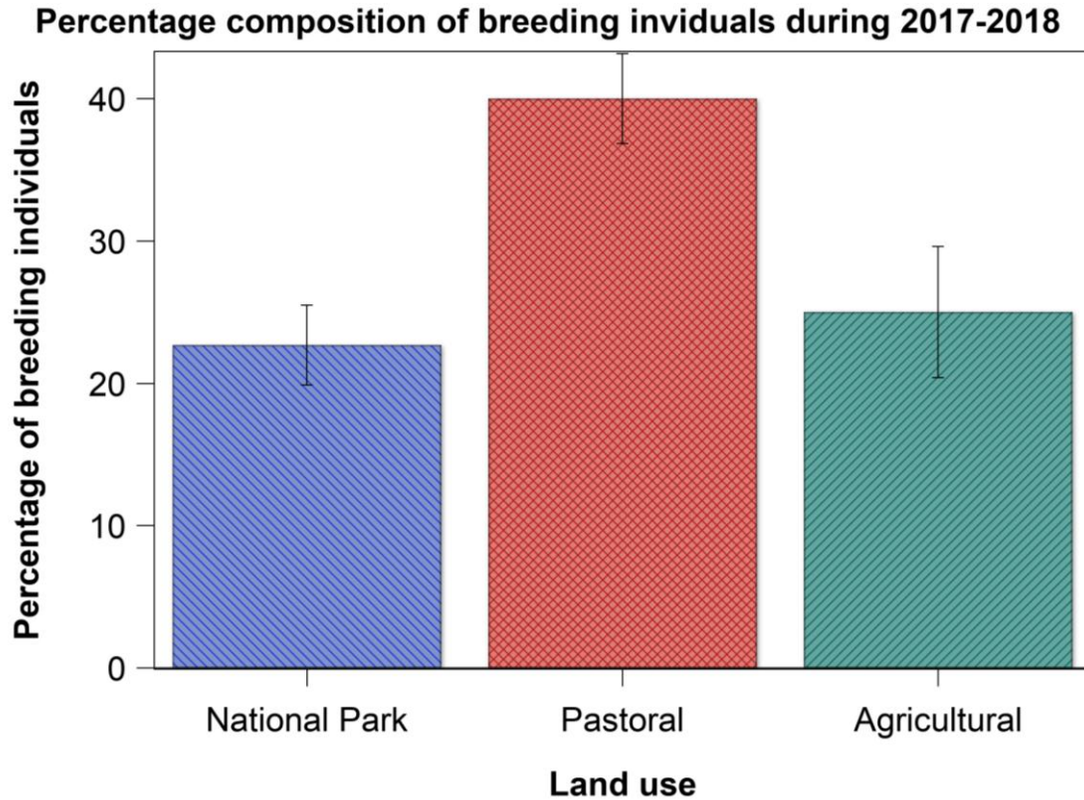


Figure 3. Percentage and the associated standard deviation of the actively breeding individuals captured on three land use types in the Serengeti Ecosystem during 2017 and 2018.

Table 4. Tests of fixed effects of land use and season (the interaction effect between the two factors was dropped because it was insignificant), on variation in juvenile recruitment of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model. NDF and DDF are the numerator and denominator degrees of freedom, respectively.

Effect	NDF	DDF	F Value	Pr > F
Landuse	2	8	13.72	0.0026
Season	1	8	6.05	0.0393

Table 5. Adjusted mean recruitment probabilities, standard errors and 95% confidence limits (CL) for small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	Landuse	Season	Mean probability	Standard error	95% Lower CL	95% Upper CL
Landuse	Agricultural		0.049481	2.037×10^{-6}	0.018462	0.125943
Landuse	National park		0.023883	9.123×10^{-7}	0.009191	0.060625
Landuse	Pastoral		0.095993	4.344×10^{-6}	0.058714	0.153104
Season		Dry	0.117526	2.138×10^{-6}	0.076605	0.183297
Season		Wet	0.046776	1.309×10^{-6}	0.02044	0.104624

three contrasting land use types.

