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# Threats to Site Occupation of Carnivores: A Spatiotemporal Encroachment of Non-native Species on the Native Carnivore Community in A Human-dominated Protected Area

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Interspecific interactions of commensal non-native species such as domestic dogs and livestock with native wildlife are evident issues in protected areas (PAs). We studied spatiotemporal interactions by combining camera trap photographic sampling over three years. We used a generic multiseason occupancy and cooccurrence analysis and kernel density estimates of temporal activity. We accumulated a total of 1,305 and 1,557 independent photo-captures respectively for non-native and carnivore species during 26,216 trap nights. We found that non-native and carnivore species did not show substantial changes in occupancy rate over time. Yet both kinds of species were frequently detected. Carnivores had lower values of occupancy equilibrium than non-native species in seasons one and two. Domestic dogs directly occurred with native carnivores (except with leopards in season one), while the human and livestock presence displayed direct (Species Interaction Factors > 1) and indirect (Species Interaction Factors close to 1) cooccurrence, respectively, with the leopard and two mesocarnivores. The leopard cat was the least spatially influenced carnivore by the non-native species interactions. Furthermore, the leopard had higher temporal overlap (high  $\Delta_4$ ) with all non-native species than the leopard cat and red fox (low  $\Delta_4$ ). Our study exposes the negative impact of free-ranging non-native species across this protected area on native carnivore occupancy. An evaluation of the interconnections among non-native and carnivore species across diverse PA management regimes is crucial to develop robust landscape-scale conservation strategies.

**Key words:** Conservation and management, Non-native and carnivore species, Overlap, Protected area, Spatiotemporal patterns.

#### BACKGROUND

Protected Areas (PAs) are the most widely-known and well-accepted strategy for protecting ecosystems and biodiversity (Chape et al. 2005; Dudley 2008; Mwakatobe et al. 2013). Protected areas have proven to be effective in protecting species' habitats from landuse changes occurring outside of the protected areas (Andam et al. 2008). They represent one of the most significant global surface areas dedicated to a common

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goal (after areas used for food production), that of preserving global biodiversity (Palomo et al. 2014). In their regular management, PAs typically face many difficulties (Cromsigt et al. 2013; Zaman et al. 2020). One of the most common issues is human interactions with wildlife, which can manifest in different forms (Treves and Karanth 2003; Okello et al. 2014) and often generates misunderstandings between PA managers and the local population. Carnivores need large living spaces for their daily activities, which include hunting and walking (Treves and Karanth 2003). To meet these needs, large carnivores typically use resources outside of PAs (Hansen et al. 2002; Treves and Karanth 2003), potentially bringing them into contact with human landuse activities such as livestock and crop production (Morehouse and Boyce 2017). Conflicts between nonnative and native carnivore species can lead to largescale ecosystem disturbances.

The presence of non-native species in PAs poses a threat to native carnivorous species in both space usage and temporal activities. The human activities within and around PAs within landscapes lead to the presence of domestic dogs (Farris et al. 2017) and livestock species (Vanak and Gompper 2010), which have harmful consequences on wildlife. The most cited effects of this include carnivores preying upon invasive livestock (Treves and Karanth 2003), causing non-native species to destabilise (Morehouse and Boyce 2017), and spreading diseases to native carnivore species in the PAs (Okello et al. 2014). They also create edge effects and habitat fragmentation (Gerber et al. 2012a; Sleeman 2013; Vanak et al. 2013; Brodie et al. 2015; Chanchani et al. 2016). Non-native species (especially carnivores) often significantly increase pressure on native carnivores by modifying their spatiotemporal activity patterns and habitat use (Gerber et al. 2012b; Farris et al. 2015) and decreasing prey availability (Young et al. 2011). When non-native species are livestock, native carnivores can alter their natural diet because herds will seem like easy prey (Okello et al. 2014; Ciucci et al. 2018). Cited effects (disease transmission, spatio-temporal destabilisation, prey decrease, preying on livestock) can lead to decreased carnivore survival within the habitat and generate negative consequences on the whole landscape with the reduction of the carrying capacity in the ecosystem. Carnivores are greatly sensitive to disturbances inside and outside of PAs and are negatively affected by edge effects and human presence (poaching or human-wildlife conflict) occurring around PAs (Wearn et al. 2012; Sleeman 2013; Hua et al. 2020). Therefore, carnivore population wellbeing is expected to be driven by species-specific tolerance to microenvironment change, habitat fragmentation and PA edge effects, human presence, non-native species density,

and prey availability. All these facts are essential in and around PAs and must be well managed to ensure species sustainability (Farris et al. 2017).

The Tieqiaoshan Provincial Nature Reserve (TPNR) contains some of the largest North China leopard (*Panthera pardus japonensis*, Gray 1862) populations in China (Vitekere et al. 2020a; Zhu et al. 2021). Most of the North China leopard populations are within 22 Nature Reserves situated in the Taihang Mountains and some nearby regions (Song et al. 2014), but most of these PAs have not been surveyed and accurate population estimates are lacking.

Thus, little is known about this leopard subspecies living in a guild with two mesocarnivores, the leopard cat (Prionailurus bengalensis, Kerr 1792) and the red fox (Vulpes vulpes, Linnaeus 1758), within the TPNR ecosystem (Hua et al. 2020). This landscape hosts a considerable human population density (local people), with livestock activities facilitating the invasion of the integral conservation zone of the TPNR by dogs, livestock, and humans (e.g., Farris et al. 2017). Few studies have examined the conservation status of these carnivores and the effects of non-native species introduced by the shepherds in the PAs. Hua et al. (2020) highlighted the effect of human presence on the detectability of these carnivores. Still, the longterm effects and the coexistence with non-native species remain unstudied within the TPNR landscape.

We carried out a multi-year study on the coexistence between non-native species-the domestic dog (dog), livestock and herdsmen (human)-and native carnivores-North China leopard (leopard), leopard cat and red fox (fox)-within TPNR, one of the Taihang Mountains landscapes. We first hypothesized that carnivores would not depict the stability of occupancy contrarily to non-native species' over the years. We also hypothesized that all three native carnivores would have a real coexistence defined as "apparent co-occurrence" or "direct interaction" with non-native species, and they would portray increasing Species Interaction Factors (SIF: a parameter describing the spatial interactions of two species in an area, previously computed by Alexander et al. (2016) and Farris et al. (2020)) over the years. Human detection, particularly, would markedly affect native species since the two others (livestock and dog) are human presence-dependent. We finally hypothesized that non-native and carnivore species would not temporally overlap, since non-native and native carnivore species diurnal and nocturnal, respectively.

Studying these multi-year spatiotemporal interactions in a landscape with human invasion (bringing invasive species) will help assess the effectiveness of management policies for the

#### MATERIALS AND METHODS

#### Study Area

The TPNR is a reserve in China that hosts the North China leopard sub-species. This PA has GPS coordinates:  $111^{\circ}25$ 'E to  $114^{\circ}17$ 'E and  $36^{\circ}39$ 'N to  $38^{\circ}06$ 'N (Fig. 1), with an elevation ranging from 1300 to 1827 m. The TPNR is a Protected Area, approved by the provincial administration by document No. 124. Officially established in 2009, this PA was assigned a total of  $353.52 \text{ km}^2$  (Zhu et al. 2021) divided into an area of integral protection (139.5 km<sup>2</sup>), a buffer zone (74.2 km<sup>2</sup>), and a multi-use zone. The annual average of rainfall varied from 500 mm to 700 mm, with heavy rains in the period July-September (Hua et al. 2020). This area did not present high daily and yearly thermal amplitude, with mean temperatures of  $10^{\circ}$ C and  $6^{\circ}$ C, respectively (Zheng et al. 2009). One of the principal

characteristics of this PA is that it hosts humans (Zhu et al. 2021), an estimated 2,000 of them, meaning that the PA had always experienced disturbances from humans (Hua et al. 2020). A significant part of the area is a mixture of primary and secondary forests, and the remaining parts are shrubs. Wildlife is abundant within the PA, including mammals (carnivores: leopard, leopard cat, fox, badger, etc.; artiodactyl: many kinds of deer and the wild boar), and a diversified fauna of reptiles, amphibians, and birds (Song et al. 2014).

