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Naturally-occurring tooth wear, tooth fracture, and cranial injuries in large carnivores from Zambia

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ABSTRACT

Determining the incidence and causes of craniodental damage in wild carnivores is often constrained by limited access to specimens with associated ecological data, such as prey type and abundance. We assessed dental condition and cranial injuries in lion, leopard, and spotted hyena in relation to prey and predator populations in Zambia's Luangwa Valley, where large prey are more abundant and lion and leopard more numerous, and the Greater Kafue Ecosystem, where smaller prey species are more prevalent and lion and leopard less common. In Luangwa, lions had significantly higher rates of tooth fracture, and blunt trauma injuries attributable to prey-handling, compared to Kafue lions. In contrast, leopards in both regions had similar rates of tooth wear and breakage. Overall, lions showed a significantly higher tooth fracture rate than leopards on a per tooth basis. Spotted hyenas had the highest rates of tooth wear and fracture among all three carnivores, and greatly exceeded previously recorded rates based on historical samples. Despite larger numbers of lion and leopard in Luangwa, there was no difference in incidence of intraspecific injuries between regions. These results are consistent with a greater abundance of large prey species, especially buffalo, in the diets of Luangwa lions, and previous work showing a reliance on smaller prey species in Kafue throughout the large carnivore guild.

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INTRODUCTION

Large (>21 kg) carnivores lead a risky life. Because of their size, they need to kill prey that are as large, or larger, than themselves (*Carbone et al., 1999; Carbone, Teacher & Rowcliffe, 2007*). Their typical prey, ungulates, are difficult to subdue, wielding weapons (hooves, horns, antlers) that can and do injure their attackers, fracturing their limbs and skulls in some cases (e.g., *Rausch, 1967; Creel & Creel, 2002; Mukherjee & Heithaus, 2013*).

The teeth of large carnivores are often broken, due to stresses incurred while killing prey, consuming bone, and in combat with conspecifics as well as other competitors (*Van Valkenburgh, 1988, 2009; Mann, Van Valkenburgh & Hayward, 2017*). Undoubtedly, the risks of being a large carnivore have shaped their evolution, but we have little data on the

frequency and severity of injuries sustained by large predators. Injury incidence is difficult to quantify from field observations; broken bones mend and cracked teeth are not easily seen from a distance. Nevertheless, skeletal and dental trauma is often recorded in the bones and teeth of individuals, and surveys of natural history collections have provided insights into the frequency and distribution of trauma in extinct and extant carnivores (e.g., [Wobeser, 1992](#); [Wilkins et al., 2007](#); [Binder, Thompson & Van Valkenburgh, 2009](#); [Losey et al., 2014](#); [Brown et al., 2017](#); [Collados, Garcia & Rice, 2018](#); [Tong et al., 2020](#)).

When considering craniodental injuries, free-ranging specimens are required because dental pathologies, tooth wear, and fracture vary significantly in captive versus wild carnivores ([Wenker et al., 1999](#); [Curtis et al., 2018](#)). Museum collections represent valuable archives of skeletal material for comparative osteological and dental investigations, but historic specimens often lack logistical and/or natural history information ([De la Sancha, Boyle & Patterson, 2017](#)), limiting interpretation of results. As an alternative source of contemporary specimens, many species of carnivores are legally hunted, and the associated permitting procedures require reporting of relevant data such as date and location. However, hunted specimens are usually sequestered in private collections, making them difficult to access and consequently they remain underutilized for purposes of scientific investigations.

Here, we report on tooth wear, tooth breakage, and cranial injuries in free-ranging lion (*Panthera leo*), leopard (*P. pardus*), and spotted hyena (*Crocuta crocuta*) from two regions, the Luangwa Valley (LV) and the Greater Kafue Ecosystem (GKE) in Zambia between 2007 and 2012. Cranial material was obtained primarily from hunted specimens with the exception of two lions that were killed by Zambia's Department of National Parks and Wildlife (DNPW) as problem animals owing to threats to humans or livestock. Tooth wear and fracture were recorded for each specimen, as well as evidence of antemortem osteological pathologies such as blunt trauma and bite marks, indicative of injuries potentially associated with prey-handling and intraspecific conflict, respectively.

Previous work on dental trauma in large carnivores suggests that, within age classes, variation in tooth wear and tooth breakage among species is largely reflective of diet, especially bone consumption. Species such as spotted hyenas that frequently consume large bones have higher rates of tooth fracture and heavier wear than species that consume less bone, such as cheetahs (*Acinonyx jubatus*) ([Van Valkenburgh, 1988, 2009](#); [Mann, Van Valkenburgh & Hayward, 2017](#)). Moreover, tooth fracture frequency within a species is likely to increase as large carcasses are consumed more completely, a feeding behavior that is likely to be favored when prey are difficult to acquire ([Mech & Frenzel, 1971](#); [Carbone et al., 2005](#); [Vucetich, Vucetich & Peterson, 2012](#)). A recent comparative study of extant gray wolf (*Canis lupus*) populations found that wolves from regions or time periods characterized by high numbers of prey had significantly less tooth wear and fracture, while wolf populations with reduced prey availability were associated with greater carcass utilization, higher bone consumption, and more tooth damage ([Van Valkenburgh et al., 2019](#)).

