Distribution and density of oxpeckers on giraffes in

² Hwange National Park, Zimbabwe

Roxanne Gagnon¹, Cheryl T. Mabika², and Christophe Bonenfant^{3,4}

¹Université de Sherbrooke, Département de biologie, 2500 Boulevard de l'Université, Sherbrooke, Québec J1K 2R1, Canada.

²Zimbabwe Parks and Wildlife Management Authority, Hwange Cluster, Main Camp, Zimbabwe.

 ³ ³Université Claude Bernard – Lyon 1; UMR CNRS 5558, Laboratoire de Biométrie et Biologie Évolutive, F-69 622, Villeurbanne CEDEX, France.
 ORCID: 0000-0002-9924-419X
 ⁴LTSER France, Zone Atelier "Hwange", Hwange National Park, Bag 62, Dete, Zimbabwe – CNRS HERD (Hwange Environmental Research Development) program

4 Correspondence

- 5 C. Bonenfant
- 6 Email: christophe.bonenfant@univ-lyon1.fr

Oxpeckers (Buphagus sp.) are two bird species closely associated to large Abstract mammals, including giraffes (Giraffa camelopardalis). Here, we counted every oxpeckers on giraffe's body from pictures to infer on the foraging behaviour of birds. We tested whether oxpeckers distributed themselves at random or aggregated on individual giraffes. We then quantified the distribution of oxpeckers over predefined zones on the giraffe body, to test the prediction that birds would preferentially be located on the host's body parts with the greatest amount of ticks. Oxpeckers displayed a strong aggregation behaviour with few hosts carrying many birds while many carried a limited number or no birds, a pattern that differed between sexes. Oxpeckers were disproportionately found on the mane and back, where the density of ticks is presumably the highest. This high aggregation level of birds is typical of parasitic species and therefore suggests that, in some cases, oxpeckers behave more as parasites rather than a mutualistic species. Alternatively this particular oxpecker distribution may mirror the distribution of ticks on giraffes. Abundance of ticks appears as a major driver of the oxpecker foraging behaviour, and the oxpecker-large herbivores system proves to be highly relevant for the study of host-parasite dynamics.

The distribution of animals in the environment results from a complex sequence of 2 behavioural decisions aiming at satisfying the energy requirements of individuals while 3 minimizing costs of movements, competition with con-specifics or other species, in balance with the perceived risks of predation for prey species (Krebs, 1972). From an evolutionary 5 perspective, animals should select habitats with the highest suitability (sensu Fretwell & 6 Lucas, 1969), *i.e.* those habitats in which the fitness of individuals will be the greatest. 7 Variation in habitat suitability, in space and time, is of prime importance to the ecology of 8 species with consequences on its distribution range (MacArthur, 1972), its mating tactic 9 (Emlen & Oring, 1977), or its population dynamics (Lack, 1966; Ostfeld & Keesing, 2000). 10 At the population level, when and where resources are available is a strong predictor of 11 the abundance of animals. Assuming an homogeneous distribution of discrete resources 12 patches in the landscape and a random walk of foraging animal trajectories, the expected 13 number of foragers per patch is given by a Poisson distribution (Hutchinson & Waser, 14 2007). This simple model of encounters between motile entities underpins most 15 multi-species interaction models, including the Lotka-Volterra (Lotka, 1956; Hutchinson & 16 Waser, 2007) or Nicholson-Bailey models (May, 1978) for, respectively, predators-prey and 17 hosts-parasitoid dynamics. Any deviation from the Poisson distribution is usually 18 interpreted as a sign of aggregation (over-dispersion) or avoidance (under-dispersion) of 19 individuals (Pielou (1969, p. 96) but see Taylor et al. (1979) or Sjöberg et al. (2000) for a 20 discussion and alternatives). Several indices have been proposed to quantify aggregation 21 levels among populations or species from count data (see Kretzschmar and Adler 1993 for 22 a review), the most widely used being the aggregation index k (Shaw et al., 1998). 23 Aggregation may arise from the animal behaviour such as social interactions (Wittenberger, 24 1981), constrain on mobility among patches (Gueron & Levin, 1995), or if animals perform 25 area-restricted search of food patches (Morales et al., 2010) or do copy what the other 26 con-specifics do when using public information (Clark & Mangel, 1986). 27 In multi-species interactions such as in bird vs. large mammals associations, the 28

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distribution of the birds is first guided by the one of the mammalian hosts, conceptually 1 equivalent to a resource patch. This scenario fits the association between oxpeckers 2 (Buphagus sp.) and large herbivores, described in the early 20th century (Moreau, 1933). 3 Oxpeckers are two bird species of sub-Saharan Africa associated to savanna ecosystems 4 (Hustler, 1987; Plantan, 2009; Palmer & Packer, 2018). They live and feed almost 5 exclusively on the body of large herbivores such as African buffaloes (Syncerus caffer 6 caffer), giraffes (Giraffa camelopardalis), black rhinoceros (Diceros bicornis), white 7 rhinoceros (Ceratotherium simum), impalas (Aepyceros melampus), greater kudus 8 (Tragelaphus strepsiceros), common elands (Taurotragus oryx) and sable antelopes 9 (Hippotragus niger) (Hustler, 1987; Stutterheim et al., 1988; Palmer & Packer, 2018). 10 Oxpeckers mainly prey upon ectoparasites of their large mammalian hosts although they 11 sometimes can snatch tissues from their host (Bezuidenhout & Stutterheim, 1980). In 12 terms of resource selection, the foraging behavior of oxpeckers can be decomposed into 13 two main steps. The first step for birds is to localize large mammals in the landscape which, 14 for oxpeckers represent motile and widely dispersed resources patches of varying size. The 15 second event takes place on the host's body where oxpeckers will search for the most 16 suitable body part in terms of ectoparasites. These two sequential steps, taking place at 17 different spatio-temporal scales (see Johnson, 1980), will drive the distribution of oxpeckers 18 among hosts and on their body. 19