For this small mammal community, breeding activity was higher during the wet than the dry season. This

reflects the underlying peak in vegetation cover, drinking water and food availability and quality in the wet season, associated with higher rainfall.

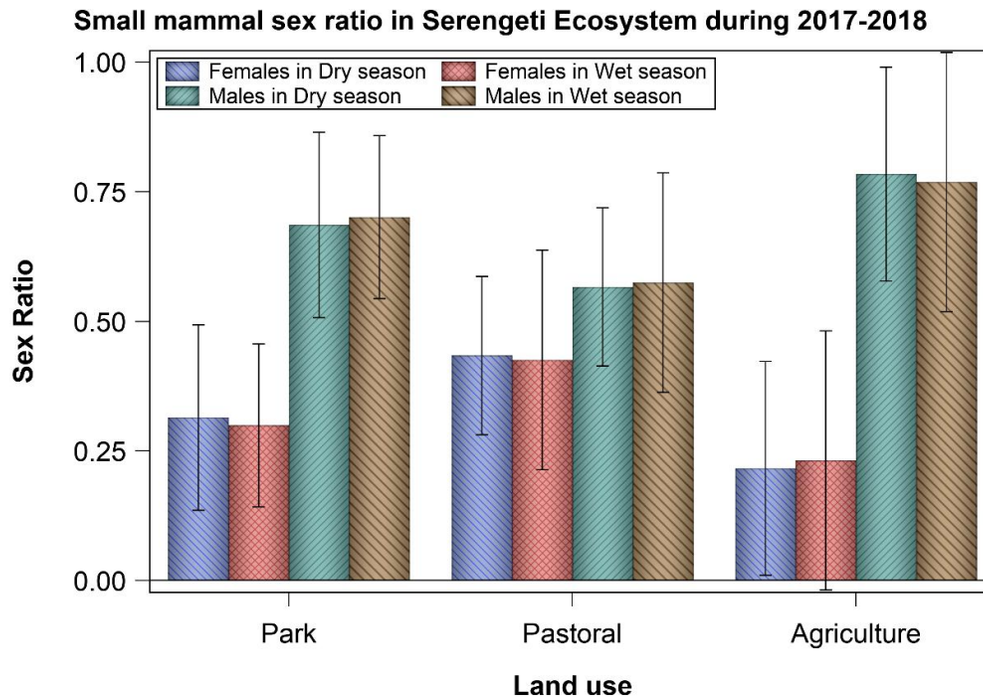


Figure 4. Small mammal sex ratio and the associated standard deviation across land use types and seasons in the Serengeti Ecosystem during 2017 and 2018.

Table 6. Tests of fixed effects of land use, season, sex and their interactions on variation in sex ratio of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	NDF	DDF	F Value	Pr > F
Landuse	2	2	0	1.0000
Season	1	2	0	1.0000
Landuse×Season	2	2	0	1.0000
Sex	1	2	133.44	0.0074
Landuse×Sex	2	2	21.7	0.0441
Season×Sex	1	2	0.18	0.7144

NDF and DDF are the numerator and denominator degrees of freedom, respectively.

The wet season breeding peak is consistent with expectation and findings of several earlier studies, suggesting that breeding in African small mammals peaks in the wet season with abundant and nutritionally sufficient forage for lactation and rearing young (Bantihun and Bekele, 2015; Makundi et al., 2007; Massawe et al., 2008, 2012; Nicolas and Colyn, 2003). Through its influence on the availability and nutritional sufficiency of forage, rainfall seasonality influences life history strategies and individual fitness of small mammals (Makundi et al., 2007). Rainfall seasonality governs reproductive success in small mammals by indirectly affecting their food supply, such as insects (Linzey and Kesner, 1997), or directly controlling the production of fresh, succulent vegetation and availability of adequate

water for lactation (Reichman and Van De Graaff, 1975; Soholt, 1977). Therefore, the small mammal species apparently time their reproduction so that the offspring are weaned at the time of peak resource abundance and quality.