#### Methods

#### **Data Collection**

We collected data on the presence and absence of three native carnivore species and three non-native species for a total of 383 days over 3 years: 130 (March– July 2017), 119 (September–December 2018) and 134 (March–June 2019), comprising three sampling periods (SP<sub>1</sub>, SP<sub>2</sub>, and SP<sub>3</sub>) and two interseasons (IS<sub>1</sub> and IS<sub>2</sub>), according to Farris et al. (2017). We used two brands of cameras: the Eastern Red Hawk E1B 6210M (Shenzhen Ereagle Technology Co. Ltd, Shenzhen, China) and the LTL6210MM (LTL Acorn Trail Camera, United



Fig. 1. The data collection sites within the Tieqiaoshan Provincial Nature Reserve and villages housing herdsmen in and around the Protected Area.

Kingdom). Both brands were triggered remotely by an infrared sensor to record animal activity, following Karanth et al. (2003) and Swann et al. (2011). SP<sub>1</sub> contained 81 cameras while SP<sub>2</sub> and SP<sub>3</sub> contained 62 cameras. Cameras were deployed within 27 4 km × 4 km quadrants in the study area, attached to trees at an average height of 0.5 m. Each data collection site had two or three cameras installed to face each other. Where possible, cameras were placed on trails, but otherwise they were placed at points thought to maximise the visibility of the animals. Cameras were set to capture data with the time and date automatically displayed on the photos (*e.g.*, Barrull et al. 2014). Each camera operated for at least 100 consecutive days in each sampling period.

#### **Data Analysis**

#### Spatial patterns

We first aggregated data from more than 100 days of capture within each sampling period into multiple shorter intervals. In SP<sub>1</sub> and SP<sub>3</sub>, each survey (sampling occasion) was two weeks long, while in SP<sub>2</sub> is was 10 days. This data aggregation allowed standardisation of the three sampling periods (*e.g.*, Bu et al. 2016; Farris et al. 2017) and provided multiple surveys within each sampling period, as required for occupancy modelling (Alexander et al. 2016; MacKenzie et al. 2006). We considered a species to be detected if it was present at a site in an interval of aggregated days. Otherwise, it was not detected.

We then performed two different occupancy analyses to investigate interactions between native carnivore and non-native species: the multiseason occupancy and the multiseason co-occurrence analyses. Occupancy analysis is based on collecting occurrence data for studied species within data collection sites. Species can be imperfectly detected; therefore, *n* sites are visited on t sampling occasions, and the presence/ absence of each targeted species is recorded on each occasion (MacKenzie et al. 2003). We used the software PRESENCE (version 5.8 < 130315.0823 > by James E Hines). This modelling using presence/absence data for different species allowed us to estimate changes in occupancy for all species over three years to test if there is an apparent spatial co-occurrence between native carnivores and non-native species, and finally to assess the influence of non-native species on the occupancy and detection probabilities of native species.

In the first step (multiseason occupancy), parameters estimated can depict variations across time: the probability of occupancy ( $\psi$ ), the probability of detection (p), the colonisation rate ( $\gamma$ ), and the local extirpation rate ( $\varepsilon$ ); all estimate values were accompanied by their standard errors. We estimated these parameters to determine the general trend of site occupation of species across years. Following MacKenzie et al. (2006), two derived parameters were also computed for the interpretation of the occupancy dynamics: (1) the rate of change in occupancy:

$$\lambda'_{t} = \frac{\psi_{t+1}/(1-\psi_{t+1})}{\psi_{t}/(1-\psi_{t})} \qquad (eq_{1} \text{ MacKenzie et al. 2006})$$

(2) the occupancy equilibrium:

 $\psi_{equilibrium} = \gamma / (\gamma + \varepsilon)$  (eq<sub>2</sub> MacKenzie et al. 2006)

In the second step (multiseason co-occurrence), we computed the co-occupancy and co-detection parameters (Table 1), and the SIF ( $\varphi$ ). This latter parameter describes the species' interactions in an area, where  $\varphi < 1$  indicates no interaction or avoidance,  $\varphi > 1$ indicates direct interaction or apparent co-occurrence and  $\varphi = 1$  indicates independent or indirect cooccurrence (Alexander et al. 2016; Farris et al. 2020). We then used the Chi-squared test of independence to verify if the occupancy of a carnivore computed as single species is independent from the same species' occupancy in a co-occurrence (two species) pattern.

#### **Temporal patterns**

The time that a species was captured was automatically stamped on species photographs. We removed any subsequent photos of the same species at the same site that occurred within 30 minutes to minimize pseudo-replication biases (e.g., Monterroso et al. 2013; Sunarto et al. 2015; Farris et al. 2017). All photos of a species remaining in the dataset after the application of this filter were considered to be independent observations. We compared the day and night visibility of species by calculating the proportion of detections that occurred in two intervals: 06:00-18:00 for daytime and 18:00-06:00 for night-time to test the species' "nocturnality." A species was categorized to prefer night-time or daytime when its detection rate was  $\geq$  70%, respectively, in the 06:00–18:00 and 18:00-6:00 time interval (e.g., Dias et al. 2018; Hua et al. 2020). To analyze the temporal activity overlap between non-native species and native carnivore, we calculated the kernel density estimates (KDEs) using the R package overlap (Oliveira-Santos et al. 2013; Meredith and Ridout 2014). We assessed the temporal overlap coefficient ( $\Delta_4$ ) to compare activity patterns of all pairwise species combinations of non-native species and native carnivores. The overlap coefficient is a metric that ranges from zero, meaning the absence

of overlap, to one, meaning complete overlap (Linkie and Ridout 2011; Meredith and Ridout 2014). As our sample was large (n > 75), we estimated the overlap coefficient using the estimator Dhat4 (*e.g.*, Guerisoli et al. 2019) denoted  $\Delta_4$ . We computed the 95% confidence interval (hereafter, 95% CI) from 999 bootstrap samples to obtain this estimator's precision (Dias et al. 2019; Mori et al. 2020). The species' overlap coefficient was considered low if  $\Delta_4 < 0.50$ , intermediate if  $0.50 < \Delta_4$ < 0.70, and high if  $\Delta_4 > 0.70$  (*e.g.*, Monterroso et al. 2014).

### The spatiotemporal value (STV) for species' interactions

We wanted to use an approach that explore species' spatiotemporal overlap, which was depicted with both non-native species and native carnivores within the study area. At this end, we made a combination of results from spatial patterns; multiseason two-species co-occurrence analysis and the temporal activity overlap between these two kinds of species. The first parameter used was the SIF. We combined it to the temporal overlap coefficient ( $\Delta_4$ ) performed from the kernel density estimator. Both SIF and  $\Delta_4$ are probabilities, thus the probability of SIF "and"  $\Delta_4$ implies multiplication. Therefore, we computed the STV by multiplying the SIF value by the  $\Delta_4$  value *i.e.*, STV = SIF \*  $\Delta_4$ , as previously used by Farris et al. (2020). These two parameters (SIF and  $\Delta_4$ ) varied between the different paring species (non-native species and native carnivores). This combination was useful as sites that have a temporal activity overlap among species are not clearly defined. Also the spatial co-occurrence analyses of species can display bias, particularly when considering that investigated sites may be occupied by one or both species (for the paring species used), but these two species can be undetected. Such cases can imply the imperfection detection concept of MacKenzie et al. (2006). Accordingly, the STV shows a degree of overlap for each species pairing, thus 0 designates no overlap for both spatial and temporal patterns and as the STV rises this confirms the aggregation of spatial and temporal overlap, which depicts direct interactions between species (Farris et al. 2020).