At the largest spatial scale, the distribution of oxpeckers among the different species of 20 mammalian hosts has been documented for decades (Moreau, 1933; Grobler, 1980; 21 Hustler, 1987; Ndlovu & Combrink, 2015). Clear preferences were demonstrated and, with 22 a large body and a high tolerance to the presence of birds, the giraffe appears to be one of 23 the key host for two oxpecker species (Grobler, 1980; Veríssimo et al., 2017). Much less is 24 known about how oxpeckers are distributed on the different individuals of a given host 25 species, with the exception of maximum records (e.g. 51 birds on one side of a single 26 giraffe, Veríssimo et al. (2017)). At the host level of resource selection, oxpeckers seems to 27 favor some body parts of their hosts but what body parts is most attractive seems to be 28 different according to the host species (Palmer & Packer, 2018). For instance, Ndlovu & 29

1 Combrink (2015) reported that red-billed oxpeckers (Buphagus erythrorhynchus) were most

² frequently perch on the back and the head of buffaloes and white rhinoceros and that the

³ neck was preferred on giraffes.

In this paper we investigated the among-host and within-host distribution of oxpeckers
 on giraffes at Hwange National Park, Zimbabwe. We extracted the number and location of
 yellow-billed (*Buphagus africanus*) and red-billed oxpeckers on *n* = 683 giraffes from 500
 pictures collected since 2013. We aimed at testing the random encounter model between
 foragers and hosts, and the ideal free distribution (IFD, Fretwell & Lucas, 1969) with
 oxpeckers as foragers and giraffes as their primary large mammalian hosts. We tested the
 three following predictions:

1. Distribution of oxpeckers among giraffes: By comparing the observed distribution of 11 the number of oxpeckers per giraffe with the expected prediction from theoretical 12 models, one may infer the underlying behaviour of resource selection by birds and 13 their movements. If oxpeckers search for giraffes at random in the landscape the 14 random encounter model predicts a Poisson distribution of birds among hosts 15 (Hutchinson & Waser, 2007). Alternatively, if oxpeckers aggregate preferentially on 16 some particular giraffes because of marked differences in the parasite load or 17 because of copying behaviour, the model predicts a negative binomial distribution of 18 birds among hosts (Pielou, 1969); 19

2. Distribution of oxpeckers on giraffe body: According to the IFD (Fretwell & Lucas, 20 1969), oxpeckers should be distributed on the giraffe body parts proportionally to the 21 local ectoparasite load. Therefore, if the ectoparasite density is homogeneous over 22 the whole giraffe body, the IFD predicts a homogeneous number of oxpeckers per 23 area unit. Alternatively, if ectoparasites concentrate on some specific body parts, the 24 IFD predicts a heterogeneous distribution of birds, with higher densities of oxpeckers 25 on giraffe body parts with the higher ectoparasite burden (Horak et al., 1983; 26 Mysterud et al., 2014). From previous observations (Plantan, 2009; Ndlovu & 27 Combrink, 2015), giraffe body parts with the highest oxpecker number should be, in 28

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decreasing order, the mane, the neck, the scapula and the back;

3. Sex-differences in bird load of giraffes: Many studies evidenced that the ectoparasite
 load is proportional to the body mass and skin surface of the host (Horak et al., 1983, 1987; Koenig, 1997). Consequently a bigger host should carry more ectoparasites
 and hence, more birds than a smaller one. Sexual size dimorphism is observed
 among many species and is particularly observable between male and female
 giraffes, reaching a 43% difference for fully grown individuals (Shorrocks, 2016). We
 will therefore test the prediction that more birds are present on male than on female
 giraffes;

10 2 | MATERIAL AND METHODS

11 2.1 | Study site

This study was undertaken around Main Camp in the northeast of Hwange National Park, 12 the main biological reserve of Zimbabwe (HNP; 19 °00'S, 26 °30'E, extending from Main 13 Camp to Giraffe's spring and Ngweshla pans; Fig. 1). This park covers 14650 km² and 14 supports a population of approx. 2800 giraffes (Shorrocks, 2016). The Bulawayo-Victoria 15 Falls railway line defines the western boundary of HNP while border with Botswana draws 16 the eastern boundary. The long-term mean annual rainfall is ca. 600 mm (CV = 25%) and 17 generally falls between October and April to form seasonal wetlands. Because of this 18 relatively low annual rainfall, a xerophile vegetation covers most of HNP. The woodland 19 vegetation consists primarily of African teak (Baikiaea plurijuga) intersected with patches of 20 camel thorn (Acacia erioloba) or leadwood (Combretum imberbe). Bushland savanna with 21 patches of grassland makes 64% of the area, mainly around the many artificially 22 maintained waterholes. HNP hosts many large and mega-herbivore species attractive to 23 oxpeckers, including giraffes, plain zebras (Equus guagga), African buffaloes, wildbeest 24 (Connochaetes taurinus), greater kudus, waterbuck (Kobus ellipsiprymnus) and impala, 25 and the less abundant sable and roan antelopes (*H. equinus*). 26