There were more active breeding females than males across all seasons and land use types, thus contradicting the prediction that there should be more breeding males than females. The preponderance of females implies that potentially more juveniles would be recruited into the community if many females survive to successfully mate and reproduce. The number of actively breeding females has been reported to peak at the onset of the breeding season when females search for potential mates but drop from the middle to the end of the breeding season as the

majority of the females are either pregnant or taking care of young ones (Geary, 2015). Therefore, the larger number of breeding females relative to males in the breeding season suggests that the time of capture likely coincided with the beginning of the breeding season for most of the small mammal species. Alternatively, the short gestation period and postpartum oestrus of some of the small mammal species may have resulted in the presence of many actively breeding females in the population across seasons and land use types.

The skew in operational sex ratio (the sex ratio of ready-to-mate or sexually active individuals) towards females among the sexually active individuals suggests that females were probably competing for male mates whereas the males increased their mating input. This follows from the expectation that competition for mates is likely to occur when operational sex ratio is skewed in favour of one sex (Geary, 2015; Zhang and Zhang, 2003). Specifically, when operational sex ratio is male skewed, then males may compete for mates, but when it is female skewed, then males tend to increase their seasonal mating input (Zhang and Zhang, 2003). However, since we focus on breeding at the community level, further data would be required to establish whether operational sex ratio is similarly biased in favour of females for breeding populations of the individual species.

In the breeding (wet) season, the mean breeding probability of the small mammal community varied across land use types and was higher for the pastoral than the protected land. This contradicts the expectation that this probability should be the highest for the protected land. The relatively large proportion of breeding individuals in the pastoral land is due to the preponderance of *A. niloticus* and *M. natalensis* there. These two species are common in human-disturbed habitats and often numerically dominate other sympatric species (Avenant, 2003; Byrom et al., 2015; Crespín et al., 2008; Magige, 2016). Their numerical dominance in the pastoral land likely reflects disturbance by livestock grazing that creates conditions they favour. Furthermore, since *A. niloticus* and *M. natalensis* species are serious pests in most parts of Africa, their numerical dominance in the shrubland and cropland habitats in the pastoral land implies that habitat conversion by human activities tends to favour pest species. As a result, habitat conversion increases the likelihood of destruction of crops and other properties. It is noteworthy that, although human-disturbed habitats favour breeding of generalist species, protection supports many species that avoid disturbed areas in the breeding season.

Juvenile recruitment into the community was higher during the dry than the wet season as a result of the higher breeding activity during the wet season. This implies that the small mammals match their breeding activities with rainfall patterns, resulting in more juveniles in the dry season. This supports the prediction that

juvenile recruitment in the ecosystem should be significantly higher in the dry than the wet season. Similar patterns have been reported by other studies (Gentile et al., 2000; Mulungu et al., 2013; Williams et al., 2017; Wirminghaus and Perrin, 1993) and imply that seasonality in juvenile recruitment is driven by resource seasonality such that juveniles join the community in the late wet and early dry seasons when high-quality resources critical for their survival and contributed by the previous rainfall season are abundant. This corroborates observations of other mammals and birds that recruitment is typically timed to match seasonal peaks in resource availability and quality to maximize fitness (Visser and Both, 2005) and enhance population growth (Habtamu and Bekele, 2008; Lima et al., 2003). Although the majority of the juveniles was recorded in the community in the dry season, few were also found in the wet season, reflecting inter-specific distinctions in the length and timing of breeding and growth rates (Oli and Dobson, 1999). Therefore, monthly monitoring would be necessary to accurately determine responses to resource seasonality of juvenile recruitment, operational sex ratio and breeding activity patterns in populations of individual small mammal species.

