ABSTRACT

Robust measures of animal densities are necessary for effective wildlife management. Leopards (*Panthera pardus*) and spotted hyenas (*Crocuta crocuta*) are higher order predators that are data deficient across much of their East African range and in Uganda, excepting for one peer-reviewed study on hyenas, there are presently no credible population estimates for these species. A lack of information on the population status and even baseline densities of these species has ramifications as leopards are drawcards for the photo-tourism industry, and along with hyenas are often responsible for livestock depredations from pastoralist communities. Leopards are also sometimes hunted for sport. Establishing baseline density estimates for these species is urgently needed not only for population monitoring purposes, but in the design of sustainable management offtakes, and in assessing certain conservation interventions like financial compensation for livestock depredation. Accordingly, we ran a single-season survey of these carnivores in the Lake Mburo National Park of south-western Uganda using 60 remote camera traps distributed in a paired format at 30 locations. We analysed hyena and leopard detections under a Bayesian spatially explicit capture-recapture (SECR) modelling framework to estimate their densities. This small national park (370 km$^2$) is surrounded by Bahima pastoralist communities with high densities of cattle on the park edge (with regular park incursions). Leopard densities were estimated at 6.31 individuals/100 km$^2$ (posterior SD = 1.47, 95% CI [3.75–9.20]), and spotted hyena densities were 10.99
individuals/100 km$^2$, but with wide confidence intervals (posterior SD = 3.35, 95% CI [5.63–17.37]). Leopard and spotted hyena abundance within the boundaries of the national park were 24.87 (posterior SD 7.78) and 39.07 individuals (posterior = SD 13.51) respectively. Leopard densities were on the middle end of SECR studies published in the peer-reviewed literature over the last 5 years while spotted hyena densities were some of the first reported in the literature using SECR, and similar to a study in Botswana which reported 11.80 spotted hyenas/100 km$^2$. Densities were not noticeably lower at the park edge, and in the southwest of our study site, despite repeated cattle incursions into these areas. We postulate that the relatively high densities of both species in the region could be owed to impala *Aepyceros melampus* densities ranging from 16.6–25.6 impala/km$^2$. Another, potential explanatory variable (albeit a speculative one) is the absence of interspecific competition from African lions (*Panthera leo*), which became functionally extinct (there is only one male lion present) in the park nearly two decades ago. This study provides the first robust population estimate of these species anywhere in Uganda and suggests leopards and spotted hyenas continue to persist in the highly modified landscape of Lake Mburo National Park.

**Subjects** Conservation Biology, Ecology, Zoology  
**Keywords** *Panthera pardus*, *Crocuta crocuta*, Spatially explicit capture-recapture, Population size, East Africa, Human-carnivore conflict

## INTRODUCTION

Precise measures of animal densities represent one of the most fundamental precursors for effective wildlife management (*Karanth, 1995; White & Burnham, 1999; Duangchantrasiri et al., 2016; Rayan & Linkie, 2015*). Density estimates assist *inter alia* with species assessments (*Jacobson et al., 2016*), the setting of harvest quotas (*Balme, Slotow & Hunter, 2009*), and in gauging the viability of individual populations (*Sollmann et al., 2011*). Measures of animal abundance and density are becoming increasingly critical for species that are exposed to significant anthropogenic pressures, are constrained to small habitat patches, and are important to the economies of developing nations (*O’Bryan et al., 2018*).

Large carnivores naturally occur at relatively low densities and have large space requirements (*Balme, Slotow & Hunter, 2009; Gopalaswamy et al., 2012b*). Anthropogenic sources of mortality at the edges of small reserves can therefore depress carnivore densities, even within protected areas because animals move beyond their boundaries and are killed (e.g., *Balme, Slotow & Hunter, 2009; Woodroffe & Ginsberg, 1998*). In Uganda, most protected areas are relatively small, isolated and have high human pressures at their edges (*Venter et al., 2016*). Additionally, most Ugandan national parks and wildlife reserves are bordered by livestock rearing communities, and large carnivores regularly kill livestock in these areas (*Ochieng, Ahebwa & Visseren-Hamakers, 2015*). Consequently, large carnivores are often killed in retaliation for livestock killing, and damage through poisoning, trapping or shooting (*Tweheyo et al., 2012*).
Leopard (*Panthera pardus*) and spotted hyena (*Crocuta crocuta*) are examples of species which have impacts on the livelihoods of local communities in Uganda (*Ochieng, Ahebwa & Visseren-Hamakers, 2015*). Both species were responsible for 1,102 attacks on cattle, sheep and goats on the edge of Lake Mburo National Park (hereafter LMNP) (spotted hyenas \( n = 762 \) or 69%, leopards \( n = 340 \) or 31% between January 2009–December 2018, *Braczkowski et al., 2020c*). Such conflict between these species and pastoralists may have ramifications and at least 19 leopards were killed on the boundary of LMNP in a 4-year period from 2003–2006 (CITES CoP 14 Proposal 3), and two hyena clans (each >14 individuals in size) that were regularly viewed by tourists were poisoned in 2007 (R Schenk, 2018, personal communication). However, both species are also important for the wildlife-viewing tourism (*Van der Meer, Badza & Ndlovu, 2016*) and in Uganda in 2018 alone, 1,585 people purchased a night game drive permit for leopard viewing in LMNP, equating to US$47,550 in revenue for the Ugandan Wildlife Authority (A Kule, 2018, personal communication). This often leads to contradictory management goals, where one entity seeks higher densities to maximize tourism revenue, and the other seeks lower densities due to livelihood losses accrued from conflict. However, a lack of robust information on the population status of leopards and spotted hyenas inhibits the design of sustainable management offtakes and also in assessing the impact of conservation interventions on carnivores and communities (*e.g.*, Financial compensation, the erection of livestock protection bomas etc.).

To address these concerns, we sought to estimate the population abundance and densities of leopards and spotted hyenas in the LMNP, south-western Uganda. LMNP is a small, protected area that lacks much of the charismatic megafauna found elsewhere in the country *e.g.*, mountain gorillas (*Gorilla beringei beringei*), chimpanzees (*Pan troglodytes*), African elephants (*Loxodonta africana*) and lions (*Panthera leo*). Consequently, leopards and spotted hyenas are important tourism draws for the region. This is even more important as African lions became functionally extinct in LMNP in the early 2000’s (*Uganda Wildlife Authority, 2010*). There is also legal trophy hunting of leopards on LMNP’s edge and high rates of human-leopard conflict on its boundary (*Braczkowski et al., 2020c*). This study represents the first assessment of leopards undertaken in a protected area system in Uganda and provides one of the first spatially explicit estimates of spotted hyena densities in the literature. This study produces a baseline single season snapshot into the population densities for both species to inform conservation management in the region and to better track the impacts of conservation interventions.

**STUDY AREA**

We studied leopards and spotted hyenas in the LMNP (370 km\(^2\)), Kiruhura district, Western Uganda (30°47’–31°04’E, 0°30’–0°30’S, Fig. 1). The LMNP forms part of the Akagera savanna ecosystem which extends from Rwanda and north-western Tanzania down into south-western Uganda (*Menaut, 1983; Van de Weghe, 1990*). LMNP experiences a bimodal annual rainfall pattern (October–December and February–June) and annual rainfall and temperatures average 800 mm and 28 °C, respectively (*Moe et al., 2016*). The woody vegetation in the park is characterized by dry *Acacia* savanna dominated
by *Acacia hockii*, woodlands, thickets and swamps which occur on the edges of Lake Kachera and Mburo (*Rannestad et al., 2006*). The most common grasses include (*Loudetia kagerensis*), (*Chloris gayana*), and (*Sporobolus pyramidalis*). LMNP supports one of two remaining population of impala (*Aepyceros melampus*) in Uganda, the most common and preferred prey of the African leopard (*Hayward et al., 2006*). The park also harbours Plains zebra (*Equus quagga*), Cape buffalo (*Syncerus caffer*), Defassa waterbuck (*Kobus ellipsiprymnus defassa*), bushbuck (*Tragelaphus scriptus*) and warthog (*Phacochoerus africanus*, *Rannestad et al., 2006*). There is only one male lion (≥10 years old) in LMNP (a vagrant thought to have come from Akagera National Park, in neighbouring Rwanda). LMNP is bordered by a matrix of small human settlements, small-scale subsistence crops, dairy ranches and communal grazing lands (*Ochieng, Ahebwa & Visseren-Hamakers, 2015*).
Park history, introduction of trophy hunting and human-carnivore conflict

Although the national park itself is small, much of the former park area—which is now mainly used as cattle rangeland—still has considerable woodlands, thickets and natural vegetation and Rannestad et al. (2006) noted higher densities of bushbuck, impala, reedbuck (Redunca redunca), waterbuck and zebra outside of the national park’s borders during the wet season. The region surrounding LMNP has a trophy hunting scheme which was initiated due to increasing complaints by communities, stating that the increasing wildlife was a nuisance (Ochieng, Ahebwa & Visseren-Hamakers, 2015). The leopard is only allowed to be hunted when a problem animal tag is made available by Ugandan Wildlife Authority (hereafter UWA) attributed to repeated stock killing and damage. Although harvests of leopards since 2007 have been low in Uganda (17 skins, skulls and trophies exported from 2009–2017), attempts were made to have the species downgraded from CITES Appendix 1 to Appendix 2 and proposed a quota of 50 leopards annually (despite the lack of even a single abundance estimate anywhere in the country, CITES CoP 14 Proposal 3). Currently, 28 leopards are available annually on quota country-wide. Contrastingly, in Africa, hyenas are often taken opportunistically by trophy hunters rather than as prized trophy animals and we could not find any evidence that they are an actively hunted species in Uganda (see for example: http://www.uganda-wildlife-safaris.com).