1 2.2 Oxpecker biology

Red-billed oxpecker (RBO) and yellow-billed oxpecker (YBO) are two sympatric species, 2 strictly african, that can be mostly found in Botswana, Namibia, South Africa and Zimbabwe 3 (Hustler, 1987; Plantan, 2009; Palmer & Packer, 2018). Although those birds feed mainly 4 on ticks, many studies have reported that oxpeckers can feed on wounded tissue, mucus, 5 saliva, ear wax, hair and blood (Bezuidenhout & Stutterheim, 1980; Weeks, 1999; Plantan, 6 2009). Some authors definitely think that the interaction between oxpekers and mammalian 7 hosts is parasitism, while others support mutualism even though they admit oxpeckers can 8 cause injuries to hosts (van Someren, 1951; Samish & Rehacek, 1999). In fact the 9 relationship between oxpeckers and hosts could be context-dependent, where birds can be 10 opportunists under particular biotic and abiotic conditions (Moreau, 1933; Nunn et al., 11 2011; Plantan, 2009). For instance, a mutualistic relationship may develop when the 12 ectoparasite load is high on the host, but oxpeckers may become parasitic when hosts 13 carry few ticks with too many birds, and leads to numerous open wounds. 14

15 2.3 Data collection

Oxpecker data derived from the study of giraffe ecology carried out at HNP. We have been 16 monitoring the giraffe population opportunistically in 2012 and 2013, and on a regular basis 17 since 2014, aiming at the photo-identification of individuals. Each year we drove the HNP 18 road network daily (Fig. 1) for at least three weeks in a row and shot pictures of every 19 encountered giraffes. For all encounters, we recorded the date, the location and the time of 20 observation along with group size and composition the individuals belonged to. We sexed 21 giraffes based on the presence of bare skin at the ossicone tip for males, and on shape of 22 the skull or visible male genitals when possible. We classified giraffes into four age-classes 23 (calf, juvenile, sub-adult and adult) assessed in the field from their size and coat color since 24 giraffes darken with age (Dagg, 2014). In this study we used pictures taken between 2013 25 and 2015, yielding a total of 500 photographs and a sample size of 683 giraffes. Although 26 we avoided to analyze a sequence of continuous pictures, re-observations of the same 27

individual in time might have occurred, either during the same field session or from one 1 year to another. Doing so, we assumed very little consequences of pseudo-replication on 2 the results because of the rapid exchanges and movements of oxpeckers among hosts. We 3 then counted oxpeckers on every giraffes seen on pictures and assigned each detected 4 bird to one of the 14 predefined parts of the giraffe body (see Fig. 3A). Giraffe body parts 5 were chosen so that they could easily be identified from landmark points whatever the point 6 of view. We excluded approx. 50% of all pictures because giraffes were too small to reliably 7 spot oxpeckers which roughly corresponded to a giraffe relative size smaller than 1/3 of the 8 picture height, or because of a poor picture guality. For a subset of pictures we repeated 9 oxpecker counts twice with two different observers (RG and CB) to estimate bird detection 10 probability. 11

12 2.4 Data analyses

We first estimated the detection probability of individual oxpeckers from pictures by setting 13 a double-observer experiment (Nichols et al., 2000). The two observers reported the total 14 number of detected birds they found (noted x_{11} and x_{22}) from which we calculated the 15 number of birds seen by observer 1 and missed by 2 (x_{12}) and conversely (x_{21}) . The 16 double-observer method returns the detection probability of observer 1 and 2 (respectively 17 p_1 and p_2), as well as the average detection probability of birds p. The estimated number of 18 oxpeckers per giraffe is then given by $\hat{N} = (x_{11} + x_{22} + x_{12} + x_{21})/\hat{p}$. We fit the 19 double-observer model to our data with the unmarked library (Fiske & Chandler, 2011) and 20 tested for the effect of host sex on the detection probabilities. 21

²² We derived the aggregation index *k* from a particular parameterization of the negative ²³ binomial distribution (where the variance *V* is related to the mean by $V = \lambda + \lambda^2/k$) that ²⁴ takes values close to zero with increasing levels of aggregation. To compare with previous ²⁵ studies, we also estimated the preference of oxpeckers for given hosts with a preference ²⁶ index (PI), calculated as the number of hosts counted divided by the number of oxpeckers ²⁷ counted. A PI of 5 means one bird is seen every fifth counted hosts on average. We ²⁸ computed confidence limits of point estimates of PI with a non-parametric bootstrap.

We estimated the relative distribution of oxpeckers on giraffe body parts with a 1 multinomial logistic regression. This particular type of GLM estimates the 14 probabilities 2 (noted π_i) of birds to be located on each body part of the giraffe holding the constrain that 3 $\sum_{i=1}^{14} \pi_i = 1$. To control for the imbalanced number of males and female giraffes among 4 years, we entered year as a categorical variable before testing for the effect the host sex (a 5 2-levels categorical variable) on the relative density of oxpeckers on body parts with 6 likelihood ratio tests using the nnet library (Venables & Ripley, 2002). 7 Finally, we modeled the number of oxpeckers per giraffe with generalized linear models 8 (GLMs), testing for effect of host sex (a 2-levels categorical variable), accounting for the 9 effects of time (year as a categorical variable), with likelihood ratio tests using the MASS 10 library (Venables & Ripley, 2002). We ran all analyses in the R 3.4 statistical software (R 11 Core Team, 2018). Unless otherwise stated, we reported all estimated parameters as 12 mean \pm sd and predicted probabilities as the mean with its associated 95% confidence 13 interval in brackets. 14

15 3 | RESULTS

From a subsample of n = 117 giraffes, the overall detection probability of oxpeckers was 16 $\hat{p} = 0.99 (0.97, 1.00)$ but differed substantially between the two observers (RG: 17 $\hat{p_1} = 0.97 (0.95, 0.98)$; CB: $\hat{p_2} = 0.85 (0.81, 0.89)$). Although oxperckers have 1.3 more 18 chances to be seen on a male than on female giraffe, detection probability did not differ 19 significantly between the host sex (estimated difference between male and female hosts on 20 the logit scale: $\beta = 0.26 \pm 0.34$, P = 0.45). On average we estimated oxpecker density to be 21 $D_O = 2.91$ (2.63, 3.23) birds per giraffe once we accounted for imperfect detections. In the 22 following analyses, RG did analyze all pictures. 23