#### RESULTS

## Overall Trend in Capture and Multi-Year Changes in Species Estimates

For non-native species, investigations in the TPNR produced over the three seasons a total of 526, 631, and 145 independent photographs of human, livestock, and dog, respectively. For carnivore species, there were 128, 154, and 412 independent photos of leopard, leopard cat, and fox, respectively. In total, the survey lasted 383 days, used 205 cameras, and yielded 26,216 trap nights. The overall trap success rates (which is the total number

**Table 1.** Parameters computed in the multiseason co-occurrence models to verify the effect of the invasive species' presence on the occupancy, detection, colonization, and local extirpation of carnivores in the Tieqiaoshan Provincial Natural Reserve (Par: Parameters)

Par	Definitions
$\psi^{\rm BA}$	The probability that species B initially occupies the area, given that species A is also present
$\psi^{\mathrm{Ba}}$	The probability that species B initially occupies the area, given that species A is not present
$\gamma^{\rm BAA}$	The probability that the area is colonised by species B in the interval t, $t_{+1}$ , given that species A is present in survey t and species A persists in the interval t, $t_{+1}$
$\gamma^{BAa}$	The probability that the area is colonised by species B in the interval t, $t_{+1}$ , given that species A is present in survey t and species A goes extinct in the interval t, $t_{+1}$
$\gamma^{\text{Baa}}$	The probability the area is colonised by species B in the interval t, $t_{+1}$ , given that species A is not present in survey t and species A does not colonise in the interval t, $t_{+1}$
$\varepsilon^{\scriptscriptstyle \mathrm{BAA}}$	The probability that species B becomes extinct in the area in the interval t, $t_{+1}$ , given that species A is present in survey t and species A persists in the interval t, $t_{+1}$
$\varepsilon^{^{\mathrm{BAa}}}$	The probability that species B becomes extinct in the area in the interval t, $t_{+1}$ , given that species A is present in survey t and species A goes extinct in the interval t, $t_{+1}$
$arepsilon^{\mathrm{Baa}}$	The probability that species B becomes extinct in the area in the interval t, $t_{+1}$ , given that species A is not present in survey t and species A does not colonise in the interval t- $t_{+1}$
$r^{\rm BA}$	The probability of detecting species B, given that both are present and species A detected
$r^{\mathrm{Ba}}$	The probability of detecting species B, given that both are present and species A not detected
Ø	Species Interactions Factor (SIF)

of independent captured photographs for a species divide by the total number of night-traps) for non-native species were 2.01%, 2.40% and 0.55% for human, livestock, and dog, respectively; and 0.48%, 0.58%, and 1.57% for leopard, leopard cat, and fox, respectively.

#### **Spatial Multi-Year Patterns**

For the multi-year occupancy estimates for nonnative species, livestock had the highest occupancy of  $0.78 \pm 0.06$  in SP<sub>2</sub> followed by human with  $0.61 \pm 0.03$ in the SP<sub>1</sub>, which also had the lowest site occupation among all non-native species ( $0.43 \pm 0.09$ ) in SP<sub>2</sub>. Dog had  $0.53 \pm 0.10$  as the highest site occupation estimates in SP<sub>3</sub> (Fig. 2). The leopard cat showed the highest probability of site occupation for carnivores, particularly SP<sub>3</sub>, with  $0.82 \pm 0.11$ . The fox had the second-highest site occupation probability of  $0.74 \pm 0.08$  in SP<sub>1</sub>. In the previous study (Vitekere et al. 2020b), the leopard depicted an average site occupation probability with its highest estimate found in SP<sub>1</sub> ( $0.54 \pm 0.09$ ). The lowest value found was for the leopard ( $0.44 \pm 0.10$ ) in SP<sub>2</sub> (Fig. 2). Neither non-native nor native carnivore species showed a substantial rate of change in occupancy (all values < 2, Table 2). However, carnivores had lower occupancy equilibrium values than non-native species in IS<sub>1</sub> (Table 2), except human, which represented the lowest value (0.22) (Table 2) when the pattern in IS<sub>2</sub> was different. The dog was the only species found at equilibrium in both IS according to its equilibrium occupancy values. The fox also showed an equilibrium in the SP<sub>2</sub> (Table 2).

In general, the fox had the highest detection probability, followed by human. The lowest detection pertained to the dog (Fig. 3). The human presence in the SP<sub>1</sub> (0.47  $\pm$  0.03) was the highest detection probability among non-native species, followed by livestock (0.32  $\pm$  0.03) in the SP<sub>1</sub>. The lowest detection was for the dog (0.20  $\pm$  0.03) in the SP<sub>2</sub>. For the carnivore, the fox had the highest detection probability (0.64  $\pm$  0.03) followed by the leopard with 0.36  $\pm$  0.05 in the SP<sub>1</sub>. The lowest detection was for the set detection was for the leopard cat (0.24  $\pm$  0.03) in the SP<sub>2</sub> (Vitekere et al. 2020b).

The highest colonisation rates were  $0.54 \pm 0.15$ ,



Fig. 2. Estimated site occupation probability across three years (2017–2019) of non-native species (human, livestock and dog) and carnivore species (leopard, cat and fox) in the Tieqiaoshan Provincial Natural Reserve; outputs of multi-year analysis computed in PRESENCE software. Carnivores' data have been previously used by Vitekere et al. (2020b).

**Table 2.** The occupancy equilibrium and rate of change in occupancy of non-native species (human, livestock, and dog) and carnivore species (leopard, cat, and fox), computed as derived parameters from equation one  $(eq_1)$  and equation two  $(eq_2)$ , respectively, to show trends in species occupancy from 2017 to 2019 in the Tieqiaoshan Provincial Natural Reserve

Species	$OE_1$	$\mathrm{RC}_1$	OE <sub>2</sub>	$RC_2$
human	0.22	0.48	0.52	1.68
livestock	0.58	1.82	0.45	0.72
dog	0.70	1.08	0.79	1.17
leopard	0.55	0.96	0.45	1.08
leopard cat	0.36	0.88	0.56	1.28
fox	0.49	0.5	0.69	1.41

OE: occupancy equilibrium and RC: rate of change in occupancy, one and two are IS1 and IS2.

 $0.42 \pm 0.12$ , respectively, for fox and dog in IS<sub>2</sub> and  $0.32 \pm 0.06$  for livestock in the IS<sub>1</sub> (Fig. 4). The lowest colonisation rates were in the IS<sub>1</sub> for both native carnivores and non-native species ( $0.15 \pm 0.09$  for leopard cat and  $0.13 \pm 0.08$  for human). Two non-native species had the highest extirpation rates (Fig. 5) for both categories of species, human in IS<sub>1</sub> and livestock in the IS<sub>2</sub>, with  $0.45 \pm 0.09$  and  $0.30 \pm 0.05$ , respectively. The lowest extirpation rate was for the dog in the IS<sub>2</sub> ( $0.11 \pm 0.09$ ). The fox had the highest extirpation rate among carnivore species, followed by the leopard cat both in the IS<sub>1</sub> ( $0.29 \pm 0.10$  and  $0.26 \pm 0.15$ , respectively). The lowest extirpated carnivore was leopard in the IS<sub>1</sub> ( $0.21 \pm 0.15$ ).