METHODS

Camera trapping

This research was granted approval by the Uganda Wildlife Authority under permit number: UWA/COD/96/05 as approved by the Executive Director Mr Stephen Masaba. We implemented one single season camera-trap survey for 53 days in the LMNP from 26 July 2018–16 September 2018 using Cuddeback™ 20-megapixel Long Range IR camera traps (powered by 8 AA batteries each) set in a paired format. The survey encompassed 30 camera trap sites distributed across the national park (Fig. 1), but we omitted camera traps in the far western sector of the park due to a lack of road access. Each camera trap site consisted of two camera traps, each mounted to a 1 m steel pole 40 cm from the ground. We positioned each camera perpendicular to a vehicle track or game trail at a 60–75°-angle to facilitate early detection of leopards and spotted hyenas. We set our camera traps on roads, vehicle tracks, trails and drainage lines, as these are regularly used by leopards and spotted hyenas as travel and hunting routes (Balme, Hunter & Slotow, 2009; Balme, Slotow & Hunter, 2009; Henschel, Malanda & Hunter, 2014).

We checked traps every 4–7 days to correct for animal damage, replace memory cards and to assess battery functionality (Braczkowski et al., 2016). Camera traps were set to burst mode and took five images every time the infrared sensor was triggered. We set camera traps in a way as to ensure that at least one camera-trap site was present in an area corresponding to the smallest female leopard home-range recorded in the literature (30 km²; Bailey, 1993, and 23 km² in Fattebert et al., 2016), as these are smaller than male
leopards and spotted hyenas. Our camera spacing was 2.1 km (5–7 camera stations per female home range). We chose this camera spacing in order to ensure that no animal had a zero probability of capture (Karanth & Nichols, 1998). The identity of individual leopards and spotted hyenas was determined by their unique rosette and spot patterns (Miththapala et al., 1989; O’Brien & Kinnaird, 2011). For leopards, we were able to classify the sex of individuals by using distinctive morphological cues such as the presence of testes and the enlarged dewlap and sagittal crest in males (Balme, Hunter & Braczkowski, 2012; Braczkowski et al., 2015a).

The first and third author assigned individual identity to temporally unique photographs and only included into the final density estimation process individuals for which there was consensus (Bahaa-el-din et al., 2016). We excluded images that were blurred, were too far away from the camera trap and those where observers could not agree on identity. For the purpose of building capture histories with known unique individual identities, we used both flanks of leopards in our analysis (Fig. 2). Spotted hyenas, however, often walked around cameras and did not present a clear flank on

**Figure 2** Individual identification of spotted hyenas and leopards from camera traps. Individual identification information extracted from leopards and spotted hyenas in the LMNP, 2018. Slide 1 (top) denotes a female leopard captured at trap location five on sampling occasion two and 10 respectively. Rosette patterns and facial spots were extracted during these two occasions. Slide 2 (bottom) denotes the spot pattern extracted from a spotted hyena captured at location 12 and 27 on sampling occasions 5 and 22 respectively.
both sides of a single animal, and several individuals moved around a single camera at the same time. To avoid mismatching flanks and mistakenly double-count individuals, we chose the flank of hyenas with the highest number of photographs recorded during our survey (Henschel, Malanda & Hunter, 2014).

**SECR modelling**

We estimated leopard and spotted hyena densities and abundance in LMNP using Bayesian spatially explicit capture-recapture modelling. By incorporating spatial information into the detection process, the method does not suffer from the “edge effects” common to non-spatial estimators (Gopalaswamy et al., 2012a). The modelling approach uses a state (leopard and spotted hyena population size and locations in the landscape) and observation process (Royle et al., 2009; Gopalaswamy et al., 2012b). To accurately estimate the densities and home-range centres of both species we generated potential activity centres across our study area (370 km$^2$) in the form of 0.336 km$^2$ (i.e., 580 m × 580 m, Gopalaswamy et al., 2012a) equally spaced pixels. This state-space assumes the number of leopards and spotted hyenas found in these pixels are defined by a binomial process, but because spotted hyenas are often found in groups, the state process allows for ≥2 spotted hyenas to have an activity centre in the same pixel (Gopalaswamy et al., 2012a). The state space encompassed the LMNP, and a buffer of 25 km around it (including the eastern rangelands bordering the park, Kanyaryeru and the southern farmlands). We masked out all human settlements and water bodies inside and surrounding the national park, as leopards and spotted hyenas are unlikely to have their home-range centres directly in such unsuitable habitats (Royle et al., 2009; Gopalaswamy et al., 2012a; Braczkowski et al., 2016). We used a classical capture re-capture sampling design and created a standard capture re-capture matrix (trap locations, individual leopards or hyenas and sampling occasions, e.g., du Preez, Loveridge & Macdonald, 2014; Braczkowski et al., 2016; Williams et al., 2017). Large terrestrial carnivores regularly feature differences at the sex-level in their home-range sizes and capture probability (Gopalaswamy et al., 2012b; Braczkowski et al., 2016). Differences in movements of animals based upon sex can affect the observation process in spatial capture-recapture (Sollmann et al., 2011). To factor this into our models, we included a sex-specific covariate in the observation process and accounted for different capture probability for leopards. We did not do this for hyenas as the female spotted hyenas feature a pseudo-scrotum which makes sexing difficult, and the visibility of males’ testes was often obscured by their large tail (Hamilton, Tilson & Frank, 1986).

In SECR modelling, $\sigma$ is the scale parameter, and represents the rate of decline in the detection rate as the location of the animal’s activity centre moves away from a camera trap station. $\lambda_0$ is the basal encounter rate and can be defined as the encounter rate of an animal whose activity centre lies exactly at a camera trap station. The detection rates of an individual animal decline with increasing distance between its activity centre and camera trap location (Borchers & Efford, 2008; Royle et al., 2009) and the parameter $\theta$ defines the shape of the detection function. If this parameter is estimated from the given data, the shape of the detection function could define how an animal utilizes space or
resources in its environment (Elliot & Gopalaswamy, 2017). In practice, since encounter rates are so small, they are approximately equal to detection probabilities (Efford, 2019). We either used a fixed $\theta$ at 0.75 (Elliot & Gopalaswamy, 2017) and 1 (Gaussian form, Royle et al., 2009) or estimated a continuous $\theta$ parameter from the data.

The complementary log-log link was used to convert encounter rates to Bernoulli detections, therefore, in our models, the probability of detecting a leopard or hyena $i$ in pixel $j$ is defined by a complementary log-log function of covariates.

We assessed six a priori models for leopards, and two for spotted hyenas (parameter definitions are presented in Table 1). Model 1 estimated the detection function (this is defined by $\theta$) and assumed that detection probability is sex specific:

$$cloglog (\pi_{ij}) = \log(\lambda_0) + \beta_{sex} - f[dist(i, j)|\theta, \sigma_{sex}]$$

where, $\pi_{ij}$ describes the detection probability on a given sampling occasion, which is a function of the basal encounter rate $\lambda_0$ and distance between the activity center of individual $i$ and pixel $j$, $\theta$ and sex-specific $\sigma_{sex}$. The specific form of this detection function is:

$$f[dist(i, j)|\theta, \sigma_{sex}] = \exp \left[ \frac{-dist(i, j)^{2\theta}}{2\sigma_{sex}^2} \right]$$

Model 2 was based on the assumption that detection probability is not dependent on sex, (i.e., $\beta_{sex}$ was fixed at 0). The rate of decline in detection probability ($\sigma$) however, remained sex specific because this parameter is also linked to the movement of animals.