²⁴ When using n = 683 giraffes, mean oxpecker density was 2.16 ± 3.01 birds per host ²⁵ without accounting for detection probability. The overall preference index (PI) is 0.46 ± 0.10 ²⁶ with a maximum number of oxpeckers counted on a single host of 17 (Fig. 2). In support of ²⁷ the aggregation hypothesis, the estimated aggregation coefficient $\hat{k} = 0.68$ (0.58, 0.79) ²⁸ suggested a strong aggregation of oxpeckers on individual giraffes. Because the estimated aggregation coefficient k approaches zero, the negative binomial distribution converges to

- ² the logarithmic series distribution, with a strong skew toward giraffe carrying no bird
- ³ (Fig. 2). Overall, our results lend support to the hypothesis of a non-random association
- ⁴ between oxpeckers and giraffes at HNP (H_1).
- 5 The relative distribution of oxpeckers on giraffe's body deviated strongly from uniformity
- ⁶ with some body parts being much more used than others (Fig. 3B). Supporting our
- ⁷ hypothesis H₂, birds gathered principally on the neck (π = 0.18 (0.15, 0.20)) and mane
- $_{\circ}$ (π = 0.25 (0.21, 0.27)) of giraffes, but were rarely seen on the lower limbs (π < 0.01) or on
- ⁹ the tail (π < 0.01). Oxpeckers did not use the giraffe's body differently according to the host

sex (likelihood ratio test: χ^2 = 28.68, df = 13, *P* = 0.07; Fig. 3B), although relatively more

birds used the ano-genital (β = 0.40 \pm 0.56, *P* = 0.47) and the scapula areas

 $_{12}$ (β = 0.45 ± 0.53, *P* = 0.39) of females compared to males. Conversely more birds were

- seen on the head (β = 1.53 ± 0.57, *P* < 0.001) and rump (β = 1.17 ± 0.54, *P* = 0.03) of
- ¹⁴ male giraffes. Our results hence confirm the marked heterogeneous distribution of

¹⁵ oxpeckers on the body of large mammalian hosts.

Although GLM with a Poisson distribution and a logarithmic link are usually 16 recommended for count data (Agresti 2002), a preliminary goodness-of-fit (GOF) test 17 suggested an over-dispersion of the data compared to a Poisson distribution (χ^2 = 2877.21, 18 df = 682, P < 0.001). A GLM a with negative binomial distribution did fit the data better 19 than with a Poisson distribution (GOF test: $\chi^2 = 609.10$, df = 682, P = 0.98). As expected 20 from our last hypothesis H₃, the number of oxpeckers was larger on the giraffes exposing 21 the largest body area to the birds (Fig 4). Accordingly, we found that oxpeckers were 20% 22 more numerous on males than on females (density of 2.60 ± 0.25 and 2.07 ± 0.14 birds per 23 giraffe respectively: $\beta = 0.23 \pm 0.12$, $\chi^2 = 3.75$, df = 1, P = 0.05). The sex-specific 24 aggregation coefficient reads $k = 0.85 \pm 0.09$ and $k = 0.62 \pm 0.07$ for females and males 25 respectively, and was significantly smaller for male giraffes (bootstrap test: $\beta = 0.23 \pm 0.13$, 26 P = 0.02). 27

1 4 DISCUSSION

The foraging behaviour and type of interaction between oxpeckers and their large 2 mammalian hosts is poorly understood and still debated (Weeks, 2000; Nunn et al., 2011; 3 Welsh et al., 2019). A closer look at the distribution of birds among and within giraffes at HNP clearly shows how heterogeneous it is at the host level with many carrying no bird 5 while a few has > 10 birds on them. We also provide empirical evidences for non-random 6 choice of host body part by oxpeckers, a behaviour likely driven by the amount of ticks birds 7 can find and share with conspecifics. Overall the observed oxpecker distribution among 8 giraffes at HNP matches with distributions generally observed in parasitic organisms, 9 although we reckon it may only mirror the tick load of individual hosts. 10 The distribution of oxpeckers on giraffes results, sequentially from the choice of an 11 individual giraffe in the landscape followed by the choice of a body part of this host, to prey 12 ultimately upon ectoparasites. At the largest spatio-temporal scale, oxpeckers have to 13 chose among large herbivore species, which is reflected by the host's preference index 14 (PI). The observed PI = 0.46 ± 0.10 for giraffes at HNP is similar to previously reported 15 values in Kruger National Park (KNP), South Africa (0.90 for Grobler (1980); 0.54 for 16 Ndlovu & Combrink (2015); 0.51 for Welsh et al. (2019) in Kenya). Surprisingly, Hustler 17 (1987) found PI of 5.39 and 6.71 for giraffes in two separate areas of HNP. Host availability 18 should indeed influence oxpeckers' choice because the decrease in abundance of a key 19 host such as the giraffe may force birds to switch to another less preferred but more 20 numerous host with little fitness costs (Pyke, 1984; Hustler, 1987; Welsh et al., 2019). 21 Here, host size plays a major role in host detection in flight and giraffes – like other large 22 mammalian hosts – are easier to detect compared with smaller species (Grobler, 1980; 23 Koenig, 1997). This is the main reason why the key host of oxpeckers alternates between 24 buffaloes (Hustler, 1987), white rhinoceros (Ndlovu & Combrink, 2015) and giraffes 25 (Grobler, 1980; Ndlovu & Combrink, 2015). The past high PI of giraffes at HNP suggests 26 that birds must have exploited others hosts in the 80ies such as black rhinoceros, white 27 rhinoceros, roans and sables (see Hustler, 1987, for details). This interpretation is 28

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supported by change in the composition of HNP's community of large herbivores over the

² last decades with, notably, the recent loss of the white rhinoceros (Valeix et al., 2008),

³ which is consistent with the low PI values we report here for giraffes.