Moreover, predators confront greater danger from injury when tackling large prey species ([Rausch, 1967](#)). In Zambia, the LV and GKE differ significantly in the relative

abundance of prey, in particular, very large (>500 kg) herbivore species with the former region having greater numbers of Cape buffalo (*Syncerus caffer*), hippopotamus (*Hippopotamus amphibius*), and elephant (*Loxodonta africana*) as well as a small population of endemic Thornicroft's giraffe (*Giraffa camelopardalis thornicrofti*) (DNPW, 2016; IUCN, 2016), and each of these species factor importantly into the diet of LV lions (Zambian Carnivore Programme unpublished, in Creel et al. (2018)). In contrast, small to medium-sized herbivores are currently the most numerous prey species in GKE (DNPW, 2016; Creel et al., 2018). Therefore, we hypothesized that capturing, killing, and feeding (including scavenging) on very large prey might result in elevated tooth fracture rates, as well as increased incidents of blunt trauma in LV than GKE carnivores of comparable ages. Lions (IUCN, 2006) and possibly leopards (Rosenblatt et al., 2016) are also more numerous in LV than GKE, and increased intraspecific competition and conflict in LV might be reflected in a greater number of cranial injuries indicative of fighting, such as bite wounds and claw marks.

Our purpose in this paper is to explore the array and frequency of craniodental injuries that are apparent in three species of large African carnivores in each of two regions in Zambia.

MATERIALS & METHODS

Study area

The Republic of Zambia occupies 752,614 km² in southcentral Africa within 15° 00' S and 30° 00' E. Samples were obtained from two regions, Luangwa Valley (LV) in the east and the Greater Kafue Ecosystem (GKE) in the west (Fig. 1). Both regions are comprised of national parks (NPs) and adjacent game management areas (GMAs). The LV contains four NPs totaling 15,630 km² that, combined with the adjacent GMAs, covers 62,038 km² (Astle, 1999). The GKE is an approximately 66,547 km² area comprised of Kafue NP (KNP) (22,400 km²) and adjacent GMAs (Siamudaala, Nyirenda & Saiwana, 2009). There are no fences or other barriers, and animals routinely cross between NPs and GMAs. Legal trophy hunting for lion, leopard, and spotted hyena occurs in most GMAs. Two lions killed by the wildlife authority (DNPW) as problem animals that were 10–15 km outside of LV ($n = 1$) and GKE ($n = 1$) were assigned to the geographically closest region.

Obtaining specimens

Between 2007 and 2012, one of us (PAW) examined and photographed skulls and teeth of hunted lion, leopard, and spotted hyena as part of a larger study on age estimation of trophy hunted lions (White & Belant, 2016; White et al., 2016). Skulls of hunted carnivores were made available by professional hunters and taxidermists. Skulls of two lions killed as problem animals were made available by DNPW. All osteological material was examined and photographic data collected in Zambia in partnership with DNPW (formerly Zambia Wildlife Authority, Research/Employment Permit No. #008872). A total of 118 lion (LV 60; GKE 58), 45 leopard (LV 18; GKE 27) and 13 spotted hyena (LV 6; GKE 7) skulls were evaluated in this study. Due to post-mortem damage or incomplete