Higher juvenile recruitment for the pastoral land than for both the agricultural and the protected lands contradicts the prediction that it should be higher for the protected land. This is largely due to the presence of the larger number of actively breeding individuals in the pastoral land in the wet season. Notably, the two-generalist species (*A. niloticus* and *M. natalensis*) contributed > 50% of actively breeding individuals in the pastoral land. Thus, the higher juvenile recruitment among the generalist species in the pastoral land reflects the influence of human activities, supporting the observation that these two species thrive better in human-disturbed habitats (Assefa and Chelmala, 2019; Getachew et al., 2016; Magige, 2016). Livestock grazing and agriculture, the two predominant forms of human activities in the pastoral land, typically alter plant structure and create simplified habitats that favour generalist species (Jones and Longland, 1999; Keesing, 1998; Luza et al., 2016; Tabeni and Ojeda, 2005). Consequently, intense grazing in the pastoral land apparently simplify habitats, thereby favouring breeding by generalist species and leading to a higher mean juvenile recruitment relative to the other two land use types. Low recruitment in the agricultural land could be due primarily to the cropping systems used. It typically takes less than a month to prepare land by oxen and replant because most of the farmers cultivate crops for cash income (Makundi et al., 1999; Massawe et al., 2003). Thus, the cropping system used in the agricultural land likely contributed to a relatively less stable supply and availability of food and shelter for the small mammal species after harvests (Massawe et al., 2006, 2003). Alternatively, the lower juvenile recruitment in the

protected and agricultural than in the pastoral land can be interpreted as indicating that many species probably breed in both land use types at other times of the year not covered by this study.

Sex ratio was male biased, both in the wet and dry seasons. This contradicts the prediction that females should be more abundant than males in the wet season but less abundant than males in the dry season. The male-biased sex ratio is probably due to the tendency of males travelling long distances than females, resulting in a higher probability of getting trapped (Bantihun and Bekele, 2015; Clapperton, 2006; D'Andrea et al., 1999; Duque et al., 2005; Phelps, 2006). Although sex ratio was male skewed for both seasons, it might be due to differences in the amount of reproductive effort invested between sexes such that females spend more time caring for the young than males (Breedveld et al., 2019; Johnson et al., 2001). In addition, females become more risk-averse during pregnancy or lactation, and hence more wary when encountering unfamiliar situations which would make them less trappable than males (Dickman, 1999). Furthermore, the skew in sex ratio towards males in the agricultural and protected lands than in the pastoral land contradicts the prediction that males should be more abundant in the pastoral land whereas females should favour the agricultural and protected lands. However, the higher sex ratio in favour of females in the pastoral than the agricultural or protected land agrees with the prediction that females in good body condition tend to give birth to more males than nutritionally stressed females (Trivers and Willard, 1973). It follows that females in the pastoral land (with likely food shortages) might be giving birth to more females whereas females in the protected and agricultural lands (with likely greater food availability) to more males. This is expected because livestock grazing reduces food quality and quantity compared to protection, while agricultural land provides abundant but more seasonal food supply to small mammals especially when crops ripen and just after harvests (Caro, 2001; Jones and Longland, 1999; Keesing, 1998; Tabeni and Ojeda, 2005). Consequently, females probably have better body condition due to their access to relatively higher food availability in the protected and agricultural lands than in the pastoral land. When population sex ratio is male skewed it may affect the entire population by further reducing the number of available females through stress and competition for mates (Rankin et al., 2011). Also, it can be more problematic because higher male-male competition and the production of fewer offspring are all detrimental to population growth and viability (Grayson et al., 2014; López-Sepulcre et al., 2009).