Focusing on the choice of individual giraffes by oxpeckers, we found a marked asymmetric distribution of birds (exponential distribution) whereby many carried no bird and 5 a few ones were seen with up to 17 birds simultaneously (Fig. 2). This non-random 6 distribution of oxpeckers among individual giraffes usually likely results from an aggregation 7 behaviours (Palmer & Packer, 2018). The aggregation coefficient k we found for the 8 oxpecker distribution at HNP, close to 0, is typical of parasitic infections where only a few 9 individuals are massively infested (Shaw et al., 1998). That oxpeckers similarly aggregate 10 on a few giraffes would suggest they behave like parasites with their host in agreement with 11 previous studies (Plantan, 2009; Nunn et al., 2011). Birds could use public information like 12 conspecific density to chose a giraffe in a group (Doligez et al., 2004). Because oxpeckers 13 mostly feed on ectoparasites, the marked aggregation of birds could indirectly mirror the 14 distribution of ticks among giraffes. In mammals, infestation is indeed highly variable 15 among hosts (e.g. Shaw et al., 1998; Brunner & Ostfeld, 2008). For instance, in roe deer 16 (Capreolus capreolus), bank vole (Clethrionomys glareolus) and mountain hare (Lepus 17 timidus), most host individuals bear few ticks and only a few individuals bear many (Horak 18 et al., 1983; Talleklint & Jaenson, 1997; Mysterud et al., 2014). From this hypothesis, one 19 could make indirect inference on tick burden of individual giraffes from the number of 20 hosted oxpeckers, given the birds distribute themselves according to the ideal free 21 distribution. 22

Within giraffes, we clearly found preferences for some body parts by oxpeckers. At HNP, oxpeckers were mostly found on the neck and the back of giraffes followed by the head, the abdomen, the lower limbs and the tail (see also Plantan, 2009; Ndlovu & Combrink, 2015, for similar results). The mane seems the most preferred giraffes' body part of oxpeckers (Koenig, 1997). This row of hairs seems to be a favourable habitat for ectoparasites by providing shelter from predators (Ndlovu & Combrink, 2015) although oxpeckers use a scissoring behaviour to easily pick parasites from the hairs Koenig (1997). It has been

noted that oxpeckers gather at the bigger host's mane to hide from predators or when they 1 are alarmed, which may contribute to increase their number in this area. However, because 2 giraffes rarely feed directly on the ground (Seeber et al., 2012) we expected a relatively low 3 density of ticks on the giraffe's head. Our results support this assertion but oxpeckers could 4 sometimes forage the head seeking for other food resources such as saliva, mucus, earwax 5 (Ndlovu & Combrink, 2015) or wounds. This also could be the case for female genitalia 6 where oxpeckers can also feed on mucus and secretion of their hosts (Weeks, 1999; 7 Plantan, 2009). The abdomen, groins, thighs and tail present the lowest density of 8 oxpeckers. Unlike mane, these areas are parts that can be easily groomed by giraffes, 9 depleting tick quickly and making this area less preferred for oxpeckers (Koenig, 1997; 10 Ndlovu & Combrink, 2015). Assuming that abundance or presence of ticks is highest in the 11 mane running on the neck and back of giraffes, our results would concur with the 12 distribution of oxpeckers as predicted by the optimal foraging theory (Pyke, 1984). 13 Density of oxpeckers differed substantially according to the sex of the giraffe, with male 14

hosts carrying 20% more birds than females (Fig. 4). This higher abundance of oxpeckers 15 on male than on female giraffes likely proceed from their higher load in ectoparasites and 16 hence, larger food resources for the birds. The simplest explanation is that male giraffes 17 being larger than females, more birds can feed on a male holding a constant per capita 18 food rate. Alternative explanations may also account for the preference of male hosts by 19 oxpeckers, ranging from male-specific parasitic load, to the aggressive behaviour. Previous 20 studies who investigated the effect of host sex on the ectoparasite load are equivocal in 21 mammals. For instance, Horak et al. (1987) reported more ticks (Amblyomma hebraeum) 22 on male kudus, which could make the female less attractive to oxpeckers. 23

²⁴ Conversely, another study carried out in Scandinavia found that tick load was similar
²⁵ whatever the sex and age of red deer (*Cervus elaphus*) (Mysterud et al., 2014). A
²⁶ proximate mechanism for why male giraffes would carry more ticks than females is
²⁷ intra-sexual fights for reproduction. Neck fight, opponent chasing and female mounting
²⁸ indeed result in males having many injuries and open wounds all over their body (Nunn
²⁹ et al., 2011). Being opportunistic feeders, oxpeckers benefit from the higher wound- and

tick-feeding opportunities on male giraffes (Plantan, 2009). Alternatively, the handicap
principle of the sexual selection (Zahavi, 1975) proposes that, by surviving with many
parasites or extravagant sexually selected traits, males would display honestly their intrinsic
quality to females. Hence, oxpeckers could play the role of in indirect signal of ectoparasite
load of male giraffes.