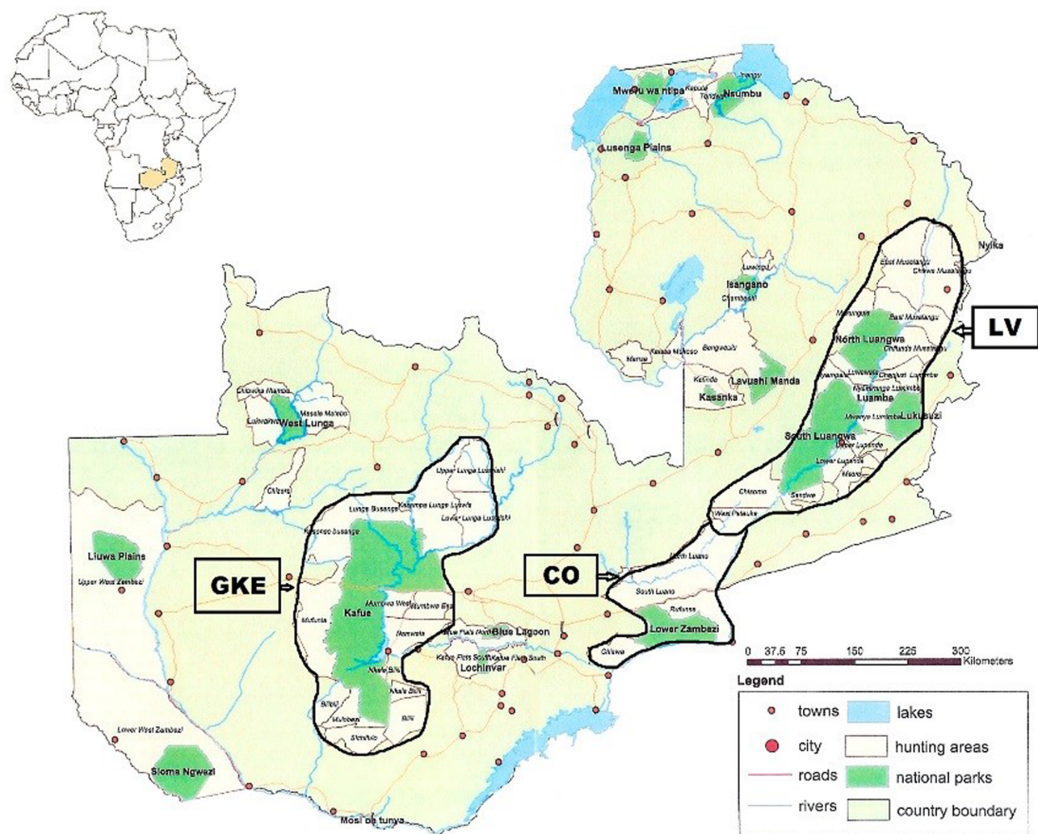


Figure 1 Map showing location of Luangwa Valley and Greater Kafue Ecosystem in Zambia. Location map of Zambia in southcentral Africa showing the Luangwa Valley (LV) and the Greater Kafue Ecosystem (GKE). The corridor area (CO) consists of contiguous habitat between the Luangwa Valley and Lower Zambezi river. [Full-size !\[\]\(b345a1c4255362eec3746050dd71ccac_img.jpg\) DOI: 10.7717/peerj.11313/fig-1](https://doi.org/10.7717/peerj.11313/fig-1)

photographic views, not all lion skulls could be used in every analysis. Thus, samples sizes for lion varied slightly as follows: tooth wear and breakage 115 lions (LV 58; GKE 57); cranial injury 116 lions (LV 60; GKE 56).

Lion hunting in Zambia is restricted to males. Sex of sampled lions was confirmed from hides or trophy photographs. We restricted our sample of problem lions to include only males. Leopard hunting in Zambia is restricted to males, and sampled leopards were assumed to be males. However, trophy photographs or hides were not available for all sampled leopards, and the possibility cannot be excluded that some hunted individuals were females (*Spong, Hellborg & Creel, 2000*). Because spotted hyena are encountered less frequently, the species is taken more opportunistically and hunter selectivity is lower than for lion or leopard. Consequently, the spotted hyena sample likely consisted of both female and male specimens. For each of the three species, we assumed that hunter bias in selectivity was similar among the two regions.

Photographs

We captured digital images of each skull using a Nikon 35mm D3300 digital camera with Nikkor AF 70–300 mm f/4.5-6.3 lens (Nikon USA Inc., Melville, NY, USA).

We photographed skulls with the mandible articulated from left and right lateral sides, anterior, occipital and dorsal views. Thereafter, we disarticulated the mandible and inverted the cranium to obtain a palatal view and dorsal and lateral views of the mandibles. Digital images were organized by a field ID number assigned to each specimen and stored on SD cards for later examination on a computer and display monitor.

Age estimation of samples

Because the probability of having broken a tooth or suffered an injury increases with age, it is important to control for age differences between samples from the two regions. We assigned age classes to each sampled lion, leopard, and spotted hyena based on published standards (below) that represent the best available methods of age estimation for each species. We assigned specimens to one of three age classes initially with skulls in-hand and verified age assignments later using digital images.

In lions, crown wear of the upper second premolar (P2) has been shown to increase with age ([Schaller, 1972](#); [Smuts, Anderson & Austin, 1978](#); [White & Belant, 2016](#)). We visually scored P2 crown wear as: (1) no wear/sharp, (2) moderate wear/rounded, or (3) heavy wear/flat, and used P2 scores in conjunction with overall tooth wear features (following [Smuts, Anderson & Austin, 1978](#); [Miller et al., 2016](#)) in assigning each lion to one of three age classes: young adult < 5 years; mature adult $\geq 5-7 <$ years; and old adult $\geq 7+$ years ([Fig. S1](#)). Assignment to the mature adult age class was confirmed in most cases by the partially to fully obliterated closure of the interfrontal suture after [Smuts, Anderson & Austin \(1978\)](#) ([Fig. S2](#)). We assigned each leopard to an age class based on wear of the upper and lower canines (C1, c1) and premolars (P3, p3,4) following the criteria of [Stander \(1997\)](#) with the modification that younger age classes were combined as follows: young adult ≤ 4 years; mature adult 5–6 years, and old adult 7–10 years ([Fig. S3](#)). We assigned spotted hyena to one of three age classes (young 1–3 years; mature > 3–6 years; old > 6–16 years) based on wear of the lower third premolar (p3) after [Kruuk \(1972\)](#) ([Fig. S4](#)). We tested for differences in age structure of sampled lion and leopard from LV vs. GKE using chi-square performed in SPSS v.26 with significance levels set at $P < 0.050$.

Scoring of dental condition

For the regional comparisons, we scored tooth breakage and overall wear from digital photographs alone. Scoring of tooth breakage and wear was performed subsequent to, and independent of, age estimation, and involves a more comprehensive review of wear across the entire tooth row as opposed to focusing on one or a few specific teeth (e.g. canines, premolars). Notably, PAW did the age estimation and BVV did the subsequent dental wear and fracture analyses, blind to the age estimates. As in earlier studies ([Van Valkenburgh, 1988](#); [Van Valkenburgh & Hertel, 1993](#); [Van Valkenburgh, 2009](#)), breakage was scored by tooth position and type, i.e., incisors, canines, premolars, and molars. A tooth was counted as broken in life (antemortem) only if there was clear evidence of subsequent wear on the tooth following breakage ([Fig. S5](#)). Teeth broken at time of death or post-mortem with apparently unworn, sharp edges were not counted as broken and were excluded from analyses.