#### Multi-Year Co-occurrence Estimates

For all comparisons, there were no significant differences between the occupancy of a single species'

analysis and the two species co-occurrence analysis as all p-values were greater than 0.05. In the humancarnivores coexistence (Table 3A), the leopard occupancy remained almost the same with human absence except in the  $SP_2$ . The leopard cat's site occupation was low in human presence except in SP<sub>3</sub>. The fox depicted unchanged estimates for occupancy in human presence except in SP<sub>2</sub>. Human absence positively and negatively influenced colonisation and local extirpation of the leopard and the leopard cat, respectively (except the local extirpation in the  $IS_2$ ). For the fox, the colonisation slightly diminished in both presence and absence of human, and the extirpation rate remains unchanged in the human absence in both IS. Its occupancy did not change in human presence and absence for the first two SPs. The human presence influenced the detectability of the leopard in all three SPs. Simultaneously, the detection of the leopard cat did not markedly change in human presence, and human



Fig. 3. Estimated detection probabilities across three years (2017–2019) of non-native species (human, livestock and dog) and carnivore species (leopard, cat and fox) in the Tieqiaoshan Provincial Natural Reserve; outputs of multi-year analysis performed in PRESENCE software. Carnivores' data have been previously used by Vitekere et al. (2020b).



Fig. 4. Estimated colonisation rates across three years (2017–2019) of non-native species (human, livestock and dog) and carnivore species (leopard, cat and fox) in the Tieqiaoshan Provincial Natural Reserve; outputs of multi-year analysis performed in PRESENCE software, col1: colonisation in interseason 1 and col2: colonisation in interseason 2. Carnivores' data have been previously used by Vitekere et al. (2020b).

presence did not influence the fox's detectability.

Referring to the livestock-carnivore interaction (Table 3B), livestock presence brought a lesser influence to the occupancy of leopard and the occupancy of the leopard cat was modified (except the SP<sub>1</sub>) and the fox occupancy changed in the SP<sub>3</sub>. The colonisation rates of carnivores were influenced by livestock presence; as for the leopard and the fox, their occupancy slightly changed, especially in IS<sub>2</sub>. Surprisingly, livestock absence positively influenced the leopard's local extirpation when for leopard cat and fox, they were negatively influenced. Detection probability of leopard was lesser with livestock absence, for leopard cat and fox did not markedly change with livestock presence.

For dog-carnivore coexistence (Table 3C) the leopard and leopard cat occupancies slightly changed with dog presence when it markedly changed for the fox, especially in the SP<sub>3</sub>. In general, the dog presence influenced the colonization rates of the leopard and fox in both IS. The dog absence impacted negatively the leopard's local extirpation, while its presence changed the local extirpation for leopard cat and fox. The detection was slightly less for all species with the dog presence.

#### **Temporal Patterns**

All the non-native species portrayed a marked preference for daytime, with very low nocturnalities (proportion of observations between 18:00 and 06:00) of 0.11, 0.12 and 0.16 for human, livestock, and dog, respectively. The leopard was crepuscular with a broad peak of activity in the morning and a more sharply defined evening peak around 18:00–19:00, with a nocturnality of 0.41; the leopard cat and the fox preferred night-time with nocturnalities of 0.71 and 0.69, respectively, and peak activity around midnight. Non-native species had noteworthy peaks of activity in the morning (7:00) and in the early evening (18:00), whereas they were observed less frequently around noon (Fig. 6).

The activity time overlap coefficients ( $\Delta_4$ ) were low for all pairwise species combinations of non-native species with the leopard cat and the fox (the highest value was 0.48 between livestock and fox). However, overlap of non-native species with the leopard was higher (human-leopard: 0.73; livestock-leopard: 0.75 and dog-leopard: 0.74, Table 4). The spatiotemporal value (STV) (Table 4) concomitantly indicates the spatial and the temporal overlapping in one value, with the highest value for the species pairing livestockleopard (0.97) and the lowest for the pair humanleopard cat (0.41).

#### DISCUSSION

### Overall Species' Site and Time Occupation Patterns

Among all non-native species, livestock depicted the highest occupancy across all SPs, followed by human in SP<sub>1</sub> and SP<sub>3</sub> and dog. The non-native species were predominantly diurnal with very low nocturnality (close to 10%). Except for the leopard, which intensely selected crepuscular time, carnivore species preferred the night for their diel activity time. As is well known, humans always prefer daytime for their activities (Mori et al. 2020), inducing some invasive species (dog and livestock mainly) to have a similar pattern of time



Fig. 5. Estimated extirpation rates across three years (2017–2019) of non-native species (human, livestock, and dog) and carnivore species (leopard, cat, and fox) in the Tieqiaoshan Provincial Natural Reserve; outputs of multi-year analysis performed in PRESENCE, ext1: local extirpation in interseason 1 and ext2: local extirpation interseason 2. Carnivores' data have been previously used by Vitekere et al. (2020b).

activity as they are mostly human-dependent (Vanak and Gompper 2010; Farris et al. 2015 2016 2017). The presence of the non-native species was quantified by their occupancy estimates (> 0.40 for all species), which indicated how these species are permanent in PAs, although human activities are forbidden in most declared conservation ecosystems (Chape et al. 2005; Dudley 2008). Non-native species remain real threats

 Table 3. Co-occurrence results for non-native species (species A) and carnivores (species B)—including the species interaction factor (SIF)—within the Tieqiaoshan Provincial Nature Reserve (2017–2019)