To evaluate the reliability of the oxpecker detection and location on giraffes from 6 pictures, we carried out a double-observer experiment on a sub-sample of our images. We 7 found the overall detection of the birds from photos to be very high (99%) but ideally all 8 should be analyzed by 2 observers. Although not perfect, one person only (RG) scrutinized 9 the 500 photographs henceforth keeping the detection and condition of observation similar 10 for the complete data set. A major advantage of oxpecker counts from pictures is to make 11 the counts and analyses repeatable and, contrary to what one might think, the use of one 12 side of giraffes to locate and to count birds is an advantage because it avoids the issue of 13 double counting. That said, we acknowledge that the major bias of our study was the 14 location where the giraffe pictures were taken *i.e.* mainly around the many artificially 15 maintained waterholes where large herbivores come to drink. When a large herbivore stand 16 on the shore of waterholes, oxpeckers often use it as platform to rest, to sunbath, and to 17 reach water to drink as well (Stutterheim, 1976). Consequently, the maximum number of 18 oxpeckers per giraffe may be higher than elsewhere in HNP. To avoid this bias some 19 studies tend to limit counting within 500 meters of water points (Grobler, 1980) but because 20 HNP is densely covered with trees, observations of giraffes away from waterholes remained 21 very difficult. 22

23 5 CONCLUSION

Our study puts forward that the distribution and abundance of oxpeckers were surprisingly heterogeneous among and within giraffes. Some host body parts are clearly preferred for foraging by birds such as the neck and the mane because those areas could be suitable habitats for ticks. Gregarious hosts (female giraffes, buffaloes) travel and forage as a group thereby increasing local abundance and transmission of ticks (Koenig, 1997) to which

- 1 oxpeckers could be excellent control agents on wild large herbivores and on domestic ones
- ² too (Ndlovu & Combrink, 2015). From an ecological point of view, the oxpecker-large
- ³ herbivores system proves to be highly relevant and useful for the study of host-parasite
- 4 dynamics.

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10 References

- ¹¹ Bezuidenhout, J. D., & Stutterheim, C. (1980). A critical evaluation of the role played by the
- red-billed oxpecker *Buphagus erythrorhynchus* in the biological control of ticks. *The*
- ¹³ Onderstepoort Journal of Veterinary Research, 47, 51–75.
- Brunner, J. L., & Ostfeld, R. S. (2008). Multiple causes of variable tick burdens on small-mammal
 hosts. *Ecology*, *89*, 2259–2272.
- ¹⁶ Clark, C. W., & Mangel, M. (1986). The evolutionary advantages of group foraging. *Theoretical*
- 17 population biology, 30, 45–75.
- ¹⁸ Dagg, A. I. (2014). *Giraffe: biology, behaviour and conservation*. Cambridge University Press.
- ¹⁹ Doligez, B., Pärt, T., Danchin, E., Clobert, J., & Gustafsson, L. (2004). Availability and use of public
- ²⁰ information and conspecific density for settlement decisions in the collared flycatcher. *Journal of*
- 21 Animal Ecology, 73, 75–87.
- ²² Emlen, S. T., & Oring, L. W. (1977). Ecology, sexual selection, and the evolution of mating systems.
- 23 Science, 197, 215–223.
- ²⁴ Fiske, I., & Chandler, R. (2011). unmarked: An R package for fitting hierarchical models of wildlife
- occurrence and abundance. *Journal of Statistical Software*, 43, 1–23.
- ²⁶ Fretwell, S. D., & Lucas, H. L. (1969). On territorial behavior and other factors influencing habitat
- distribution in birds. Acta Biotheoretica, 19, 16–36. doi:10.1007/bf01601953

- Grobler, J. H. (1980). Host selection and species preference of the red-billed oxpecker Buphagus
- 2 erythrorhynchus in the Kruger National Park. Koedoe, 23. doi:10.4102/koedoe.v23i1.637
- ³ Gueron, S., & Levin, S. A. (1995). The dynamics of group formation. *Mathematical Biosciences*,
- 4 *128*, 243–264.
- 5 Horak, I., MacIvor, K., Petney, T., & De, V. V. (1987). Some avian and mammalian hosts of
- 6 Amblyomma hebraeum and Amblyomma marmoreum (Acari: Ixodidae). The Onderstepoort
- Journal of Veterinary Research, 54, 397–403.
- 8 Horak, I., Potgieter, F., Walker, J. B., De Vos, V., & Boomker, J. (1983). The ixodid tick burdens of
- ⁹ various large ruminant species in South African nature reserves. *Onderstepoort Journal of*
- ¹⁰ *Veterinary Research*, *50*, 221–228.
- Hustler, K. (1987). Host preference of oxpeckers in the Hwange National Park, Zimbabwe. African
- Journal of Ecology, 25, 241–245. doi:10.1111/j.1365-2028.1987.tb01115.x
- Hutchinson, J. M., & Waser, P. M. (2007). Use, misuse and extensions of "ideal gas" models of
- animal encounter. *Biological Reviews*, *82*, 335–359.
- Johnson, D. H. (1980). The comparison of usage and availability measurements for evaluating
- resource preference. *Ecology*, *61*, 65–71.
- ¹⁷ Koenig, W. D. (1997). Host preferences and behaviour of oxpeckers: co-existence of similar species
- in a fragmented landscape. *Evolutionary Ecology*, *11*, 91–104. doi:10.1023/a:1018439614008
- ¹⁹ Krebs, C. J. (1972). The experimental analysis of distribution and abundance. *New York: Harper*
- 20 and Row.
- Lack, D. L. (1966). Population studies of birds. Clarendon P.
- ²² Lotka, A. J. (1956). Elements of mathematical biology.
- MacArthur, R. H. (1972). *Geographical ecology: patterns in the distribution of species*. Princeton
 University Press.
- ²⁵ May, R. M. (1978). Host-parasitoid systems in patchy environments: a phenomenological model.
- Journal of Animal Ecology, 47, 833–843.
- ²⁷ Morales, J. M., Moorcroft, P. R., Matthiopoulos, J., Frair, J. L., Kie, J. G., Powell, R. A., Merrill, E. H.,
- ²⁸ & Haydon, D. T. (2010). Building the bridge between animal movement and population dynamics.
- 29 Philosophical Transactions of the Royal Society B: Biological Sciences, 365, 2289–2301.
- ³⁰ Moreau, R. E. (1933). The food of the red-billed oxpecker, *Buphagus erythrorhynchus* (Stanley).
- Bulletin of Entomological Research, 24, 325. doi:10.1017/s000748530003162x
- 32 Mysterud, A., Hatlegjerde, I. L., & Sørensen, O. J. (2014). Attachment site selection of life stages of