We assigned one of five tooth wear stages to each individual: (1) little or no apparent wear observed on shear facets or blunting of cusps, (2) shear facets apparent on carnassial teeth and cusps slightly blunted on some teeth, (3) shear facets apparent on carnassial teeth and cusps blunted on most teeth, (4) carnassial teeth exhibiting strong shear facets and moderate blunting of premolar cusps, or (5) carnassial teeth exhibiting strong shear facets and/or blunted cusps, and premolars and molars with well-rounded cusps. The five categories were then reduced to three for the analysis, (1) “slight” (stage 1 only), (2) “moderate” (stages 2 and 3), and (3) “heavy” (stages 4 and 5) (Fig. S6).

We summed tooth wear stage distribution and fracture frequencies by species and location (LV or GKE) and, for lion and leopard, compared distribution of wear stage between regions using chi-square with significance set at $P < 0.050$. Interspecific comparisons using the entire dataset were made on (1) number of broken teeth as a percentage of the total number of teeth, (2) percentage of individuals with one or more broken teeth, and (3) percentage of tooth fracture by tooth position (incisors, canines, premolars, carnassials). For lion and leopard, the percentage of broken teeth and breakage by tooth position for each species were compared between samples from LV and GKE using chi-square with significance set at $P < 0.050$. Sample size of spotted hyena ($n = 13$) was insufficient to compare between the two regions.

We then compared the results of tooth damage in modern Zambia carnivores to historical samples for all three species obtained from various museums as described in [Mann, Van Valkenburgh & Hayward \(2017\)](#) using chi-square with significance set at $P < 0.050$. Due to methodological differences in assessing tooth wear, i.e., Zambia samples from photographs; historical samples in-hand, we restricted our comparisons of modern with historic specimens to tooth breakage. Further, it should be noted that our Zambia felid sample consists of males, whereas [Mann, Van Valkenburgh & Hayward \(2017\)](#) present data for the sexes combined. In addition, the historical samples span more years (several decades) than our 6-year Zambia study, and thus are not strictly comparable.

Scoring of cranial injuries

We examined skulls for evidence of past injury including healed or healing fractures or other pathologies excluding those directly associated with dental breakage, i.e., missing bone at the alveolar margin of a broken tooth. Damages that were associated with time of death were readily determined, e.g., freshly splintered bone, and excluded from analyses. We initially assessed cranial injuries with skulls in-hand and subsequently quantified damages using digital images.

We scored injuries as resulting from either blunt trauma or intraspecific conflict as follows: (1) major fractures, defined as fractures that caused misalignment of one or more bones sometimes resulting in remodeling, and shallow depressions or gouges with irregular shapes were presumed to have resulted from blunt trauma sustained during prey capture and handling, such as a kick or impact of a horn boss; or (2) deep circular depressions and linear scratches, consistent with bite and claw marks, respectively, were judged to have been caused by intraspecific conflict (Fig. 2). Because some judgements were subjective, it is possible that prey-inflicted injuries vs. intraspecific conflict were