A: human-carnivores

$ \begin{array}{c c c c c c c c c c c c c c c c c c c $		$\psi^{\rm BA}$	$\psi^{Ba}$	$\gamma^{\rm BAA}$	$\gamma^{\rm BAa}$	$\gamma^{\rm Baa}$	$\epsilon^{\text{BAA}}$	$\epsilon^{BAa}$	$\epsilon^{Baa}$	$\mathbf{r}^{\mathrm{BA}}$	$r^{Ba}$	φ
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	SP											
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	human-leopard											
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	one	$0.46\pm0.09$	$0.58\pm0.05$	$0.23\pm0.07$	$0.26\pm0.09$	$0.30\pm0.10$	$0.29\pm0.08$	$0.27\pm0.13$	$0.15\pm0.05$	$0.19\pm0.08$	$0.30\pm0.02$	$1.16\pm0.03$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	two	$0.42 \pm 0.15$	$0.39\pm0.01$	$0.15\pm0.03$	$0.18\pm0.06$	$0.22\pm0.08$	$0.32\pm0.10$	$0.21\pm0.08$	$0.18\pm0.14$	$0.20\pm0.10$	$0.14\pm0.12$	$1.18\pm0.09$
	three	$0.44 \pm 0.11$	$0.50\pm0.07$	NA	NA	NA	NA	NA	NA	$0.18\pm0.09$	$0.25\pm0.03$	$1.21\pm0.10$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	human-leopard cat											
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	one	$0.78 \pm 0.07$	$0.81 \pm 0.09$	$0.11 \pm 0.04$	$0.20 \pm 0.06$	$0.19 \pm 0.06$	$0.31\pm0.16$	$0.25 \pm 0.08$	$0.23 \pm 0.05$	$0.29 \pm 0.11$	$0.36\pm0.05$	$1.05 \pm 0.11$
	two	$0.63 \pm 0.13$	$0.71 \pm 0.04$	$0.14 \pm 0.08$	$0.18 \pm 0.12$	$0.22 \pm 0.07$	$0.22 \pm 0.11$	$0.19 \pm 0.09$	$0.20 \pm 0.04$	$0.33 \pm 0.08$	$0.49 \pm 0.14$	$1.08 \pm 0.15$
$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c}$	three	$0.78 \pm 0.11$	$0.67 \pm 0.08$	NA	NA	NA	NA	NA	NA	$0.31 \pm 0.10$	$0.30 \pm 0.12$	$1.11 \pm 0.05$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	human-fox											
$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c}$	one	$0.70 \pm 0.01$	$0.76 \pm 0.08$	$0.21 \pm 0.10$	$0.19 \pm 0.11$	$0.26 \pm 0.02$	$0.36 \pm 0.06$	$0.31 \pm 0.02$	$0.28 \pm 0.13$	$0.50 \pm 0.14$	$0.55 \pm 0.10$	$1.05 \pm 0.09$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	two	$0.42 \pm 0.09$	$0.58 \pm 0.11$	0.21 = 0.10 $0.28 \pm 0.07$	0.19 = 0.11 $0.36 \pm 0.05$	0.20 = 0.02 $0.48 \pm 0.03$	$0.32 \pm 0.01$	$0.32 \pm 0.04$	0.20 = 0.13 $0.23 \pm 0.02$	$0.35 \pm 0.13$	0.00 = 0.10 $0.48 \pm 0.05$	1.02 = 0.05 $1.11 \pm 0.05$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	three	$0.64 \pm 0.09$	$0.66 \pm 0.05$	NA	NA	NA	NA	NA	NA	$0.26 \pm 0.11$	$0.28 \pm 0.09$	$1.17 \pm 0.02$
$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	Di livastaali aami	Vorac										
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $		voies										
$ \begin{array}{c} \text{SP} \\ \text{Iivestock-leopard} \\ \text{one} & 0.46 \pm 0.09 & 0.49 \pm 0.05 & 0.44 \pm 0.02 & 0.24 \pm 0.07 & 0.20 \pm 0.13 & 0.16 \pm 0.08 & 0.29 \pm 0.02 & 0.28 \pm 0.05 & 0.24 \pm 0.15 & 0.25 \pm 0.17 & 1.29 \pm 0.02 \\ \text{two} & 0.48 \pm 0.10 & 0.41 \pm 0.09 & 0.39 \pm 0.05 & 0.32 \pm 0.12 & 0.18 \pm 0.09 & 0.21 \pm 0.05 & 0.24 \pm 0.03 & 0.20 \pm 0.10 & 0.25 \pm 0.08 & 1.34 \pm 0.09 \\ \text{three} & 0.55 \pm 0.08 & 0.45 \pm 0.07 & \text{NA} & \text{OA} & 0.47 \pm 0.08 & 0.19 \pm 0.13 & 1.27 \pm 0.11 \\ \text{ivestock-leopard} & & & & & & & & & & & & & & & & & & &$		$\psi^{\rm BA}$	$\psi^{\rm Ba}$	$\gamma^{\rm BAA}$	$\gamma^{BAa}$	$\gamma^{\mathrm{Baa}}$	$\epsilon^{\rm BAA}$	$\epsilon^{BAa}$	$\epsilon^{Baa}$	$r^{BA}$	$r^{Ba}$	$\varphi$
	SP											
$ \begin{array}{c} \mbox{one} & 0.46 \pm 0.09 & 0.49 \pm 0.05 & 0.44 \pm 0.02 & 0.24 \pm 0.07 & 0.20 \pm 0.13 & 0.16 \pm 0.08 & 0.29 \pm 0.02 & 0.28 \pm 0.05 & 0.24 \pm 0.15 & 0.25 \pm 0.17 & 1.29 \pm 0.02 \\ \mbox{two} & 0.48 \pm 0.10 & 0.14 \pm 0.09 & 0.39 \pm 0.05 & 0.32 \pm 0.12 & 0.18 \pm 0.09 & 0.21 \pm 0.05 & 0.24 \pm 0.03 & 0.00 & 0.02 \pm 0.08 & 0.20 \pm 0.01 & 0.25 \pm 0.08 & 1.24 \pm 0.09 \\ \mbox{three} & 0.56 \pm 0.08 & 0.45 \pm 0.07 & NA & N$	livestock-leopard											
$ \begin{array}{c} \mathrm{two} & 0.48 \pm 0.10 & 0.41 \pm 0.09 & 0.39 \pm 0.05 & 0.32 \pm 0.12 & 0.18 \pm 0.09 & 0.21 \pm 0.05 & 0.24 \pm 0.03 & 0.30 \pm 0.08 & 0.20 \pm 0.10 & 0.25 \pm 0.08 & 1.34 \pm 0.09 \\ \mathrm{three} & 0.56 \pm 0.08 & 0.45 \pm 0.07 & \mathrm{NA} & NA$	one	$0.46\pm0.09$	$0.49\pm0.05$	$0.44\pm0.02$	$0.24\pm0.07$	$0.20\pm0.13$	$0.16\pm0.08$	$0.29\pm0.02$	$0.28\pm0.05$	$0.24\pm0.15$	$0.25\pm0.17$	$1.29\pm0.02$
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	two	$0.48\pm0.10$	$0.41\pm0.09$	$0.39\pm0.05$	$0.32\pm0.12$	$0.18\pm0.09$	$0.21\pm0.05$	$0.24\pm0.03$	$0.30\pm0.08$	$0.20\pm0.10$	$0.25\pm0.08$	$1.34\pm0.09$
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	three	$0.56\pm0.08$	$0.45\pm0.07$	NA	NA	NA	NA	NA	NA	$0.47\pm0.08$	$0.19\pm0.13$	$1.27\pm0.11$
cat one 0.81 ± 0.07 0.78 ± 0.08 0.14 ± 0.08 0.28 ± 0.13 0.28 ± 0.10 0.14 ± 0.05 0.13 ± 0.08 0.10 ± 0.03 0.28 ± 0.11 0.36 ± 0.12 1.32 ± 0.07 two 0.67 ± 0.02 0.52 ± 0.08 0.17 ± 0.04 0.19 ± 0.11 0.23 ± 0.09 0.28 ± 0.07 0.23 ± 0.09 0.15 ± 0.08 0.19 ± 0.08 0.31 ± 0.10 1.08 ± 0.03 three 0.57 ± 0.10 0.46 ± 0.06 NA NA NA NA NA NA NA NA NA 0.33 ± 0.10 0.32 ± 0.09 1.06 ± 0.08 livestock-fox one 0.76 ± 0.08 0.70 ± 0.03 0.27 ± 0.08 0.25 ± 0.05 0.30 ± 0.08 0.32 ± 0.09 0.29 ± 0.11 0.23 ± 0.07 0.66 ± 0.05 0.60 ± 0.07 1.04 ± 0.04 two 0.60 ± 0.01 0.47 ± 0.05 0.48 ± 0.03 0.50 ± 0.03 0.55 ± 0.11 0.27 ± 0.13 0.25 ± 0.08 0.18 ± 0.10 0.50 ± 0.01 0.57 ± 0.11 1.14 ± 0.09 three 0.75 ± 0.10 0.49 ± 0.09 NA NA NA NA NA NA NA NA 0.32 ± 0.04 0.39 ± 0.06 1.10 ± 0.09 C: dog-carnivores C: dog-carnivores SP dog-leopard one 0.43 ± 0.04 0.48 ± 0.01 0.36 ± 0.08 0.39 ± 0.07 0.33 ± 0.01 0.25 ± 0.10 0.23 ± 0.05 0.18 ± 0.02 0.29 ± 0.07 0.33 ± 0.02 0.93 ± 0.02 three 0.55 ± 0.08 0.31 ± 0.03 NA NA NA NA NA NA NA NA 0.23 ± 0.06 1.19 ± 0.04 0.22 ± 0.08 1.18 ± 0.02 0.29 ± 0.07 0.33 ± 0.02 0.93 ± 0.02 three 0.55 ± 0.08 0.31 ± 0.03 NA NA NA NA NA NA NA NA NA 0.23 ± 0.06 0.19 ± 0.04 0.22 ± 0.08 1.36 ± 0.03 dog-leopard one 0.43 ± 0.04 0.48 ± 0.01 0.36 ± 0.08 0.39 ± 0.07 0.33 ± 0.01 0.25 ± 0.10 0.23 ± 0.05 0.18 ± 0.02 0.29 ± 0.07 0.33 ± 0.02 0.93 ± 0.02 two 0.61 ± 0.01 0.32 ± 0.03 NA NA NA NA NA NA NA NA 0.23 ± 0.08 0.37 ± 0.10 1.29 ± 0.07 dog-leopard car one 0.83 ± 0.05 0.78 ± 0.01 0.16 ± 0.06 0.13 ± 0.10 0.20 ± 0.09 0.19 ± 0.07 0.23 ± 0.01 0.15 ± 0.08 0.24 ± 0.07 0.31 ± 0.08 1.17 ± 0.11 two 0.78 ± 0.08 0.60 ± 0.09 0.19 ± 0.04 0.28 ± 0.08 0.29 ± 0.02 0.18 ± 0.05 0.21 ± 0.01 0.21 ± 0.06 0.28 ± 0.03 1.16 ± 0.05 three 0.81 ± 0.03 0.77 ± 0.07 NA 0.25 ± 0.12 0.32 ± 0.09 1.31 ± 0.04 dog-fox one 0.83 ± 0.10 0.66 ± 0.09 0.42 ± 0.08 0.38 ± 0.08 0.29 ± 0.06 0.32 ± 0.01 0.15 ± 0.06 0.58 ± 0.10 0.50 ± 0.08 1.32 ± 0.03 two 0.60 ± 0.08 0.35 ± 0.11 0.44 ± 0.03 0.56 ± 0.07 0.32 ± 0.01 0.29 ± 0.05 0.21 ± 0.01 0.51 ± 0.07 0.58 ±	livestock-leopard											
$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c}$	cat											
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	one	$0.81 \pm 0.07$	$0.78 \pm 0.08$	$0.14 \pm 0.08$	$0.28 \pm 0.13$	$0.28 \pm 0.10$	$0.14 \pm 0.05$	$0.13 \pm 0.08$	$0.10 \pm 0.03$	$0.28 \pm 0.11$	$0.36 \pm 0.12$	$1.32 \pm 0.07$
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	two	$0.67 \pm 0.02$	$0.52 \pm 0.08$	$0.17 \pm 0.04$	$0.19 \pm 0.11$	$0.23 \pm 0.09$	$0.28 \pm 0.07$	$0.23 \pm 0.09$	$0.15 \pm 0.08$	$0.19 \pm 0.08$	0.30 = 0.12 0.31 + 0.10	1.02 = 0.07 $1.08 \pm 0.03$
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	three	$0.67 \pm 0.02$ $0.57 \pm 0.10$	$0.52 \pm 0.06$ $0.46 \pm 0.06$	NA	0.17 ± 0.11	NA	0.20 ± 0.07	NA	NA	$0.19 \pm 0.00$ $0.33 \pm 0.10$	$0.31 \pm 0.10$ $0.32 \pm 0.09$	$1.06 \pm 0.03$ $1.06 \pm 0.08$
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	livestock-fox	0107 - 0110	0110 - 0100							0100 - 0110	0102 - 0107	1100 - 0100
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	one	$0.76 \pm 0.08$	$0.70 \pm 0.03$	$0.27 \pm 0.08$	$0.25 \pm 0.05$	$0.30 \pm 0.08$	$0.32 \pm 0.09$	$0.29 \pm 0.11$	$0.23 \pm 0.07$	$0.66 \pm 0.05$	$0.60 \pm 0.07$	$1.04 \pm 0.04$
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	two	$0.70 \pm 0.00$ $0.60 \pm 0.01$	$0.70 \pm 0.05$ $0.47 \pm 0.05$	$0.27 \pm 0.00$ $0.48 \pm 0.03$	$0.20 \pm 0.03$ $0.50 \pm 0.03$	$0.50 \pm 0.00$ $0.55 \pm 0.11$	$0.32 \pm 0.03$ $0.27 \pm 0.13$	$0.25 \pm 0.08$	$0.23 \pm 0.07$ $0.18 \pm 0.10$	$0.00 \pm 0.00$	$0.00 \pm 0.07$	$1.01 \pm 0.01$ $1.14 \pm 0.09$
C: dog-carnivores $\psi^{BA}$ $\psi^{Ba}$ $\gamma^{BAA}$ $\gamma^{BAa}$ $\gamma^{Baa}$ $\varepsilon^{BAA}$ $\varepsilon^{BAa}$ $\varepsilon^{Baa}$ $r^{BA}$ $r^{Ba}$ $\varphi$ SP         dog-leopard         one         0.43 \pm 0.04         0.48 \pm 0.01         0.36 \pm 0.08         0.39 \pm 0.07         0.33 \pm 0.01         0.25 \pm 0.10         0.23 \pm 0.05         0.18 \pm 0.02         0.29 \pm 0.07         0.33 \pm 0.02         0.93 \pm 0.02           two         0.61 \pm 0.01         0.32 \pm 0.05         0.20 \pm 0.03         0.25 \pm 0.09         0.23 \pm 0.04         0.28 \pm 0.08         0.27 \pm 0.08         0.16 \pm 0.06         0.19 \pm 0.07         0.33 \pm 0.02         0.93 \pm 0.02           two         0.61 \pm 0.01         0.32 \pm 0.03         NA         NA         NA         NA         NA         0.23 \pm 0.07         0.23 \pm 0.08         0.27 \pm 0.08         0.16 \pm 0.06         0.13 \pm 0.03         0.12 \pm 0.07         0.23 \pm 0.01         0.16 \pm 0.06         0.13 \pm 0.10         0.20 \pm 0.09         0.19 \pm 0.07         0.23 \pm 0.11         0.25 \pm 0.08         0.24 \pm 0.07         0.31 \pm 0.08         1.17 \pm 0.11           two         0.78 \pm 0.08         0.60 \pm 0.09         0.19 \pm 0.04         0.28 \pm 0.03         0.17 \pm 0.05         0.20 \pm 0.02         0.18 \pm 0.05         <	three	$0.00 \pm 0.01$ $0.75 \pm 0.10$	$0.47 \pm 0.09$ $0.49 \pm 0.09$	0.40 ± 0.05	0.50 ± 0.05 NA	0.55 ± 0.11 NA	0.27 ± 0.15	0.23 ± 0.00	0.10 ± 0.10 NA	$0.30 \pm 0.01$ $0.32 \pm 0.04$	$0.37 \pm 0.011$ $0.39 \pm 0.06$	$1.14 \pm 0.09$ $1.10 \pm 0.09$
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	C: dog-carnivores											
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$				DAA	DA-		BAA					
SP         dog-leopard         0.43 ± 0.04         0.48 ± 0.01         0.36 ± 0.08         0.39 ± 0.07         0.33 ± 0.01         0.25 ± 0.10         0.23 ± 0.05         0.18 ± 0.02         0.29 ± 0.07         0.33 ± 0.02         0.93 ± 0.02           two         0.61 ± 0.01         0.32 ± 0.05         0.20 ± 0.03         0.25 ± 0.09         0.23 ± 0.04         0.28 ± 0.08         0.27 ± 0.08         0.16 ± 0.06         0.19 ± 0.04         0.22 ± 0.08         1.36 ± 0.03           three         0.55 ± 0.08         0.31 ± 0.03         NA         NA         NA         NA         0.40         0.42         0.83 ± 0.05         0.19 ± 0.04         0.22 ± 0.08         1.36 ± 0.03           dog-leopard cat		$\Psi^{BA}$	$\psi^{\rm Ba}$	$\gamma^{BAA}$	$\gamma^{BAa}$	$\gamma^{\text{Baa}}$	ε	ε <sup>BAa</sup>	ε <sup>Baa</sup>	r <sup>BA</sup>	r <sup>Ba</sup>	φ
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Estimates are accompanied by standard errors, SP: sampling period and NA: not applicable because colonization and local extirpation rates are only present in IS.