Model 3 as with model 2, had $\beta_{sex}$ set at 0 while the detection function was set at $\theta = 0.75$

Model 4 was based on the assumption that basal encounter rate is dependent on sex, thus, $\beta_{sex}$ was fixed at 1. Rate of decline in detection probability ($\sigma$) also remained sex specific. The detection function parameter $\theta$ was fixed at 0.75.

| Parameter | Definition |
|-----------|------------|
| $n$       | Total number of leopards or hyenas detected during the survey period |
| $nz$      | Number of leopards augmented to $n$, so $M = n + nz$ represents the maximum number of leopards in the large state space $S$ |
| $\sigma_F$ | Rate of decline in detection probability with increasing distance between the activity center of a leopardess and the location at which female leopard was found |
| $\sigma_M$ | Rate of decline in detection probability with increasing distance between the activity center of a leopard and the location at which male leopard was found |
| $\beta_{sex}$ | Difference of the complementary log-log value of detection probability between a male and female leopard |
| $\lambda_0$ | Basal encounter rate of a leopard whose activity center is located exactly at the centroid of a grid cell |
| $\psi$ | Ratio of the true number of individuals in the population compared with the data-augmented population $M$ |
| $N_{super}$ | Total number of leopards in the larger state space $S$ |
| $\psi_{sex}$ | Proportion of leopards that are female ($1 - \psi_{sex}/\psi_{sex}$) |
| $\theta$ | Determines the shape of the estimated detection function, value $\theta$ ranges from 0.5 (exponential form) to 1 (Gaussian) |
| $D$ | Estimated density of leopards per 100 km$^2$ |
Model 5 assumed basal encounter rate is dependent on sex but rate of decline in
detection probability was independent of sex. The detection function parameter was fixed
at $\theta = 0.75$.

Model 6 was the same as model 1 but the detection function parameter ($\theta$) was
fixed at 1.

For the spotted hyenas’ density assessment, we only used model 1 and model 6 due to
the lack of a sex covariate.

We used Bayesian Markov Chain Monte Carlo (MCMC) simulation and the
Metropolis-Hastings algorithm (Tierney, 1994) to run our models in the package SCRbayes
(https://github.com/jaroyle/SCRbayes) in the programming environment R Version 3.6.1
(R Development Core Team, 2019). We set each model to run for 20,000 iterations
including a burn-in of 5,000 iterations but we adjusted this further if we did not arrive at a
standing distribution, (refining burn-in period and initial iterations further). Each model
was set to run for four chains (Elliot & Gopalaswamy, 2017). Model adequacy was
determined by examining the Bayesian $p$-value on individual encounters (Royle et al.,
2009). MCMC convergence was assessed using the Gelman–Rubin diagnostic (Gelman &
Rubin, 1992). The five input files necessary to run these analyses and accompanying
R scripts are provided in the supporting information section of this manuscript
(Information S3). Although we were principally interested in estimating density,
we also computed posterior mean abundance across the study area of the greater LMNP
system.

RESULTS
We recorded a total of 1,444 trap nights during the 53-day survey period. Cameras
were not functional due to animal interference and battery failures for 146 trap nights, and
these were not included in the SECR analysis. We recorded a total of 61 temporally
independent (i.e., animals counted only once in a 24-h period) detections of leopards
during our camera trap survey, and 51 spotted hyena detections (Table 2). From these
we recorded 112 and 42 useable flanks for leopards and hyenas respectively (51 right
hyena flanks vs 32 left flanks; nine excluded due to not identifiable or juvenile hyena).
We identified 20 unique leopards (six adult males and 14 adult females), and 27 (no sex
noted) spotted hyenas. This equates to a detection rate of 1.38 leopards and 1.87 spotted
hyenas per 100 trap-nights.

| Species       | Number of left flanks | Number of right flanks | Number of useable flanks for analysis | Unique individuals identified | Unique individuals recaptured |
|---------------|-----------------------|------------------------|---------------------------------------|------------------------------|------------------------------|
| Spotted hyena | 32                    | 51                     | 42                                    | 27                           | 8                            |
| Leopard       | 57                    | 55                     | 112                                   | 20                           | 13                           |

Table 2  Number of flanks of hyenas and leopards recorded during camera trapping in LMNP with
total recaptures.
Density estimates and model diagnostics

Bayesian p-values for all our leopard density models ranged from 0.61–0.76 (Table 3), indicating an adequate model fit (extremities 0.15–0.85). Convergence of models was indicated by a mean potential shrink reduction factor of <1.2 for each parameter for each model (Gelman & Rubin, 1992, Information S1). The same assessment of model adequacy was recorded for a model where sigma was estimated without a sex effect for the estimates of spotted hyena density (Bayesian p = 0.61 and shrink reduction factor for all parameters <1.2, Information S2). Model selection using marginal likelihood from Dey, Delampady & Gopalaswamy (2019) indicated that model 5, which considered basal encounter rate to be dependent on sex but detection probability independent of sex had the highest log likelihood score (log likelihood = −55,615.56, Table 3).

Leopard density estimates

Using model 5, leopard density for LMNP was estimated at 6.31 individuals/100 km$^2$ (posterior SD 1.47, 95% CI range [3.75–9.20, Table 4]). The posterior mean abundance for

| Table 3 SECR models from the Lake Mburo survey. Models used to generate our density analyses for leopards and spotted hyenas in the LMNP, Uganda, 2018. We present the model number, Bayes p-value to signify model adequacy and the marginal likelihood values used to select our models, and number of iterations used to achieve convergence. |
|-------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Species     | Model number    | Bayes p-value   | Marginal likelihood | Total iterations | Burn in required to reach convergence |
| Leopards    | 1               | 0.71            | −62,893.814       | 52,000           | 42,000           |
|             | 2               | 0.71            | −62,885.778       | 50,000           | 20,000           |
|             | 3               | 0.72            | −62,784.534       | 80,000           | 2,000            |
|             | 4               | 0.71            | −62,729.456       | 50,000           | 20,000           |
|             | 5               | 0.61            | −55,615.556       | 50,000           | 20,000           |
|             | 6               | 0.76            | −62,985.962       | 50,000           | 20,000           |
| Spotted hyenas | 1           | 0.62            | −41,030.296       | 11,000           | 6,000            |
|             | 2               | 0.64            | −41,045.548       | 11,000           | 1,000            |

| Table 4 Parameter estimates with accompanying posterior standard deviation for our spatially explicit capture recapture models estimating leopard and hyena densities in the Lake Mburo National Park, Uganda. |
|-------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Species     | Model number    | sigma (σm)      | sigma2 (σf)     | lam0 (λ0)       | beta sex (βsex) | Psi (ψ)         | ψsex           | Theta (θ)      | Density (D)   |
| Leopard     | 1               | 2.59            | 1.08            | 0.73            | 0.14            | 0.02            | 2.04            | 0.67            | 0.05            | 0.08            | 0.08            | 0.74            | 0.13            | 8.92            | 2.14            |
|             | 2               | 2.13            | 1.07            | 0.67            | 0.11            | 0.07            | 0            | 0            | 0.05            | 0.04            | 0.75            | 0.03            | 0.75            | 0.11            | 9.33            | 2.28            |
|             | 3               | 2.81            | 0.5             | 0.75            | 0.08            | 0.11            | 0            | 0            | 0.2            | 0.05            | 0.95            | 0.03            | 0.75            | 0            | 9.31            | 2.24            |
|             | 4               | 4.97            | 3.23            | 0.74            | 0.08            | 0.04            | 1.94            | 0.88            | 0.18            | 0.05            | 0.89            | 0.08            | 0.75            | 0            | 8.82            | 2.14            |
|             | 5               | 1.33            | 0.1             | 1.33            | 0.1             | 0.04            | 0.09            | 0.04            | 0.14            | 0.03            | 0.69            | 0.12            | 0.75            | 0            | 6.31            | 1.47            |
|             | 6               | 5.09            | 1.76            | 0.97            | 0.12            | 0.02            | 1.94            | 0.66            | 0.19            | 0.05            | 0.88            | 0.08            | 1              | 0              | 8.83            | 2.11            |
| Spotted hyena | 1           | 3.22            | 0.74            | 3.22            | 0.74            | 0.05            | 0.002           | 0            | 0.23            | 0.07            | 0.0003           | 0.0001           | 1              | 0              | 11.00            | 3.35            |
|             | 2               | 1.9             | 0.64            | 1.9             | 0.64            | 0.007           | 0.003           | 0            | 0.24            | 0.08            | 0.0003           | 0.0001           | 0.75            | 0.14           | 11.26            | 3.56            |
the Lake Mburo National Park was 24.87 (posterior SD 7.78) using this model. The leopard movement parameter or sigma $\sigma$ for males and females from this model was 1.33 km (this movement parameter is a measurement of how far animals travel in the landscape and is related to home range size; Braczkowski et al., 2020b). The next best-ranked candidate model (model number 4) which considered sex as a factor affecting detection probability estimated a movement parameter of 1.60 km for males and 0.59 km for females.