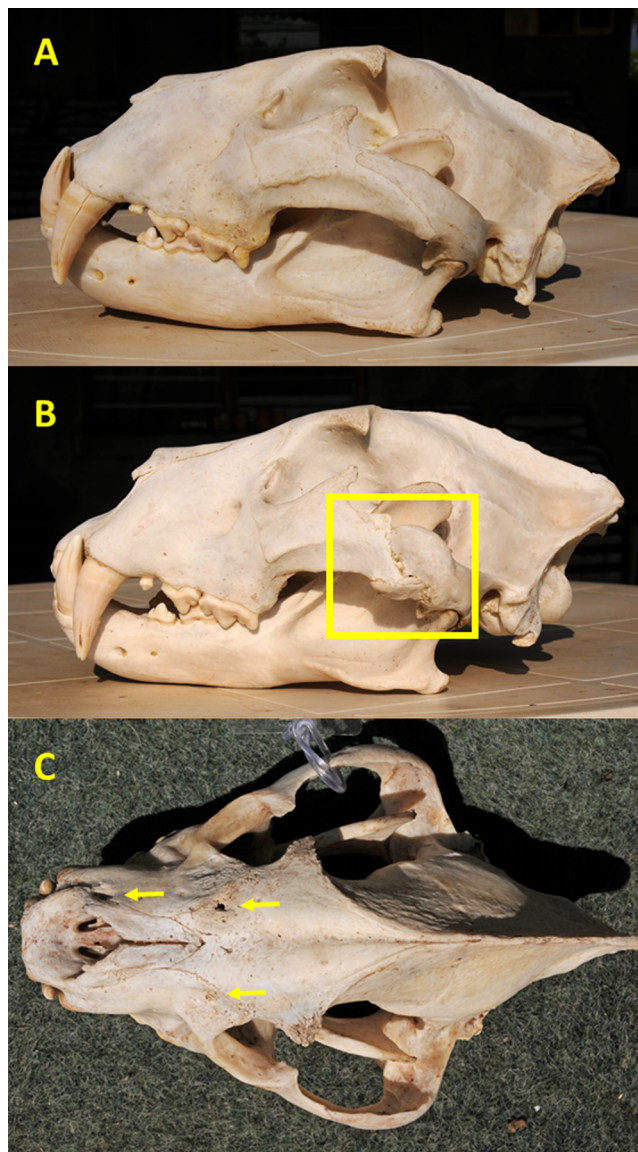


Figure 2 Examples of cranial injuries found in Zambia carnivores. (A) Lion skull with undamaged zygomatic arch. (B) Lion skull with major, partially-healed fracture and remodeled zygomatic arch (shown in square) likely caused by blunt trauma from a large prey. (C) Spotted hyena skull with one puncture and two circular depressions (arrows) from bites to the frontal, maxilla, and premaxilla bones. Photo credits: Paula A. White. [Full-size !\[\]\(5fd6ef84f97f42d7f8b34275f1b65312_img.jpg\) DOI: 10.7717/peerj.11313/fig-2](https://doi.org/10.7717/peerj.11313/fig-2)

erroneously assigned in some instances. Major fractures most commonly involved the zygomatic arches (Fig. 2) or nasals, while other injuries were found primarily on the premaxilla, maxilla, nasal, frontal, and parietal bones.

We recorded the number of incidents of traumas per skull, with an incident defined as occurrence of one type of pathology per animal because it could not be known if multiple injuries were inflicted during a single event. For example, a lion with one or more fractures (which could have occurred during a single predation event) was counted as one incident of blunt trauma attributable to prey-handling, and a lion with one or more

bites and scratches (which could have occurred during a single lion fight) was counted as one incident of intraspecific trauma, whereas a lion with one or more fractures, one bite mark and deep scratches was counted as two incidents (fracture(s) = one blunt trauma; bite and scratches = one intraspecific trauma).

Carnivore populations

From the literature, we obtained lion population size estimates during the sampled time period of 400–750 in LV (0.0064–0.012 lions/km²) with an additional approximately 50 lion inhabiting the CO, and an estimated 250–500 lion in GKE (0.0037–0.0075 lions/km²) (IUCN, 2006). We used IUCN (2006) estimates rather than available focal studies in reporting lion densities because the former represent the larger regions inclusive of the NPs and all adjoining GMAs. Lion appeared to be both more numerous and present at higher density in LV than in GKE (IUCN, 2006).

The LV is considered as having one of the highest densities of leopard in Africa (Gregg, 2019) although few empirical data exist on Zambia's leopard populations (Purchase, Mateke & Purchase, 2007). A study conducted from 2006–2008 in a 491 km² area of the LV reported leopard densities of 3.36 to 4.79 leopards/100 km² (Ray, 2011). Subsequent investigations (2012–2014) in a 313 km² area of the LV estimated from 5.08 to 8.50 leopards/100 km² (Rosenblatt et al., 2016). We found no current publications reporting on leopard density in GKE although there is no evidence to suggest that leopard density in GKE was as high, or higher, than in LV.

Scant data are available on the current status of spotted hyena populations in LV or GKE (Purchase, Mateke & Purchase, 2007); available reports suggest that the species occurs at relatively low density in both regions. Based on camera traps, spotted hyena appeared to be approximately 2–5 times less abundant than leopard in Ray's (2011) LV study area. Playback censuses conducted in 2003–2004 by Carlson, Carlson & Bercovitch (2004) estimated 18–44 spotted hyenas/1,000 km² in northern Kafue National Park (NKNP) and commented that spotted hyena were shy and rarely seen. Likewise, Creel et al. (2018) reported on spotted hyena in NKNP between 2013–2016 as present but not common. This is consistent with reports of spotted hyena occurrence in the larger GKE between 2007–2012 where sightings were relatively infrequent and most often consisted of singles or pairs (White & Kim, 2018). These limited and localized results highlight the need for updated surveys on the status of spotted hyena in LV and GKE (Purchase, Mateke & Purchase, 2007).

Prey populations

From the literature, we obtained relative abundance of prey in each region. Aerial surveys conducted by DNPW in 2015 contributed to population trend data for 12 species of large mammals over the time periods 2002–2015 in LV and 2006–2015 in GKE (DNPW, 2016). Population estimates for 12 key prey species (DNPW, 2016), as well as elephant (IUCN, 2016), and hippopotamus (Chansa & Milanzi, 2010) are provided as Supplemental Information. Because all three carnivore species consume carrion, we augmented estimates of elephant population size with counts of elephant carcasses, where available (CITES

Table 1 Age structure of sampled Zambia carnivores.