to wildlife (Gerber et al. 2012a; Sleeman 2013; Vanak et al. 2013; Brodie et al. 2015; Chanchani et al. 2016). Concomitantly, native carnivore site occupation was evident (lowest occupancy was 0.44) in the area, with the leopard cat having the highest occupancy across all SP (Fig. 2). The fox showed higher occupancy than the leopard and was captured mainly at night. Commonly, carnivores are reputed to be nocturnal (Schuette et al. 2013; Monterroso et al. 2014; Hua et al. 2020). Nevertheless, despite this nighttime preference, we put them into two categories based on the behaviors we observed. The leopard would be qualified as a "cathemeral species" as it is active during the day and night and mesocarnivores qualified as "nocturnal preferred."

Contrary to the first hypothesis, these carnivores did not reveal significant changes in site occupancy across years, a fact evidenced by the rate of change in occupancy as a noticeable rise was documented from IS<sub>1</sub> to IS<sub>2</sub>. This rate demonstrated stability in both IS (Table 2) for non-native species, yet for human, the occupancy markedly changes alternately. The recent Natural Forests Protection Program (NFPP) policies of the China national government for habitat restoration and improvement of management strategies within the TPNR would be responsible for constant carnivores



Fig. 6. Overlap of density of activity over three years between the non-native species (lines) and native carnivores species (dashed lines) based on camera trap data (2017–2019) in Tieqiaoshan Provincial Natural Reserve ("cat" represents the "leopard cat").

**Table 4.** Estimated activity overlap coefficient ( $\Delta_4$ ), associated 95% confidence interval obtained using the bootstrap method with 999 sample replications and the spatiotemporal value (STV which describes simultaneously the spatial and temporal overlap value, its obtained by multiplying the SIF by  $\Delta_4$ ) for non-native species and carnivores species the in the Tieqiaoshan Provincial Nature Reserve (2017–2019)

Species pairwise	$(\Delta_4)$	95% CI	Average STV
human-leopard	0.73	0. 64-0.80	0.86
human- cat	0.38	0.31-0.45	0.41
human- fox	0.42	0.37-0.47	0.46
livestock-leopard	0.75	0.37-0.83	0.97
livestock- cat	0.44	0.37-0.51	0.50
livestock- fox	0.48	0.44-0.53	0.52
dog-leopard	0.74	0.64-0.82	0.88
dog- cat	0.39	0.31-0.48	0.44
dog-red fox	0.44	0.37-0.50	0.64

estimates (Vitekere et al. 2020b; Zhu et al. 2021). These NFPP measures have enlarged the size and enhanced the forest quality of the PA (Xu et al. 2009; Wei et al. 2014). Furthermore, the homogeneousness effects due to the landscape features (case of the TPNR ecosystem) can also explain the native carnivores' site occupation, as found by Kass et al. (2020). The PA's outside intense disturbances would promote a type of lockdown for wildlife in the TPNR.

Regarding human occupancy in a PAs, findings opposite to ours were established by Farris et al. (2017), where this occupancy progressively increased over five years. Yet the same results revealed an alternate frequency of a lesser rising and diminution of dog occupancy over time. For our study, both non-native and native carnivore species were well detected (except the dog, but with normal detectability  $\geq 0.20$ ). The estimated detection values determine the occupancy veracity, particularly when sampling sites and incidences are small (Royle and Nichols 2003; MacKenzie et al. 2006). Therefore, when species are well detected within their habitat (detection probability  $\geq 0.30$  for repetitive sampling occasion  $\geq 5$ ), the occupancy estimates would not be considered biased (Nicholson and van Manen 2009). As these changes in species' site occupation were weaker than expected, it is perhaps worth exploring disturbances, biophysical and environmental variables effects. Nonetheless, although the gap between both non-native and native carnivore species' occupancy equilibria was insignificant (except dog), carnivores were characterised by lower occupancy equilibrium values than non-native species. Thus, it is evident that as long as native carnivores depict low occupancy equilibrium compared with non-native species within the TPNR, their long-term existence is substantially threatened since non-native species are potential sources of danger for PAs management worldwide.

#### **Spatiotemporal Overlap Patterns**

In the co-occupation framework, the pairwise human-carnivores portrayed a general trend of indirect space overlap with all carnivore species referring to the SIF estimates and contrary to our hypothesis. The leopard was the carnivore most influenced by human presence (all SIF > 1) as this species (leopard) was at equilibrium for night and daytime, inducing a high time overlap with non-native species (all  $\Delta_4 \ge 0.73$ ). Our findings revealed that, spatially, the leopard did not avoid zones with human presence as previously found with other top predators (*e.g.*, Flores-Morales et al. 2019 for the Coyote *Canis latrans*, Guerisoli et al. 2019 for Puma *Puma concolor*, and Mori et al. 2020 for Wolf *Canis lupus*). Yet, Zhu et al. (2021) found that anthropogenic disturbances affect leopard's density in this area. However, leopard's activities were generally nocturnal, and specifically crepuscular; logically in the TPNR, the leopard may avoid contact with non-native species, specifically human, concentrating its activity bouts late in the dusk hours to avoid harmful contacts. On the other hand, the leopard cat and fox depicted a higher SIF with human than the leopard did with human. Indeed, species differ in their aptitude to adapt to human presence on their degree of specialization in habitat use and ranging habits (Carricondo-Sánchez 2018). Some species can shift and create an adaptation to human presence due to their ecological behavior. Although there was a reduction in the detectability of the leopard at the sites where human was signaled ( $r^{BA}$  $< r^{\text{Ba}}$ ), the human presence had no substantial effect on leopard site use (Table 3A compared to Fig. 3). Leopard site occupation was even higher in the SP<sub>2</sub> in sites where leopards initially occupied the area; given that humans were also present ( $\psi^{BA}$ , Table 3A). Other studies have documented this (Carter et al. 2015; Farris et al. 2017) and corroborated our findings. In a PA with anthropogenic activities, human site occupation has always been followed by disturbance effects on the site, attracting small and average bodiedsize mammals. These zones in PAs would be in the vicinity of boundary areas where PA managers lack the authority, resources, and funding to establish wildlife management regulations (Lindenmayer et al. 2012; Bauer et al. 2015). Therefore, the fox is a species that may be attracted there. Thus, the presence of this carnivore would be perceived as colonisation rate since it is reflected as a conditional occupancy (MacKenzie et al. 2006). Colonisation also did not greatly fluctuate with human presence and human absence at the sites. In our investigation, the growth in human-carnivores spatiotemporal co-occurrence over these three years is a measure of human invasion related to a wide range of anthropogenic activities as resource extraction and livestock pasture.