Hyena density estimates
For spotted hyenas, right flanks were recorded with the highest frequency (Table 2). Spotted hyena density for LMNP was estimated at 11.00 individuals/100 km$^2$ (posterior SD = 0.32, 95% CI range [5.57–17.09, Table 4]) using model 1. The spotted hyena movement parameter $\sigma$ for both sexes combined was 3.15 km. The posterior mean abundance for the entire state space buffer was 39.07 spotted hyenas (posterior = SD 13.51, Table 3).

DISCUSSION
We provide a robust estimate of leopard densities and abundance in the LMNP ecosystem, southwestern Uganda, and also the first SECR assessment for spotted hyenas in Uganda as a whole (however these had wider confidence intervals when compared to leopards). These estimates are important baselines for the future monitoring of leopard and spotted hyena populations in the LMNP, one which experiences both significant levels of human-carnivore conflict and trophy hunting (Braczkowski et al., 2020c). Robust estimates of population abundance and densities are a critical cornerstone for tracking changes and trends in carnivore populations over time (e.g., Balme, Slotow & Hunter, 2009; Williams et al., 2017). In this human-carnivore conflict-prone area, it is unknown whether retaliatory killings following depredation on livestock are sustainable in the long term, especially as the LMNP is small and isolated from other larger protected areas. Previous research has shown that carnivore populations in small, isolated national parks cannot withstand the edge effects from human-carnivore conflict (e.g., from cattle farming) and trophy hunting (Woodroffe & Ginsberg, 1998; Balme, Slotow & Hunter, 2009).

Possible explanations for observed leopard and hyena densities
Leopard densities in LMNP were on the mid-tier of estimates recorded in the recent literature using SECR studies ($n = 15$ studies from 2013–2018, Table 5). The leopard densities we observed at 6.31 individuals/100 km$^2$ are somewhat surprising given (a) the small size of LMNP, and (b) the high levels of conflict between these two carnivores and the livestock rearing communities on the park edge (Braczkowski et al., 2020c). Contrastingly, the hyena densities were similar to a SECR study in uMkhuze Game Reserve, northern Kwa-Zulu Natal, South Africa (a savanna system) which estimated a density of 10.59 individuals/100 km$^2$ (posterior SD = 2.10, De Blocq, 2014), and a study in Botswana’s Moremi estimated 11.80 (posterior SD = 2.60, Rich et al., 2019). We postulate that three factors may be contributing to these densities, namely (1) the availability of
Table 5  Literature review of recent SECR leopard studies. A review of 17 recent SECR studies performed on leopards in the last 5 years from the peer-reviewed literature. Some studies used a combination of maximum likelihood and Bayesian-based modelling approaches and therefore contain two or more estimates. We excluded the following studies for the following reasons: Goswami & Ganesh (2014)--no error reporting around estimates Kittle, Watson & Fernando (2017)--SECR results of tracks places results in contention Rich et al. (2019)--estimate is not directly reported only a figure is present. *We examined the first 10 pages of Google Scholar and limited the studies in this table to (a) those using SECR and (b) being published in the last 5 years.

| Study name                   | Location                                | Habitat type                | Model used to estimate density | Density estimate (leopards/100 km²) | SD (SE) |
|------------------------------|-----------------------------------------|-----------------------------|--------------------------------|-------------------------------------|---------|
| Balme et al. (2019)          | Sabi-Sands Game Reserve, South Africa   | Semi-wooded savanna         | Borchers & Efford (2008)       | 11.80                               | 2.60    |
| Borah et al. (2014)          | Manas National Park, India              | Tropical forest and mountains | Borchers & Efford (2008)       | 3.40                                | 0.82    |
| Braczkowski et al. (2016)    | Phinda Private Game Reserve, South Africa | Savanna                   | Royle et al. (2009)            | 3.55                                | 1.04    |
| Braczkowski et al. (2016)    | Phinda Private Game Reserve, South Africa | Savanna                   | Borchers & Efford (2008)       | 3.40                                | 1.20    |
| Devens et al. (2018)         | Bavianskloof mountains, South Africa    | Mountain fynbos and forest  | Royle et al. (2009)            | 0.24                                | 0.10    |
| Devens et al. (2018)         | Langeberg mountains, South Africa       | Mountain fynbos and forest  | Royle et al. (2009)            | 1.89                                | 0.30    |
| du Preez, Loveridge & Macdonald (2014) | Bubye Valley Conservancy, Zimbabwe       | Mopane woodland (savanna)   | Borchers & Efford (2008)       | 5.28                                | 0.89    |
| du Preez, Loveridge & Macdonald (2014) | Bubye Valley Conservancy, Zimbabwe       | Mopane woodland (savanna)   | Royle et al. (2009)            | 5.46                                | 1.14    |
| Hedges et al. (2015)         | Kenyir Wildlife Corridor, Malaysia      | Dipterocarp forest          | Borchers & Efford (2008)       | 3.30                                | 1.28    |
| Hedges et al. (2015)         | Kenyir Wildlife Corridor, Malaysia      | Dipterocarp forest          | Efford (2011)                  | 3.06                                | 0.91    |
| Kittle, Watson & Fernando (2017) | Horton Plains, Sri-Lanka               | Montane forest              | Borchers & Efford (2008)       | 13.40                               | 6.3     |
| Ngoprasert, Lynam & Gale (2017) | Ban Krang, Kaeng Krachan National Park, Thailand | Evergreen forest            | Borchers & Efford (2008)       | 2.50                                | 1.20    |
| Qi et al. (2015)             | Laoye mountains, China                  | Deciduous forest            | Royle et al. (2009)            | 0.62                                | 0.15    |
| Rahman et al. (2018)         | Ujong Kulon National Park, Java, Indonesia | Tropical forest            | Borchers & Efford (2008)       | 12.80                               | 1.99    |
| Rahman et al. (2018)         | Ujong Kulon National Park, Java, Indonesia | Tropical forest            | Royle et al. (2009)            | 11.54                               | 1.22    |
| Ramesh et al. (2017)         | Ndumo Game Reserve, South Africa        | Woodland savanna            | Royle et al. (2009)            | 1.60                                | –       |
| Ramesh et al. (2017)         | Western Shores, South Africa            | Coastal savanna             | Royle et al. (2009)            | 8.40                                | –       |
| Rostro-Garcia et al. (2018)  | Srepok wildlife sanctuary, Cambodia     | Dry deciduous forest        | Royle et al. (2009)            | 1.00                                | 0.40    |
| Selvan et al. (2014)         | Pakke Tiger Reserve, India              | Tropical forest             | Borchers & Efford (2008)       | 2.82                                | 1.20    |
| Strampelli et al. (2018)     | Xonghile Game Reserve, Mozambique       | Woodlands and thickets (savanna) | Borchers & Efford (2008)       | 2.59                                | 0.96    |
| Swanepoel, Somers & Dalerum (2015)  | Farming matrix, Waterberg, South Africa | Livestock and game farms    | Borchers & Efford (2008)       | 6.59                                | 5.20    |
| Swanepoel, Somers & Dalerum (2015)  | Lapalala Game Reserve, South Africa     | Mountain bushveld (dystrophic savanna) | Borchers & Efford (2008)       | 5.35                                | 2.93    |
preferred prey, (2) the existence of a compensation scheme that reimburses ranchers after depredation events on the LMNP edge (Braczkowski et al., 2020c), and (3) the functional extinction of lions in the region dating back to over a decade ago (at the time of publication there was only one male lion (≥10 years old) in this ecosystem, a vagrant thought to have come from Akagera National Park, in neighbouring Rwanda). LMNP is one of only two protected areas in Uganda with a population of impala, the most preferred prey of leopards (Hayward et al., 2006). The most recent studies implemented using distance sampling by Rannestad et al. (2006) and Kisame et al. (2018) found significant populations of impala within LMNP and on the adjacent cattle farmlands at 25.6 ± 4.8 individuals/km² in the 2003 study of Rannestad et al. (2006), and 15.3 and 16.6 individuals/km² in the 2014 and 2016 sampling periods of Kisame et al. (2018). Importantly Rannestad et al. (2006) also found a higher number of impala groups (80 vs 58) and total individuals (348 vs 255) in the community lands adjoining the park than within the national park in the wet season of 2003. Similarly, Kisame et al. (2018) estimated that nearly half of the impala population in the LMNP and surrounding ranches was found on non-protected land. Other densities of key leopard prey species estimated in this study included 3.8 ± 0.8 individuals/km² for bushbuck (higher densities outside) and warthogs (12.3 ± 2.9 individuals/km², densities lower outside national park, Rannestad et al., 2006). The availability of these species at relatively high densities both inside and beyond the edge of LMNP could be one reason for the densities of leopards and hyenas we observed in our study. It also remains unclear whether the functional extinction of lions in the LMNP has contributed to some level of release of leopards and spotted hyenas.