Sample	N (%)	Young adult	Mature adult	Old adult
<i>Panthera leo</i> (Luangwa)	60	3 (5)	49 (81.7)	8 (13.3)
<i>Panthera leo</i> (Kafue)	58	6 (10.3)	47 (81)	5 (8.6)
<i>Panthera pardus</i> (Luangwa)	18	3 (16.7)	11 (61.1)	4 (22.2)
<i>Panthera pardus</i> (Kafue)	27	3 (11.1)	22 (81.5)	2 (7.4)
<i>Crocuta crocuta</i> (all Zambia)	13	4 (30.8)	3 (23.1)	6 (46.1)

Secretariat, 2016). Additionally, we referenced *Creel et al. (2018)* for relative abundance of prey species in NKNP from 2013–2016.

RESULTS

Age structure of sampled carnivores

When considering samples from the Luangwa Valley (LV) and Greater Kafue Ecosystem (GKE) combined, the majority of lion (81%) and leopard (73%) were mature adults, i.e., ≥ 5 years old, with 11% of lions and 13% leopards having estimated ages of ≥ 7 years, i.e., old adults (Table 1) (Data S1). There was no significant difference in the age distribution between regions either for lion ($X^2 = 1.701$, $df = 2$, $P = 0.427$) or leopard ($X^2 = 2.639$, $df = 2$, $P = 0.267$). The 13 spotted hyena specimens included young (31%), mature (23%), and old (46%) individuals (Table 1) (Data S1).

Tooth wear and fracture frequency between species

The distribution of individuals among tooth wear categories was similar between lions and leopards, with lions showing a tendency towards having more individuals in the heavy wear class (Fig. 3A) (Data S2) despite the two species having similar age distributions (Table 1). However, lions showed twice the tooth fracture rate of leopards on a per tooth basis (4% vs. 2%; $X^2 = 8.150$, $df = 2$, $P = 0.002$) (Table 2) (Table S1) (Data S2). Although lions also exceeded leopards in the percentage of individuals with one or more broken teeth (53% vs. 44%, Table 2), this difference was not significant ($X^2 = 0.957$, $df = 2$, $P = 0.211$). Across the tooth row, lion had significantly more broken incisors and premolars than leopard ($P < 0.030$) but the two species did not differ greatly in their rates of canine or carnassial tooth fracture (Fig. 3B) (Table S1) (Data S2).

As expected based on their fondness for consuming bone, spotted hyena had more tooth wear and higher tooth fracture rates than lion or leopard. Among the hyenas sampled, there were none that exhibited only slight tooth wear, and the majority were placed in the heavy wear category (Fig. 3A) (Data S2). On a per tooth basis, spotted hyena had a fracture rate of 12%, which is three times that observed for lion (4%, $X^2 = 50.117$, $df = 2$, $P < 0.001$) and six times that observed for leopard (2%, $X^2 = 71.223$, $df = 2$, $P < 0.001$) (Table 2) (Table S1). On a per individual basis, 77% of the spotted hyena had at least one broken tooth, whereas that proportion was 53% for lion and 44% for leopard (Table 2) (Data S2). The higher tooth fracture observed in spotted hyena was distributed across the