- 1 Ixodes ricinus ticks on a main large host in Europe, the red deer (Cervus elaphus). Parasites &
- ² Vectors, 7. doi:10.1186/s13071-014-0510-x
- 3 Ndlovu, M., & Combrink, L. (2015). Feeding preferences of oxpeckers in Kruger National Park,
- 4 South Africa. *Koedoe*, *57*. doi:10.4102/koedoe.v57i1.1316
- 5 Nichols, J. D., Hines, J. E., Sauer, J. R., Fallon, F. W., Fallon, J. E., & Heglund, P. J. (2000). A
- 6 double-observer approach for estimating detection probability and abundance from point counts.
- ⁷ The Auk, 117, 393–408.
- 8 Nunn, C. L., Ezenwa, V. O., Arnold, C., & Koenig, W. D. (2011). Mutualism or parasitism? Using a
- 9 phylogenetic approach to characterize the oxpecker-ungulate relationship. *Evolution*, 65,
- ¹⁰ 1297–1304. doi:10.1111/j.1558-5646.2010.01212.x
- ¹¹ Ostfeld, R. S., & Keesing, F. (2000). Pulsed resources and community dynamics of consumers in
- terrestrial ecosystems. *Trends in Ecology & Evolution*, *15*, 232–237.
- ¹³ Palmer, M. S., & Packer, C. (2018). Giraffe bed and breakfast: Camera traps reveal Tanzanian
- yellow-billed oxpeckers roosting on their large mammalian hosts. *African Journal of Ecology*.
- ¹⁵ doi:10.1111/aje.12505
- Pielou, E. (1969). An introduction to mathematical ecology. *An introduction to mathematical ecology.*
- 18 Plantan, T. B. (2009). Feeding behaviour of wild and captive oxpeckers (Buphagus spp.): a case of
- 19 conditional mutualism. Ph.D. thesis, University of Miami.
- ²⁰ Pyke, G. H. (1984). Optimal foraging theory: A critical review. Annual Review of Ecology and
- 21 Systematics, 15, 523–575. doi:10.1146/annurev.es.15.110184.002515
- 22 R Core Team (2018). R: A Language and Environment for Statistical Computing. R Foundation for
- 23 Statistical Computing, Vienna, Austria.
- ²⁴ Samish, M., & Rehacek, J. (1999). Pathogens and predators of ticks and their potential in biological
- control. Annual Review of Entomology, 44, 159–182. doi:10.1146/annurev.ento.44.1.159
- ²⁶ Seeber, P. A., Ndlovu, H. T., Duncan, P., & Ganswindt, A. (2012). Grazing behaviour of the giraffe in
- Hwange National Park, Zimbabwe. *African Journal of Ecology*, *50*, 247.
- 28 Shaw, D., Grenfell, B., & Dobson, A. (1998). Patterns of macroparasite aggregation in wildlife host
- populations. *Parasitology*, *117*, 597–610.
- ³⁰ Shorrocks, B. (2016). *The giraffe: biology, ecology, evolution and behaviour*. John Wiley & Sons.
- 31 Sjöberg, M., Albrectsen, B., & Hjältén, J. (2000). Truncated power laws: a tool for understanding
- aggregation patterns in animals? *Ecology letters*, *3*, 90–94.

- 1 Stutterheim, C. (1976). The biology of the red-billed oxpecker, Buphagus erythrorhynchus (Stanley
- ² 1814) in the Kruger National Park. Master's thesis, University of Pretoria, South African.
- 3 Stutterheim, I. M., Bezuidenhout, J. D., & Elliot, E. (1988). Comparative feeding behaviour and food
- ⁴ preferences of oxpeckers (*Buphagus erythrorhynchus* and *B. africanus*) in captivity. *The*
- 5 Onderstepoort Journal of Veterinary Research.
- 6 Talleklint, L., & Jaenson, T. G. (1997). Infestation of mammals by Ixodes ricinus ticks (Acari:
- ⁷ Ixodidae) in south-central Sweden. *Experimental & applied acarology*, 21, 755–771.
- 8 Taylor, L., Woiwod, I., & Perry, J. (1979). The negative binomial as a dynamic ecological model for
- ⁹ aggregation, and the density dependence of *k*. *The Journal of Animal Ecology*, (pp. 289–304).
- ¹⁰ Valeix, M., Fritz, H., Chamaillé-Jammes, S., Bourgarel, M., & Murindagomo, F. (2008). Fluctuations
- in abundance of large herbivore populations: insights into the influence of dry season rainfall and
- elephant numbers from long-term data. *Animal Conservation*, *11*, 391–400.
- van Someren, V. D. (1951). The red-billed oxpecker and its relation to stock in Kenya. The East
- African Agricultural Journal, 17, 1–11.
- ¹⁵ Venables, W. N., & Ripley, B. D. (2002). *Modern Applied Statistics with S.* New York: Springer,
- 16 fourth edition. ISBN 0-387-95457-0.
- ¹⁷ Veríssimo, D., Cugnière, J.-C., Cugnière, S., Cugnière, J., Cugnière, G., & Cugnière, L. (2017).
- 18 Record number of yellow-billed oxpeckers *Buphagus africanus* Linnaeus, 1766 (Aves:
- ¹⁹ Passeriformes: Buphagidae) foraging on a single host. *Journal of Threatened Taxa*, *9*, 9768.
- ²⁰ doi:10.11609/jott.3055.9.1.9768-9770
- ²¹ Weeks, P. (1999). Interactions between red-billed oxpeckers, *Buphagus erythrorhynchus*, and
- domestic cattle, *Bos taurus*, in Zimbabwe. *Animal Behaviour*, *58*, 1253–1259.
- ²³ doi:10.1006/anbe.1999.1265
- Weeks, P. (2000). Red-billed oxpeckers: vampires or tickbirds? *Behavioral Ecology*, *11*, 154–160.
 doi:10.1093/beheco/11.2.154
- ²⁶ Welsh, E. C., Keesing, F., & Allan, B. F. (2019). Oxpecker (*Buphagus erythrorhynchus, Buphagus*
- *africanus*) and tick abundances in acaricide-treated livestock areas. *African Journal of Ecology*,
- ²⁸ *57*, 155–159.
- ²⁹ Wittenberger, J. F. (1981). Animal social behavior. Duxbury Press.
- Zahavi, A. (1975). Mate selection a selection for a handicap. *Journal of theoretical Biology*, *53*, 205–214.