Conclusion

The Serengeti small mammal community showed strong

breeding seasonality with active breeding peaking in the wet season. Availability and quality of resources resulted in a peak in juvenile recruitment in the early dry season. Small mammal breeding and recruitment varied across land use types in both the wet and dry seasons, reflecting spatial distinctions in resource availability and quality related to contrasting land uses. Specifically, mean breeding and recruitment probabilities for *M. natalensis* and *A. niloticus* species were higher in the pastoral land than the protected and agricultural lands, suggesting that livestock grazing simplified small mammal habitats in the pastoral land, favouring generalist species. Adult sex ratio varied seasonally and across land use types and was male skewed, with potentially adverse consequences for juvenile recruitment, population growth and viability. However, skewed sex ratio towards females for the pastoral than the agricultural or protected land reinforces the prediction of the Trivers-Willard hypothesis that females in poor body condition should give birth to more females than the presumably nutritionally less stressed females in the protected or agricultural land. Accordingly, human alterations to natural habitats degrade and simplify habitats, favouring generalist species, including some serious pest species, and altering small mammal breeding patterns and population demography. It follows that protection, by enhancing habitat intactness helps enhance species richness by reducing opportunistic pest species that typically numerically dominate disturbed habitats.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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Table S1. Tests of pairwise differences of simple effects of season and sex (season and sex interaction) for breeding probability (logit scale) of small mammals trapped in the Serengeti Ecosystem during 2017 and 2018 based on the logistic regression model.

Statement number	Effect	Slice	Season	Season	Difference estimate	Standard error	DF	tValue	Pr> t	95% Lower CL	95% Upper CL
1	SeasonxSex	Sex F	Dry	Wet	-2.461	0.5753	11	-4.28	0.001	-3.728	-1.19
1	SeasonxSex	Sex M	Dry	Wet	-0.708	0.3075	11	-2.31	0.042	-1.386	-0.03
2	SeasonxSex	Season	Dry		1.001	0.3657	11	2.74	0.019	0.197	1.807
2	SeasonxSex	Season	Wet		2.754	0.5402	11	5.1	3x10 ⁻⁴	1.566	3.944

Pairwise comparisons are performed among the levels of one factor at a fixed level of the other factor. The null hypothesis is that the difference is zero. DF denotes degrees of freedom.

Table S2. Pairwise differences (across land use types and seasons) in mean recruitment probabilities of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	Landuse	Season	Landuse	Season	Difference Estimate	Standard error	DF	t Value	Pr> t	95% Lower CL	95% Upper CL
Landuse	Agricultural		National park		0.755	0.5952	8	1.27	0.24	-0.62	2.1276
Landuse	Agricultural		Pastoral		-1.406	0.462	8	-3.04	0.016	-2.47	-0.3406
Landuse	National park		Pastoral		-2.1611	0.459	8	-4.71	0.002	-3.22	-1.1025
Season		Dry		Wet	0.9984	0.4059	8	2.46	0.039	0.062	1.9345

Table S3. Adjusted mean sex ratio and the associated 95% confidence limits (CL) for small mammals trapped in the Serengeti Ecosystem during 2017 and 2018 based on the logistic regression model.

Effect	Landuse	Sex	Mean sex ratio	Standard error	95% Lower CL	95% Upper CL
Sex		F	0.2751	0.02402	0.1844	0.3892
Sex		M	0.7249	0.02402	0.6108	0.8156
LandusexSex	Agriculture	F	0.1563	0.03926	0.04892	0.4001
LandusexSex	Agriculture	M	0.8437	0.03926	0.5999	0.9511
LandusexSex	Park	F	0.2878	0.03055	0.1755	0.4342
LandusexSex	Park	M	0.7122	0.03055	0.5658	0.8245
LandusexSex	Pastoral	F	0.4221	0.03406	0.286	0.5712
LandusexSex	Pastoral	M	0.5779	0.03406	0.4288	0.714

F and M denote females and males, respectively.

Table S4. Pairwise differences (across land use types and sex on the logit link scale) of the mean sex ratio and the associated 95% confidence limits (CL) of small mammals trapped in the Serengeti ecosystem during 2017 and 2018 based on the logistic regression model. DF denotes degrees of freedom.