For example, from their study of leopard densities in three Kwazulu-Natal Parks, South Africa, Ramesh et al. (2017) found that where lion distribution overlapped spatially with leopards, densities of leopards decreased drastically. However this pattern of leopard suppression by lions was not observed in the Sabi Sand Game Reserve, a protected area system adjacent to the Kruger National Park where leopard-lion observations have been recorded since the 1970’s (Balme et al., 2017b).

Spotted hyenas and lions have an intricate relationship of facilitation and competition (Périquet, Fritz & Revilla, 2015). Unlike leopards, spotted hyenas do not show a negative correlation with lion presence in Africa (Périquet, Fritz & Revilla, 2015) despite intraguild predation and the negative impact that lions can have on hyena reproduction (Watts & Holekamp, 2008). Spotted hyenas may benefit from the presence of lions–and

| Study name               | Location                        | Habitat type                                      | Model used to estimate density | Density estimate (leopards/100 km²) | SD (SE) |
|-------------------------|---------------------------------|--------------------------------------------------|--------------------------------|-------------------------------------|---------|
| Swanepoel, Somers & Dalerum (2015) | Welgevonden Game Reserve, South Africa | Mountain bushveld (dystrophic savanna)            | Borchers & Efford (2008)        | 4.56                                | 1.35    |
| Shrestha et al. (2014)  | Parsa Wildlife Reserve, Nepal    | Dry deciduous forest                              | Efford (2004)                  | 3.78                                | 0.85    |
| Shrestha et al. (2014)  | Parsa Wildlife Reserve, Nepal    | Dry deciduous forest                              | Royle et al. (2009)            | 3.48                                | 0.83    |
| Williams et al., 2017  | Soutpansberg mountains, South Africa | Matrix of livestock farms, nature reserves, mountains | Royle et al. (2009)            | 5.34                                | 0.02    |
vice versa—due to the high dietary overlap between the species leading to scavenging and kleptoparasitic opportunities (Hayward, 2006; Davidson et al., 2019). Observed positive correlations in lion and spotted hyena density in many parts of Africa may also be a result of their similar preferred prey base. In Zambia, M’soka et al. (2016) found a high density of spotted hyenas in a lion-depleted ecosystem, though it was suggested that the observed density was driven by the availability of wildebeest, as in Höner et al. (2005). The spotted hyena densities we observed in our study were similar to an unpublished SECR study from uMkhuze Game Reserve, KwaZulu-Natal, South Africa (De Blocq, 2014), and a study from Botswana’s Moremi (Rich et al., 2019). Estimates of spotted hyena densities using non-SECR methods, from African savanna sites range widely from 2–20 individuals/100 km² in the Kruger National Park, South Africa (Mills, Juritz & Zucchini, 2001) to over 100 individuals/100 km² in the Ngorongoro Crater, Tanzania (Kruuk, 1972; Höner et al., 2005). The spotted hyena density from this study is similar those from protected areas in southern Africa but lower than those in other East African savannas (Holekamp & Dloniak, 2010). It is important to note that the majority of previous estimates have been produced using non-spatial methods (e.g., call-ups and mark-resight), and to our knowledge our study is one of the first to use a SECR approach for spotted hyena density estimation (Table 6). SECR densities are typically lower for large carnivores due to other methods making more generalized extrapolations over a given unit area (Noss et al., 2012) which may explain the difference between our results and those from other savanna systems in East Africa where non-spatial methods were used.

It is noteworthy that areas of high density between the species do not appear to overlap (Fig. 3). Previous studies have suggested that spotted hyenas can be significant kleptoparasites of leopard kills, forcing them to cache or avoid areas with high hyena density (Balme et al., 2017a; Davis et al., 2021). Similarly, another study detected low temporal overlap between leopards and spotted hyenas in Tanzania, which was postulated to be due to the avoidance of kleptoparasitism (Havmøller et al., 2020). Therefore, the avoidance of kleptoparasitism may drive the differences in space use between the species we detected in LMNP but would require further investigation.

### Table 6 Spotted hyena densities recorded in the literature.

| Study name                        | Location                                    | Habitat type                               | Model used to estimate density | Density estimate (hyenas/100 km²) | SD (SE) |
|-----------------------------------|---------------------------------------------|--------------------------------------------|--------------------------------|-----------------------------------|---------|
| Vissia, Wadhwa & van Langevelde (2021) | Central Tuli, Botswana                       | Riverine woodland and shrub savanna       | Borchers & Efford (2008)      | 14.90                             | 2.23    |
| Rich et al. (2019)                | Moremi Game Reserve and cattle matrix, Botswana | Semi-wooded savanna                      | Borchers & Efford (2008)      | 11.80                             | 2.60    |
| Briers-Low (2017)                 | Majete Game Reserve, Malawi                 | Tropical dry woodland/miombo savanna woodland | Royle et al. (2009)            | 2.69                              | 0.48    |
| De Blocq (2014)                   | uMkhuze Game Reserve, South Africa           | Semi-wooded savanna                      | Royle et al. (2009)            | 10.59                             | 2.1     |
| O’Brien & Kinnaird (2011)         | Mpala Ranch, Kenya                          | Semi-wooded savanna/cattle ranch          | Borchers & Efford (2008)      | 4.93                              | 1.7     |
Limitations and future monitoring of large carnivores in LMNP

Our study is limited by a lack of temporal replication. This is important as we could not generate critical population parameters such as emigration, immigration, birth and death (e.g., Karanth et al., 2006). These parameters are indicators of population trend and are ultimately required to ascertain the true trajectory of a given population. It should also be remembered that spotted hyenas live in fission-fusion clans and may move together in groups or singularly. It remains to be seen if this clan-living structure may cause biases in estimates of density and other parameters in our sampling situation. For example, López-Bao et al. (2018) show that wolf densities are not significantly affected by group living. Similarly, Bischof et al. (2020) suggest that if there are low to moderate levels of gregariousness observed in group living individuals, there is little overdispersion that occurs in the estimation of the detection function and scale parameter. However, if gregariousness is high, overdispersion may be observed in confidence intervals around parameter estimates, affecting the veracity of estimates. Although our study represents the first snapshot of this leopard and spotted hyena population, it is important as a baseline estimate from which future estimates can be made against (e.g., Balme, Slotow & Hunter, 2009). Our study also failed to quantify any relationships between hyenas and leopards, which in some sites have been shown to positively influence one another’s occupancy in a landscape (Comley et al., 2020).

Figure 3 Densities of leopards and hyenas in Lake Mburo. African leopard and spotted hyena detection frequencies (denoted in frequency by the size of spheres) and density estimates from our SECR models, LMNP, Uganda.

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There is a growing conflict between large carnivores and humans in the greater LMNP ecosystem (Braczkowski et al., 2020c). The impacts of spotted hyenas and leopards on cattle, sheep and goats in the Bahima pastoral lands adjacent to LMNP are significant, and leopards and spotted hyenas were the source of 98% (n = 1,102) of depredation events recorded between January 2009–December 2018 in the region (Braczkowski et al., 2020c). Other studies have highlighted spotted hyenas as a primary source of livestock loss, which combined with their negative public image, makes them vulnerable to retaliatory killing (Kissui, 2008; Holmern, Nyahongo & Røskaft, 2007). While spotted hyenas are behaviourally flexible, populations are slow to recover following even moderate reduction (Benhaiem et al., 2018). This pattern has also been observed for African leopards (e.g., Balme, Slotow & Hunter, 2009). For this reason, the continued monitoring of the LMNP spotted hyena and leopard population is crucial from a human-carnivore conflict perspective. Continued population monitoring of leopards is also critical in the context of trophy-hunting of leopard and leopard prey, which is allowed on properties adjoining the LMNP. Even though legal harvests of leopards in Uganda since 2007 have been low (17 skins, skulls and trophies exported from 2009–2017), and 28 leopards are available on quota country-wide annually (Braczkowski et al., 2015b), it is critical to monitor these populations annually or biannually as they can rapidly decline under even modest harvest pressures (Balme, Slotow & Hunter, 2010). The way in which quotas have been set in Uganda for leopards was also done using a non-robust method which related rainfall to leopard densities (CITES CoP 14 Proposal 3).