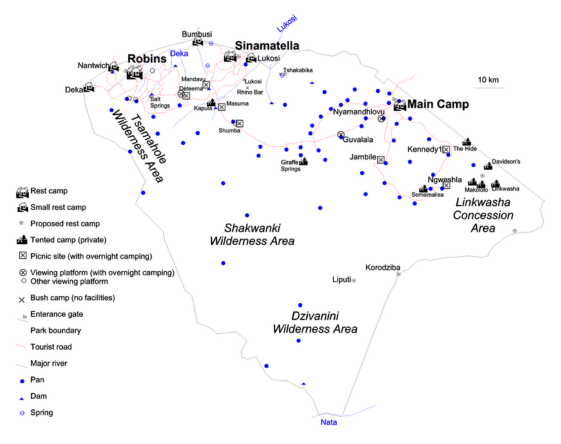


Fig. 1 Hwange National Park map, Zimbabwe. The data collected (2013-2015) from oxpeckers (*Buphagus sp.*) and giraffes (*Giraffa camelopardalis*) derived from a study located in the Main Camp area, covering the north east of the park from Ngweshla to Giraffe Springs passing through Jambile. ('Hwange National Park' 2019, in Wikipedia: The Free Encyclopedia, Wikimedia Foundation Inc., viewed 11 May 2019,

<https://en.wikipedia.org/wiki/Hwange_National_Park>)

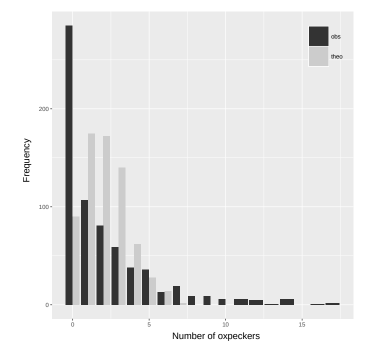
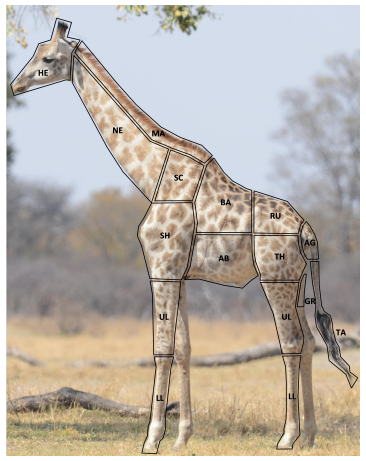


Fig. 2 Distribution of oxpeckers (*Buphagus sp.*) on individual giraffes (*Giraffa camelopardalis*) at Hwange National Park, Zimbabwe. In black is the observed distribution and in grey the expected distribution according to a Poisson distribution law taking the observed mean as parameter ($\lambda = 2.16$). Note the marked over-representation of giraffes carrying no bird and the long distribution tail of giraffes with numerous birds on their body in the observed data.



(A)

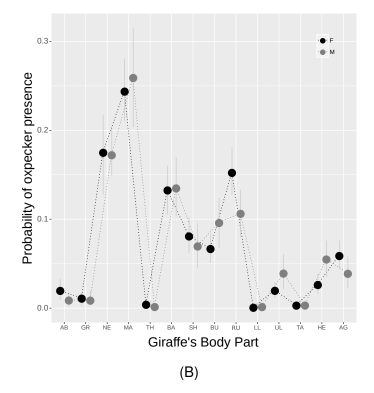


Fig. 3 (A) Defined zonation and (B) proportion of oxpeckers (*Buphagus sp.*) counted on the 14 different body parts of giraffes (*Giraffa camelopardalis*) observed at Hwange National Park, Zimbabwe. The 14 different body parts we defined were: AB: abdomen, GR: groin, NE: neck, MA: mane, TH: thigh, BA: back, SH: shoulder, RU: rump, SC: scapula, LL: lower leg, UL: upper leg, TA: tail, HE: head, AG: ano-genital. Although the general patterns were similar the use of giraffe body parts by oxpeckers differed significantly (*n* = 683 giraffes).

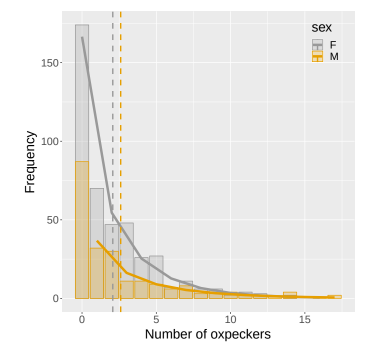


Fig. 4 Distribution of oxpeckers (*Buphagus sp.*) on male and female giraffes (*Giraffa camelopardalis*) at Hwange National Park, Zimbabwe. Curves are the predicted frequencies as given by a negative binomial distribution model which parameters have been estimated separately for the two sexes. Vertical dashed lines represent the mean number of oxpeckers carried by individual giraffes. Note that males with no oxpecker are less frequent than females, and that the largest aggregations of oxpeckers have been found on male giraffes.