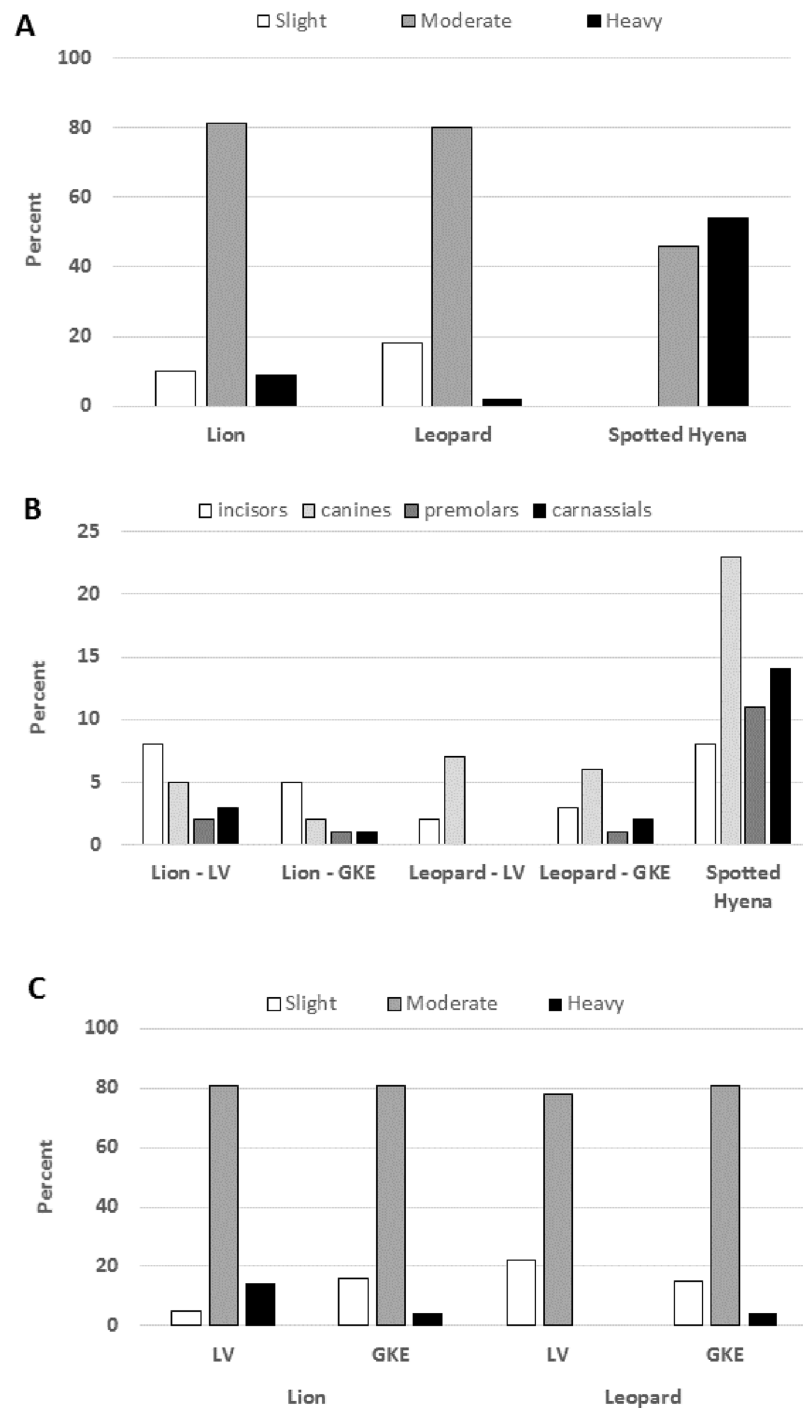


Figure 3 Tooth wear stage and tooth fracture by position in Lion, Leopard and Spotted Hyena. (A) Percent of individuals assigned to each tooth wear stage in the three species sampled in Zambia. Total sample size is 115 lions, 45 leopards and 13 spotted hyena. (B) Percent of teeth broken for each tooth position within each sample by region, Luangwa Valley (LV), Greater Kafue Ecosystem (GKE). Sample sizes for spotted hyenas were too small to compare regions. For tooth sample size, see Table S1. (C) Percent of individuals assigned to each tooth wear stage within each region. Total sample size is 58 (Lion-LV), 57 (Lion-GKE), 18 (Leopard-LV), and 27 (Leopard-GKE).

Full-size DOI: 10.7717/peerj.11313/fig-3

Table 2 Tooth fracture in Zambia carnivores from two regions and comparisons with historical samples.

Sample	N (skulls)	N (teeth)	% individuals w/≥1 brkn tooth	% broken teeth	% broken teeth w/canine teeth excluded
<i>Panthera leo</i> (Mann, Van Valkenburgh & Hayward, 2017)	133	3,491	33.8	3.5	2.8
<i>Panthera leo</i> (all Zambia)	115	3,286	53	3.8	4
<i>Panthera pardus</i> (Mann, Van Valkenburgh & Hayward, 2017)	146	3,697	39	3.7	2.4
<i>Panthera pardus</i> (all Zambia)	45	1,313	44.4	2.1	1.5
<i>Crocuta crocuta</i> (Mann, Van Valkenburgh & Hayward, 2017)	179	5,084	51	4.3	3.5
<i>Crocuta crocuta</i> (all Zambia)	13	402	76.9	11.7	10
Within Zambia					
<i>Panthera leo</i> (Luangwa)	58	1,658	62.1	5.0	5.0
<i>Panthera leo</i> (Kafue)	57	1,628	43.9	2.6	2.6
<i>Panthera pardus</i> (Luangwa)	18	517	38.9	1.9	1.0
<i>Panthera pardus</i> (Kafue)	27	796	48.1	2.3	1.5

tooth row, with canine teeth and cheek teeth being more likely to be broken than incisors (Fig. 3B).

Tooth fracture frequency within species

Sample sizes of lion and leopard were sufficient to explore differences in tooth fracture frequency between populations in LV and GKE. Among the lions, the differences between the two regions were striking with LV lion exhibiting significantly higher fracture frequencies than their GKE counterparts, both in percentage of broken teeth (5% vs. 2.6%, $X^2 = 13.213$, $df = 2$, $P < 0.001$) and percentage of individuals with one of more broken teeth (62.1% vs. 43.9%; $X^2 = 3.827$, $df = 2$, $P = 0.038$) (Table 2) (Data S2). Notably, the higher fracture frequency of LV lions was apparent across the entire tooth row, from incisors to carnassials (Fig. 3B) (Table S1). In association with their higher tooth fracture rates, LV lions also had more rapid rates of tooth wear than GKE lions, as evidenced by a smaller proportion of individuals in the slight wear class along with a larger proportion of individuals in the heavy wear class (Fig. 3C; $X^2 = 6.623$, $df = 2$, $P = 0.036$), despite the fact that the age distributions of lions did not differ between LV and GKE (Table 1). Unlike the lions, the leopards from the two regions did not differ significantly in fracture frequency on either a per tooth ($X^2 = 0.161$, $df = 2$, $P = 0.424$) or per individual basis ($X^2 = 0.375$, $df = 2$, $P = 0.381$) (Fig. 3C) (Table S1) (Data S2).