CONCLUSION

We aimed at providing the first leopard and spotted hyena population density estimates for the Lake Mburo ecosystem in Uganda, a small but regionally important national park with significant cattle farming on its edge. We found that leopard occur at a relatively high density of 6.3 individuals/100 km$^2$, probably due to a combination of factors such a high local prey density and an absence of lions. Spotted hyena densities were also relatively high, with several factors putatively at play, including abundance of prey including livestock, the absence of lions, and the general tolerance of hyenas for human disturbance. Our estimates form a robust baseline for future population monitoring to inform both the design of sustainable management offtakes, and conservation interventions for the two species in the region.

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Ralph Schenk, Nicholas Swanson and Merlin Swanson are employed by the Mihingo Lodge.

Author Contributions
- Aleksander Braczkowski conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Ralph Schenk conceived and designed the experiments, performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Dinal Samarasinghe performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Duan Biggs analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Allie Richardson performed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Nicholas Swanson conceived and designed the experiments, performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Merlin Swanson conceived and designed the experiments, performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Arjun Dheer performed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Julien Fattebert conceived and designed the experiments, performed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

Animal Ethics
The following information was supplied relating to ethical approvals (i.e., approving body and any reference numbers):
This was a camera trap study and did not involve either the handling, capture or immobilisation of any vertebrates.

Field Study Permissions
The following information was supplied relating to field study approvals (i.e., approving body and any reference numbers):
This research was granted approval by the Uganda Wildlife Authority under permit number: UWA/COD/96/05 as approved by the Executive Director Mr. Stephen Masaba.

**Data Availability**
The following information was supplied regarding data availability:
All data and code are available in the Supplemental Files.

**Supplemental Information**
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.12307#supplemental-information.

**REFERENCES**

Bahaa-el-din L, Sollmann R, Hunter LT, Slotow R, Macdonald DW, Henschel P. 2016. Effects of human land-use on Africa’s only forest-dependent felid: the African golden cat Caracal aurata. *Biological Conservation* **199**:1–9 DOI 10.1016/j.biocon.2016.04.013.

Bailey TN. 1993. *The African leopard*. New York: Columbia University Press.

Balme GA, Hunter L, Braczkowski AR. 2012. Applicability of age-based hunting regulations for African leopards. *PLOS ONE* **7**(4):e35209 DOI 10.1371/journal.pone.0035209.

Balme GA, Hunter LTB, Slotow R. 2009. Evaluating methods for counting cryptic carnivores. *Journal of Wildlife Management* **73**:433–441 DOI 10.2193/2007-368.

Balme GA, Miller JR, Pitman RT, Hunter LT. 2017a. Caching reduces kleptoparasitism in a solitary, large felid. *Journal of Animal Ecology* **86**(3):634–644 DOI 10.1111/1365-2656.12654.

Balme GA, Pitman RT, Robinson HS, Miller JR, Funston PJ, Hunter LT. 2017b. Leopard distribution and abundance is unaffected by interference competition with lions. *Behavioral Ecology* **28**(5):1348–1358 DOI 10.1093/beheco/arx098.

Balme G, Rogan M, Thomas L, Pitman R, Mann G, Whittington-Jones G, Hunter L. 2019. Big cats at large: Density, structure, and spatio-temporal patterns of a leopard population free of anthropogenic mortality. *Population Ecology* **61**(3):256–267.

Balme GA, Slotow R, Hunter LTB. 2009. Impact of conservation interventions on the dynamics and persistence of a persecuted leopard population. *Biological Conservation* **142**:2681–2690 DOI 10.1016/j.biocon.2009.06.020.

Balme GA, Slotow ROB, Hunter LT. 2010. Edge effects and the impact of non-protected areas in carnivore conservation: leopards in the Phinda-Mkhuze Complex, South Africa. *Animal Conservation* **13**(3):315–323 DOI 10.1111/j.1469-1795.2009.00342.x.

Benhaiem S, Marescot L, East ML, Kramer-Schadt S, Gimenez O, Lebreton JD, Hofer H. 2018. Slow recovery from a disease epidemic in the spotted hyena, a keystone social carnivore. *Communications Biology* **1**(1):1–12 DOI 10.1038/s42003-018-0197-1.

Bischof R, Dupont P, Milleret C, Chipperfield J, Royle JA. 2020. Consequences of ignoring group association in spatial capture-recapture analysis. *Wildlife Biology* **1**:1–10 DOI 10.2981/wlb.00649.

Borah J, Sharma T, Das D, Rabha N, Kakati N, Basumatary A, Vattakaven J. 2014. Abundance and density estimates for common leopard Panthera pardus and clouded leopard Neofelis nebulosa in Manas National Park, Assam, India. *Oryx* **48**(1):149–155.

Borchers DL, Efford MG. 2008. Spatially explicit maximum likelihood methods for capture-recapture studies. *Biometrics* **64**(2):377–385 DOI 10.1111/j.1541-0420.2007.00927.x.
Braczkowski AR, Balme GA, Dickman A, Fattebert J, Johnson P, Dickerson T, Macdonald DW, Hunter L. 2016. Scent lure effect on camera-trap based leopard density estimates. *PLOS ONE* 11(4):e0151033 DOI 10.1371/journal.pone.0151033.

Braczkowski AR, Balme GA, Dickman A, Macdonald DW, Fattebert J, Dickerson T, Johnson P, Hunter L. 2015a. Who bites the bullet first? The susceptibility of leopards Panthera pardus to trophy hunting. *PLOS ONE* 10(4):e0123100 DOI 10.1371/journal.pone.0123100.

Braczkowski AR, Dickman A, Macdonald DW, Johnson PJ, Balme GA, Lindsey PA, Hunter LT. 2015b. Rosettes, remingtons and reputation: establishing potential determinants of leopard (Panthera pardus) trophy prices across Africa. *African Journal of Wildlife Research* 45(2):158–168 DOI 10.3957/056.045.0158.

Braczkowski A, Fattebert J, Schenk R, O’Brien C, Biggs D, Maron M. 2020c. Evidence for increasing human-wildlife conflict despite a financial compensation scheme on the edge of a Ugandan National Park. *Conservation Science and Practice* 2(12):e309 DOI 10.1111/csp2.309.

Braczkowski A, Gopalaswamy AM, Nsubuga M, Allan J, Biggs D, Maron M. 2020b. Detecting early warnings of pressure on an African lion (Panthera leo) population in the Queen Elizabeth Conservation Area, Uganda. *Ecological Solutions and Evidence* 1(1):e12015 DOI 10.1002/2688-8319.12015.

Briers-Louw WD. 2017. Ecology of apex predators in Majete Wildlife Reserve. Malawi (Doctoral dissertation). South Africa: Stellenbosch University.

Comley J, Joubert CJ, Mqqatsa N, Parker DM. 2020. Lions do not change rivers: complex African savannas preclude top-down forcing by large carnivores. *Journal for Nature Conservation* 56:125844 DOI 10.1016/j.jnc.2020.125844.

Davidson Z, Dupuis-Desormeaux M, Dheer A, Pratt L, Preston E, Gilicho S, Mwololo M, Chege G, MacDonald S, Doncaster CP. 2019. Borrowing from Peter to pay Paul: managing threatened predators of endangered and declining prey species. *PeerJ* 7(2):e7916 DOI 10.7717/peerj.7916.

Davis RS, Yarnell RW, Gentle LK, Uzal A, Mgoolla WO, Stone EL. 2021. Prey availability and intraguild competition regulate the spatiotemporal dynamics of a modified large carnivore guild. *Ecology and Evolution* 11(12):7890–7904 DOI 10.1002/ece3.7620.

De Blocq AD. 2014. Estimating spotted hyaena (Crocuta crocuta) population density using camera trap data in a spatially explicit capture-recapture framework (Doctoral dissertation, University of Cape Town).