The leopard showed direct interactions within the site co-occupation with livestock (all SIF > 1). At the same time, the leopard cat and the fox were characterised by independent interactions (SIF close to 1), with the exception of the leopard cat in SP<sub>1</sub>. Also, the temporal overlap between the leopard and livestock was apparent ( $\Delta_4 = 0.75$ ) and can be elucidated by previous findings in the TPNR since the leopard feeds regularly on livestock (Consolee et al. 2020; Vitekere et al. 2020a), I don't understand this clause. Such cooccurrence has been seen by Lovari et al. (2015) in an analysis of common leopard and livestock coexistence in an area of Nepal. Other similar studies stated that livestock had been revealed as the second important prey of the top predators (Okello et al. 2014; Ciucci et al. 2018). Livestock becomes a central component of the prey of leopards probably because there is a lack of defence by herds' keepers, particularly during the calving period inside the PAs. These findings are partially in accordance with the temporal hypothesis given that livestock and leopard use the same time and overlap leads to predation. Nevertheless, when farmers undertake their livestock defence they habitually use retaliatory procedures to harm predators (Treves and Karanth 2003; Cromsigt et al. 2013). The TPNR landscape is reputed to host more farmers, revealing that within this landscape livestock is more present and carnivore avoidance interactions are more unlikely to occur. So, direct interactions between leopard and livestock in the TPNR indirectly launches an interaction between humans and carnivores, which always ends in human-carnivore conflict (Consolee et al. 2020; Vitekere et al. 2020a). Occasionally fences are built and have proven palliative solutions to impede livestock encroachment from PAs (Lovegrove et al. 2002).

In an area characterised by limited resources due to its uniformity of habitats, it is common to observe native species being destabilised by non-native species. The heterogeneity of a landscape is somewhat responsible for ecological niche diversification (Eppstein et al. 2006; Soto and Palomares 2015), and trails play an important role in the invasion of PAs. This is the case in the TPNR, which is crossed by diverse trails. A strong and positive correlation between trails and dog presence in PAs was mostly documented when studying non-native species effects in an ecosystem (Farris et al. 2016 2017). The dog was characterized by higher spatiotemporal overlaps with leopards (all SIF > 1except in SP<sub>1</sub>, and  $\Delta_4 = 0.74$ ). A significant spatial overlap was depicted by the apparent co-occurrence between the dog and fox in all SP with the fox. While this invasive species and the fox are similar in body size, they are not ecologically similar, and the principle of competitive exclusion (MacArthur and Levins 1967) may not be applied. Many studies brought out the fox preference for human-disturbed habitat (Macdonald et al. 2004; Hughes and Macdonald 2013; Barrull et al. 2014; Mori et al. 2015; Vitekere et al. 2020b). The dog being at occupancy equilibrium in our study clearly supports inferences about its long-term presence in the TPNR. In a Madagascar study, Farris et al. (2016) found the dog excluding native carnivores in some areas within his study area. Free-ranging domestic dogs are amongst the most well-known worldwide non-native species that often affect native carnivores. This situation is a little challenging when there is a spatiotemporal overlap with native carnivores. As it is also a carnivore, the dog will interact with native carnivores by becoming

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a real competitor (Hughes and Macdonald 2013; Vanak et al. 2013; Mella-Méndez et al. 2019). It can reduce the prey biomass (Frank et al. 2014; Wierzbowska et al. 2016), transmit disease (Rasambainarivo et al. 2017; Mella-Méndez et al. 2019) and undertake direct intraguild aggression in PAs (Hughes and Macdonald 2013). Our study displays the dog's ability to influence the temporal patterns of native carnivores. Precisely, we add the evidence to the body of literature the competitive dynamics between the dog and the leopard within a human-dominated landscape.

However, while mesocarnivores were strongly bounded to the night-time, they exhibited some plasticity within this preferred temporal interval as their  $\Delta_4$  were low than 0.75. Indeed, none of these two species (leopard cat and fox) displayed a constant activity pattern in nocturnal times. Further researchers have found the wild cat and fox using night-time but with some evident activities shifted in the diurnal period (Barrull et al. 2014; Monterroso et al. 2014); consequently, Monterroso et al. (2014) qualified them as "facultative nocturnal". According to our findings, we assumed that, regardless of the high night-time rate of activities, these mesocarnivores tried to avoid the overlap activities with the big feline (leopard) to maximize their ability in the landscape and moderate the risks for antagonistic overlap. Yet, this does not necessarily mean they avoid leopard, because the nocturnality of some mediumsized carnivores is proved to coincide with their prey's activity. In contrast, with 31% of its diel activity in the daytime, the red fox did not totally avoid this period. Previous studies found the red fox performing important daytime activity (Cavallini and Lovari 1994; Adkins and Stott 1998). An equivalent observation was also made for the leopard cat, which performed 29% of its activities during this study in the daytime, corroborating other studies that found 20-21% (Germain et al. 2008; Monterroso et al. 2014). Generally, these two carnivores did not depict time overlap with non-native species, except at a small level, (20-30% only in the daytime), as all non-native species were characterized by a diel activity in daytime  $\Delta_4$  of pairwise species combination less than the 0.50. This would have some advantages in these two species' ecologies, particularly allowing flexibility in their diel activity patterns (as both mesocarnivores depicted spatial overlap with non-native species). It can permit the facilitation of accessing other affordable prey in their habitat or support, avoiding the riskiest periods of the day for both non-native species and top predators. However, Mori et al. (2020) found that the red fox is a mesocarnivore that often overlap spatially and temporally with top predators. Yet, these findings were not completely corroborated by the STV for species pairings as the fox implied the average

STV with non-native species (from 0.46 to 0.66). For the pairing species with highest STV (human-leopard, livestock-leopard, dog-leopard even dog-fox) this value reflects the state of spatiotemporal aggregation. Such scenario is the riskiest state for native species as it portrays potential threats for shared space and probably leads to more other possible vulnerabilities (negative interactions) to native species (Okello et al. 2014; Rasambainarivo et al. 2017; Mella-Méndez et al. 2019; Farris et al. 2020) since they occur in the same space at the same time.

#### CONCLUSIONS

Although the principal aim of PAs is to protect biodiversity, elusive species such as carnivores have always experienced interspecies threats that are hard to detect since carnivore studies often rely on passive sampling when investigating species' spatiotemporal threats (Farris et al. 2017 2020). In this study, the presence of invasive species was considered to be dependent on human presence. Only humans would be presumed to interact with wildlife directly, but livestock and dogs depicted clear overlaps with carnivores. Most livestock interactions with mesocarnivores were indirect, but the dog influenced the leopard and the fox's site occupations more.

In terms of the conservation implications, our results suggest that the presence of wild prey could reduce livestock depredation by the leopard and thus reduce the spatiotemporal overlap between carnivores and livestock. This would help to facilitate coexistence patterns with humans within the TNR and imply sustainable management of the reserve's landscape by reducing retaliatory measures. Such outcomes could be used to develop targeted education programs that inform local people living inside and outside the TNR or travelling with domestic dogs in the PA on the negative impacts their dogs may have on native carnivores. Considering spatiotemporal patterns using two niche dimensions can potentially lead to confusion in inferences interactions. For example, one might conclude that a native species has been impacted by non-native species when the two species do not demonstrate any temporal overlap because they use different time activities. Therefore, studies integrating landscape features (disturbances, habitat type, and diet or prey) are necessary to understand better these intrinsic interactions in species, which are essential drivers of population and community dynamics over time in Protected Areas.

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