Effect	Landuse	Sex	Landuse	Sex	Difference estimate	Standard error	DF	t Value	Pr > t	95% Lower CL	95% Upper CL
Sex		F		M	-1.9374	0.1677	2	-11.55	0.0074	-2.659	-1.2158
Landuse×Sex	Agriculture	F	Agriculture	M	-3.3725	0.4135	2	-8.16	0.0147	-5.1517	-1.5932
Landuse×Sex	Agriculture	F	Park	F	-0.7803	0.3335	2	-2.34	0.1442	-2.2153	0.6547
Landuse×Sex	Agriculture	F	Park	M	-2.5921	0.3324	2	-7.8	0.0161	-4.0225	-1.1618
Landuse×Sex	Agriculture	F	Pastoral	F	-1.3722	0.3272	2	-4.19	0.0524	-2.7802	0.03575
Landuse×Sex	Agriculture	F	Pastoral	M	-2.0002	0.3305	2	-6.05	0.0262	-3.4222	-0.5782
Landuse×Sex	Agriculture	M	Park	F	2.5921	0.3324	2	7.8	0.0161	1.1618	4.0225
Landuse×Sex	Agriculture	M	Park	M	0.7803	0.3335	2	2.34	0.1442	-0.6547	2.2153
Landuse×Sex	Agriculture	M	Pastoral	F	2.0002	0.3305	2	6.05	0.0262	0.5782	3.4222
Landuse×Sex	Agriculture	M	Pastoral	M	1.3722	0.3272	2	4.19	0.0524	-0.0358	2.7802
Landuse×Sex	Park	F	Park	M	-1.8118	0.2102	2	-8.62	0.0132	-2.7162	-0.9073
Landuse×Sex	Park	F	Pastoral	F	-0.5919	0.2055	2	-2.88	0.1024	-1.4761	0.2923
Landuse×Sex	Park	F	Pastoral	M	-1.2199	0.2029	2	-6.01	0.0266	-2.093	-0.3467
Landuse×Sex	Park	M	Pastoral	F	1.2199	0.2029	2	6.01	0.0266	0.3467	2.093
Landuse×Sex	Park	M	Pastoral	M	0.5919	0.2055	2	2.88	0.1024	-0.2923	1.4761
Landuse×Sex	Pastoral	F	Pastoral	M	-0.628	0.1929	2	-3.25	0.0828	-1.4581	0.2021

S1 Data. The total number of adult (Age=A) male (Sex=M) and female (Sex=F) breeding and non-breeding small mammals captured in three land uses in the Tanzania Serengeti Ecosystem in the wet and dry seasons of 2017 and 2018.

Year	Land use	Season	Sex	Age	Total number Breeding	Total number not breeding	Total number captured	Total trap nights	logTrapnights	Prob	Lower 95% prediction Limit for Prob	Upper 95% prediction Limit for Prob
2017	National Park	Wet	F	A	7	1	8	11280	9.33079	0.8	0.59428	0.91613
2017	National Park	Wet	M	A	3	16	19	11280	9.33079	0.16304	0.09475	0.26609
2017	National Park	Dry	F	A	1	3	4	11280	9.33079	0.18182	0.07604	0.37503
2017	National Park	Dry	M	A	1	22	23	11280	9.33079	0.06557	0.02198	0.17971
2017	Pastoral Land	Wet	F	A	22	2	24	5640	8.63764	0.94444	0.77413	0.98828
2017	Pastoral Land	Wet	M	A	7	15	22	5640	8.63764	0.36111	0.20846	0.54814
2017	Pastoral Land	Dry	F	A	4	8	12	5640	8.63764	0.4697	0.3398	0.60384