Devens C, Tshabalala T, McManus J, Smuts B. 2018. Counting the spots: The use of a spatially explicit capture-recapture technique and GPS data to estimate leopard (Panthera pardus) density in the Eastern and Western Cape, South Africa. *African Journal of Ecology* 56(4):850–859.

Dey S, Delampady M, Gopalaswamy AM. 2019. Bayesian model selection for spatial capture-recapture models. *Ecology and Evolution* 9(20):11569–11583 DOI 10.1002/ece3.5551.

du Preez BD, Loveridge AJ, Macdonald DW. 2014. To bait or not to bait: a comparison of camera-trapping methods for estimating leopard Panthera pardus density. *Biological Conservation* 176:153–161 DOI 10.1016/j.biocon.2014.05.021.

Duangchantrasiri S, Umponjan M, Simcharoen S, Pattanavibool A, Chaiwattana S, Maneerat S, Kumar NS, Jathanna D, Srivathsa A, Karanth KU. 2016. Dynamics of a low-density tiger population in Southeast Asia in the context of improved law enforcement. *Conservation Biology* 30(3):639–648 DOI 10.1111/cobi.12655.

Efford MG. 2004. Density estimation in live– trapping studies. *Oikos* 106:598–610.
Efford MG. 2011. Secr: spatially explicit capture-recapture models. R package version 3.0.2. Available at https://cran.r-project.org/web/packages/secr/index.html (accessed 1 October 2013).

Efford MG. 2019. Non-circular home ranges and the estimation of population density. Ecology 100(2):e02580 DOI 10.1002/ecy.2580.

Elliot NB, Gopalaswamy AM. 2017. Toward accurate and precise estimates of lion density. Conservation Biology 31(4):934–943 DOI 10.1111/cobi.12878.

Fattebert J, Balme GA, Robinson HS, Dickerson T, Slotow R, Hunter LT. 2016. Population recovery highlights spatial organization dynamics in adult leopards. Journal of Zoology 299(3):153–162 DOI 10.1111/jzo.12344.

Gelman A, Rubin DB. 1992. Inference from iterative simulation using multiple sequences. Statistical Science 7(4):457–472 DOI 10.1214/ss/1177011136.

Gopalaswamy AM, Royle JA, Delampady M, Nichols JD, Karanth KU, Macdonald DW. 2012a. Density estimation in tiger populations: combining information for strong inference. Ecology 93(7):1741–1751 DOI 10.1890/11-2110.1.

Gopalaswamy AM, Royle JA, Hines JE, Singh P, Jathanna D, Kumar NS, Karanth KU. 2012b. Program SPACECAP: software for estimating animal density using spatially explicit capture-recapture models. Methods in Ecology and Evolution 3(6):1067–1072 DOI 10.1111/j.2041-210X.2012.00241.x.

Goswami R, Ganesh T. 2014. Carnivore and herbivore densities in the immediate aftermath of ethno-political conflict: the case of Manas National Park, India. Tropical Conservation Science 7(3):475–487.

Hamilton WJ, Tilson RL, Frank LG. 1986. Sexual monomorphism in spotted hyenas, Crocuta crocuta. Ethology 71(1):63–73 DOI 10.1111/j.1439-0310.1986.tb00570.x.

Havmøller RW, Jacobsen NS, Scharff N, Rovero F, Zimmermann F. 2020. Assessing the activity pattern overlap among leopards (Panthera pardus), potential prey and competitors in a complex landscape in Tanzania. Journal of Zoology 311(3):175–182 DOI 10.1111/jzo.12774.

Hayward MW. 2006. Prey preferences of the spotted hyaena (Crocuta crocuta) and degree of dietary overlap with the lion (Panthera leo). Journal of Zoology 270(4):606–614 DOI 10.1111/j.1469-7998.2006.00183.x.

Hayward MW, Henschel P, O’Brien J, Hofmeyr M, Balme G, Kerley G. 2006. Prey preferences of the leopard (Panthera pardus). Journal of Zoology 270(2):298–313 DOI 10.1111/j.1469-7998.2006.00139.x.

Hedges J, Lam WY, Campos-Arceiz A, Rayan DM, Laurance WF, Latham CJ, Clements GR. 2015. Melanistic leopards reveal their spots: infrared camera traps provide a population density estimate of leopards in Malaysia. The Journal of Wildlife Management 79(5):846–853.

Henschel P, Malanda GA, Hunter L. 2014. The status of savanna carnivores in the Odzala-Kokoua National Park, northern Republic of Congo. Journal of Mammalogy 95(4):882–892 DOI 10.1644/13-MAMM-A-306.

Holekamp KE, Dloniak SM. 2010. Intraspecific variation in the behavioral ecology of a tropical carnivore, the spotted hyena. In: Advances in the Study of Behavior, Vol. 42. Cambridge: Academic Press, 189–229.

Holmern T, Nyahongo J, Roskaft E. 2007. Livestock loss caused by predators outside the Serengeti National Park, Tanzania. Biological Conservation 135(4):518–526 DOI 10.1016/j.biocon.2006.10.049.

Höner OP, Wachter B, East ML, Runyoro VA, Hofer H. 2005. The effect of prey abundance and foraging tactics on the population dynamics of a social, territorial carnivore, the spotted hyena. Oikos 108(3):544–554 DOI 10.1111/j.0030-1299.2005.13533.x.
Jacobson AP, Gerngross P, Lemeris JR Jr, Schoonover RF, Anco C, Breitenmoser-Würsten C, Durant SM, Farhadinia MS, Henschel P, Kamler JF, Laguardia A. 2016. Leopard (Panthera pardus) status, distribution, and the research efforts across its range. PeerJ 4(1):e1974 DOI 10.7717/peerj.1974.

Karanth KU. 1995. Estimating tiger Panthera tigris populations from camera-trap data using capture—recapture models. Biological Conservation 71(3):333–338 DOI 10.1016/0006-3207(94)00057-W.

Karanth KU, Nichols JD. 1998. Estimation of tiger densities in India using photographic captures and recaptures. Ecology 79(8):2852–2862 DOI 10.1890/0012-9658(1998)079[2852:EOTDII]2.0.CO;2.

Karanth KU, Nichols JD, Kumar NS, Hines JE. 2006. Assessing tiger population dynamics using photographic capture-recapture sampling. Ecology 87(11):2925–2937 DOI 10.1890/0012-9658(2006)87[2925:ATPDUP]2.0.CO;2.

Kisame FE, Wanyama F, Buhanga E, Rwetsiba A. 2018. Ground Counts for medium to large mammals in Lake Mburo Conservation Area. Uganda Wildlife Authority, Kampala, Uganda. Available at https://www.ugandawildlife.org/phocadownload/conservation-publications/survey-reports/Ground-counts-for-LMNP-2018.pdf.

Kissui BM. 2008. Livestock predation by lions, leopards, spotted hyenas, and their vulnerability to retaliatory killing in the Maasai steppe, Tanzania. Animal conservation 11(5):422–432.

Kittle AM, Watson AC, Fernando TSP. 2017. The ecology and behaviour of a protected area Sri Lankan leopard (Panthera pardus kotiya) population. Tropical Ecology 58(1):71–86.

Kruuk H. 1972. The spotted hyena: a study of predation and social behavior. Chicago: University of Chicago Press.

López-Bao JV, Godinho R, Pacheco C, Lema FJ, García E, Llaneza L, Palacios V, Jiménez J. 2018. Toward reliable population estimates of wolves by combining spatial capture-recapture models and non-invasive DNA monitoring. Scientific Reports 8(1):1–8 DOI 10.1038/s41598-018-20675-9.

Menaut JC. 1983. The vegetation of African savannas. In: Bourlière F, ed. Tropical Savannas. Vol. 13. Amsterdam: Elsevier, 109–150.

Mills MGL, Juritz JM, Zucchini W. 2001. Estimating the size of spotted hyaena (Crocuta crocuta) populations through playback recordings allowing for non-response. In: Animal Conservation Forum, Cambridge: Cambridge University Press, 335–343.

Mithathapala S, Seidensticker J, Phillips LG, Fernando SBU, Smallwood JA. 1989. Identification of individual leopards (Panthera pardus kotiya) using spot pattern variation. Journal of Zoology 218(4):527–536 DOI 10.1111/j.1469-7998.1989.tb04996.x.

Moe SR, Loe LE, Jessen M, Okullo P. 2016. Effects of mammalian herbivores and termites on the performance of native and exotic plantation tree seedlings. Journal of Applied Ecology 53(2):323–331 DOI 10.1111/1365-2664.12577.