2017	Pastoral Land	Dry	M	A	2	32	34	5640	8.63764	0.17647	0.10793	0.27512
2017	Agricultural Land	Wet	F	A	0	0	0	11280	9.33079	0.66667	0.11894	0.96735
2017	Agricultural Land	Wet	M	A	1	7	8	11280	9.33079	0.24138	0.10907	0.45264
2017	Agricultural Land	Dry	F	A	0	3	3	11280	9.33079	0.3	0.08579	0.66184
2017	Agricultural Land	Dry	M	A	7	18	25	11280	9.33079	0.21739	0.11228	0.37891
2018	National Park	Wet	F	A	17	5	22	11280	9.33079	0.8	0.59428	0.91613
2018	National Park	Wet	M	A	12	61	73	11280	9.33079	0.16304	0.09475	0.26609
2018	National Park	Dry	F	A	5	24	29	11280	9.33079	0.18182	0.07604	0.37503
2018	National Park	Dry	M	A	3	35	38	11280	9.33079	0.06557	0.02198	0.17971
2018	Pastoral Land	Wet	F	A	12	0	12	5640	8.63764	0.94444	0.77413	0.98828
2018	Pastoral Land	Wet	M	A	6	8	14	5640	8.63764	0.36111	0.20846	0.54814
2018	Pastoral Land	Dry	F	A	27	27	54	5640	8.63764	0.4697	0.3398	0.60384
2018	Pastoral Land	Dry	M	A	16	52	68	5640	8.63764	0.17647	0.10793	0.27512
2018	Agricultural Land	Wet	F	A	2	1	3	11280	9.33079	0.66667	0.11894	0.96735
2018	Agricultural Land	Wet	M	A	6	15	21	11280	9.33079	0.24138	0.10907	0.45264
2018	Agricultural Land	Dry	F	A	3	4	7	11280	9.33079	0.3	0.08579	0.66184
2018	Agricultural Land	Dry	M	A	3	18	21	11280	9.33079	0.21739	0.11228	0.37891

Also provided are the total trap nights of effort per land use and the breeding (Number breeding/Total) probability and its 95% lower and upper confidence limits. The probabilities were predicted using a logistic regression model with a binomial error distribution, a logit link function and the logarithm of trap night as the offset.

S2 Data. The total number of adult and juvenile small mammals captured in three land uses in the Tanzania Serengeti Ecosystem in the wet and dry seasons of 2017 and 2018. Also provided are the total trap nights of effort per land use and the juvenile recruitment (Juveniles/Total) probability and its 95% lower and upper confidence limits. The probabilities were predicted using a logistic regression model with a binomial error distribution, a logit link function and the logarithm of trap night as the offset.

Year	Land use	Season	Adults	Juveniles	Total	Trapnight	logTrapnight	Prob	Lower 95% Prediction	Upper 95% Prediction
									Limit for Prob	Limit for Prob
2017	National Park	Wet	30	0	30	11280	9.33079	0.01463	0.0047	0.04459
2017	National Park	Dry	32	3	35	11280	9.33079	0.03875	0.01455	0.09913
2017	Pastoral Land	Wet	46	8	54	5640	8.63764	0.06056	0.02566	0.13626
2017	Pastoral Land	Dry	46	24	70	5640	8.63764	0.14889	0.10029	0.21541
2017	Agricultural Land	Wet	8	0	8	11280	9.33079	0.03063	0.0091	0.09802
2017	Agricultural Land	Dry	28	5	33	11280	9.33079	0.07899	0.03081	0.18792

2018	National Park	Wet	102	0	102	11280	9.33079	0.01463	0.0047	0.04459
2018	National Park	Dry	67	3	70	11280	9.33079	0.03875	0.01455	0.09913
2018	Pastoral Land	Wet	28	0	28	5640	8.63764	0.06056	0.02566	0.13626
2018	Pastoral Land	Dry	123	2	125	5640	8.63764	0.14889	0.10029	0.21541
2018	Agricultural Land	Wet	28	0	28	11280	9.33079	0.03063	0.0091	0.09802
2018	Agricultural Land	Dry	28	1	29	11280	9.33079	0.07899	0.03081	0.18792