M’soka J, Creel S, Becker MS, Droge E. 2016. Spotted hyaena survival and density in a lion depleted ecosystem: the effects of prey availability, humans and competition between large carnivores in African savannahs. Biological Conservation 201(S1):348–355 DOI 10.1016/j.biocon.2016.07.011.

Ngoprasert D, Lynam AJ, Gale GA. 2017. Effects of temporary closure of a national park on leopard movement and behaviour in tropical Asia. Mammalian Biology 82(1):65–73.

Noss AJ, Gardner B, Maffei I, Cuéllar E, Montaño R, Romero-Muñoz A, Sollman R, O’Connell AF. 2012. Comparison of density estimation methods for mammal populations with...
camera traps in the Kaa-I ya del Gran Chaco landscape. *Animal Conservation* **15**(5):527–535 DOI 10.1111/j.1469-1795.2012.00545.x.

Ochieng A, Ahebwa WM, Visseren-Hamakers IJ. 2015. Hunting for conservation? The re-introduction of sport hunting in Uganda examined. In: *Institutional Arrangements for Conservation, Development and Tourism in Eastern and Southern Africa*, Dordrecht: Springer, 139–155.

O’Brien TG, Kinnaird MF. 2011. Density estimation of sympatric carnivores using spatially explicit capture-recapture methods and standard trapping grid. *Ecological Applications* **21**(8):2908–2916 DOI 10.1890/10-2284.1.

O’Brien CJ, Braczkowski AR, Beyer HL, Carter NH, Watson JE, McDonald-Madden E. 2018. The contribution of predators and scavengers to human well-being. *Nature Ecology & Evolution* **2**(2):229–236 DOI 10.1038/s41559-017-0421-2.

Périquet S, Fritz H, Revilla E. 2015. The Lion King and the Hyaena Queen: large carnivore interactions and coexistence. *Biological Reviews* **90**(4):1197–1214 DOI 10.1111/brv.12152.

Qi J, Shi Q, Wang G, Li Z, Sun Q, Hua Y, Jiang G. 2015. Spatial distribution drivers of Amur leopard density in northeast China. *Biological Conservation* **191**:258–265.

R Development Core Team. 2019. R: a language and environment for statistical computing. Vienna, Austria: R foundation for statistical computing. Available at https://www.R-project.org/.

Rahman DA, Rianti P, Muhiban M, Muhtarom A, Rahmat UM, Santosa Y, Aulagnier S. 2018. Density and spatial partitioning of endangered sympatric Javan leopard (Felidae) and dholes (Canidae) in a tropical forest landscape. *Journal of Vertebrate Biology* **67**(3–4):207–219.

Ramesh T, Kalle R, Rosenlund H, Downs CT. 2017. Low leopard populations in protected areas of Maputaland: a consequence of poaching, habitat condition, abundance of prey, and a top predator. *Ecology and Evolution* **7**(6):1964–1973 DOI 10.1002/ece3.2771.

Rannestad OT, Danielsen T, Moe SR, Stokke S. 2006. Adjacent pastoral areas support higher densities of wild ungulates during the wet season than the Lake Mburo National Park in Uganda. *Journal of Tropical Ecology* **22**(6):675–683 DOI 10.1017/S0266467406003610.

Rayan DM, Linkie M. 2015. Conserving tigers in Malaysia: a science-driven approach for eliciting conservation policy change. *Biological Conservation* **184**:18–26 DOI 10.1016/j.biocon.2014.12.024.

Rich LN, Miller DA, Muñoz DJ, Robinson HS, McNutt JW, Kelly MJ. 2019. Sampling design and analytical advances allow for simultaneous density estimation of seven sympatric carnivore species from camera trap data. *Biological Conservation* **233**:12–20 DOI 10.1016/j.biocon.2019.02.018.

Rostro-García S, Kamler JF, Crouthers R, Sopheak K, Prum S, In V, Macdonald DW. 2018. An adaptable but threatened big cat: density, diet and prey selection of the Indochinese leopard (Panthera pardus delacouri) in eastern Cambodia. *Royal Society open science* **5**(2):171187.

Royle JA, Karanth KU, Gopalaswamy AM, Kumar NS. 2009. Bayesian inference in camera trapping studies for a class of spatial capture-recapture models. *Ecology* **90**(11):3233–3244 DOI 10.1890/08-1481.1.

Selvan KM, Lyngdoh S, Habib B, Gopi GV. 2014. Population density and abundance of sympatric large carnivores in the lowland tropical evergreen forest of Indian Eastern Himalayas. *Mammalian Biology* **79**(4):254–258.

Shrestha R, Karki J, Thapa GJ, Subedi N, Pradhan NMB, Dhakal M, Khanal P, Kelly MJ. 2014. Leopard Panthera pardus fusca density in the seasonally dry, subtropical forest in the Bhabhar of Terai Arc, Nepal. *Advances in Ecology* **2014** DOI 10.1155/2014/286949.
Sollmann R, Furtado MM, Gardner B, Hofer H, Jácomo AT, Tôrres NM, Silveira L. 2011. Improving density estimates for elusive carnivores: accounting for sex-specific detection and movements using spatial capture-recapture models for jaguars in central Brazil. *Biological Conservation* **144**(3):1017–1024 DOI 10.1016/j.biocon.2010.12.011.

Strampelli P, Andresen L, Everatt KT, Somers MJ, Rowcliffe JM. 2018. Habitat use responses of the African leopard in a human-disturbed region of rural Mozambique. *Mammalian Biology* **89**:14–20.

Swanepoel LH, Somers MJ, Dalerum F. 2015. Density of leopards Panthera pardus on protected and non-protected land in the Waterberg Biosphere, South Africa. *Wildlife Biology* **21**(5):263–268.

Tierney L. 1994. Markov chains for exploring posterior distributions. *The Annals of Statistics* **22**(4):1701–1728 DOI 10.1214/aos/1176325750.

Tweheyo M, Tumusiime DM, Turyahabwe N, Asimwe A, Orikiriza L. 2012. Wildlife damage and control methods around Lake Mburo National Park, Uganda. *International Journal of Pest Management* **58**(1):25–31 DOI 10.1080/09670874.2011.641605.

Uganda Wildlife Authority. 2010. Strategic action plan for large carnivore conservation in Uganda 2010–2020. Kampala, Uganda. Available at [http://www.catsg.org/fileadmin/files/News/Conservation_Center/3.4_Strategies__Action_Plans/UWA_Strategic_Action_Plan_for_large_Carnivore_Conservation_2010-2020.pdf](http://www.catsg.org/fileadmin/files/News/Conservation_Center/3.4_Strategies__Action_Plans/UWA_Strategic_Action_Plan_for_large_Carnivore_Conservation_2010-2020.pdf).

Van de Weghe JP. 1990. *Akagera, land of water, grass and fire*. Brussels: WWF.

Van der Meer E, Badza MN, Ndhlovu A. 2016. Large carnivores as tourism flagship species for the Zimbabwe component of the Kavango Zambezi Transfrontier Conservation Area. *African Journal of Wildlife Research* **46**(2):121–134 DOI 10.3957/056.046.0121.

Venter O, Sanderson EW, Magrach A, Allan JR, Beher J, Jones KR, Possingham HP, Laurance WF, Wood P, Fekete BM, Levy MA. 2016. Sixteen years of change in the global terrestrial human footprint and implications for biodiversity conservation. *Nature Communications* **7**(1):1–11 DOI 10.1038/ncomms12558.

Vissia S, Wadhwa R, van Langevelde F. 2021. Co-occurrence of high densities of brown hyena and spotted hyena in central Tuli, Botswana. *Journal of Zoology* **314**(2):143–150 DOI 10.1111/jzo.12873.

Watts HE, Holekamp KE. 2008. Interspecific competition influences reproduction in spotted hyenas. *Journal of Zoology* **276**(4):402–410 DOI 10.1111/j.1469-7998.2008.00506.x.

White GC, Burnham KP. 1999. Program MARK: survival estimation from populations of marked animals. *Bird Study* **46**(sup1):S120–S139 DOI 10.1080/00063659909477239.

Williams PJ, Hooten MB, Womble JN, Bower MR. 2017. Estimating occupancy and abundance using aerial images with imperfect detection. *Methods in Ecology and Evolution* **8**(12):1679–1689 DOI 10.1111/2041-210X.12815.

Woodroffe R, Ginsberg JR. 1998. Edge effects and the extinction of populations inside protected areas. *Science* **280**(5372):2126–2128 DOI 10.1126/science.280.5372.2126.