Comparison with historical samples

Tooth fracture rates on a per tooth basis in male lion and leopard from Zambia were largely comparable to those observed in historical samples of both sexes of these species that originated from different parts of Africa as published in Mann, Van Valkenburgh & Hayward (2017) (Table 2). There were significant differences between the historical and Zambia samples in a few cases. Among lion, the proportion of individuals with at least one broken tooth was significantly greater in the Zambia lion (53% vs. 33.8%; $X^2 = 4.322$, $df = 2$, $P = 0.025$) as was the proportion of teeth fractured in LV lion (5% vs. 3.5%;

Table 3 Frequency of individuals with at least one injury ascribed to either blunt trauma or intraspecific conflict in lion and leopard by region and in spotted hyena (both regions).

Sample	N (%)	Blunt trauma major fractures ¹	Intraspecific bite wounds, scratches	Non-specific	Total incidents
<i>Panthera leo</i> (Luangwa)	60	31 (51.7) 15 (25)	19 (31.7)	0	50 (83.3)
<i>Panthera leo</i> (Kafue)	56	17 (30.4) 5 (8.9)	17 (30.4)	0	34 (60.7)
<i>Panthera pardus</i> (Luangwa)	18	4 (22.2)	8 (44.4)	1 (5.6)	13 (72.2)
<i>Panthera pardus</i> (Kafue)	27	5 (18.5)	11 (40.7)	5 (18.5)	21 (77.8)
<i>Crocuta crocuta</i>	13	N/A	N/A	4 (30.8)	4 (30.8)

Note:

¹ Subset of the 31 (LV) and 17 (GKE) blunt traumas in lions that consisted of major fractures.

$X^2 = 6.717$, $df = 2$, $P = 0.007$) but not GKE lion. The per tooth fracture rate for Zambia leopard was significantly less than that recorded by [Mann, Van Valkenburgh & Hayward \(2017\)](#) (2.1% vs. 3.7%; $X^2 = 8.641$, $df = 2$, $P = 0.003$). The most striking difference is in the spotted hyena. Our small sample (13) collected over 6 years in Zambia broke their teeth much more frequently on both a per tooth basis (12% vs. 4%; $X^2 = 44.816$, $df = 2$, $P = 0$) and per individual basis (77% vs. 51%; $X^2 = 694.932$, $df = 2$, $P = 0$) than the spotted hyena sample in [Mann, Van Valkenburgh & Hayward \(2017\)](#) that included 179 skulls spanning decades and multiple countries.

Cranial injuries

Over half (68/116; 59%) of all our sampled lions had at least one cranial injury ascribed to natural causes, and 16 individual lions (14%) showed evidence of both blunt trauma and intraspecific conflict ([Data S3](#)). LV lion had significantly higher rates of blunt trauma injuries attributable to prey-handling compared to lion in GKE (52% vs. 30%) ($X^2 = 5.422$, $df = 2$, $P = 0.019$) ([Table 3](#)). Among the 31 LV lions with blunt trauma injuries, 15 consisted of major healed or healing fractures ([Fig. 2](#)) as opposed to only five lions with major injuries in GKE. In contrast, the incidence of injuries ascribed to intraspecific conflict was similar for lion from LV and GKE (32% vs. 30%) ([Table 3](#)).

Among leopard, the number of incidents attributable to blunt trauma and intraspecific conflict combined were similar between regions (LV 72%; GKE 78%) ([Table 3](#)) ([Data S3](#)). No major fractures were seen in leopards. Due to small sample sizes and the number of incidents that could not be assigned to specific categories, differences in the frequency of blunt trauma injuries related to prey-handling vs. intraspecific conflict in LV vs. GKE leopards were not conclusive ([Table 3](#)).

Of the 13 spotted hyena skulls examined, four exhibited signs of significant natural trauma, including one with multiple penetrating bite wounds across its rostrum which may have been inflicted by a lion or a hyena ([Fig. 2](#)). Because the likely causes of the natural injuries in spotted hyena (prey-handling vs. intraspecific conflict vs. lion) could not be determined to a reasonable degree, all were classified as non-specific ([Table 3](#)) ([Data S3](#)).

Prey diversity, abundance and trends

Prey species diversity was similar in both regions with a few exceptions; a small population of endemic Thornicroft's giraffe occurs only in LV, and lechwe *Kobus lechwe* occur only in GKE. Relative abundance and density of prey varied between regions and among species (DNPW, 2016) (Data S4). Buffalo (DNPW, 2016), elephant (IUCN, 2016), and hippopotamus (Chansa & Milanzi, 2010) occurred in much higher abundance and density in LV compared with GKE. Data on elephant carcasses were available only from a specified survey area in LV where from 4 to 49 carcasses per year (mean of 19 carcasses/year) were reported from 2002–2012 (CITES Secretariat, 2016) (Data S5). Hippopotamus surveys conducted between 2005–2008 found that LV contained the largest concentration in Zambia, with the Luangwa river accounting for 62% (25,000) of the country's overall hippopotamus population compared to 10% (4,000) in the Kafue river in GKE (Chansa & Milanzi, 2010).

Higher numbers of several species of small to medium-sized antelopes (e.g., lechwe, puku *K. vardonii*, waterbuck *K. ellipsiprymnus*, sable *Hippotragus niger*) were recorded in GKE (Data S4). Impala *Aepyceros melampus* were the most abundant antelope by far in both systems, although density of puku and sable were also high in GKE. Lechwe populations in GKE were highly localized in the northwest corner of the ecosystem (DNPW, 2016). A slightly different picture of prey composition in Kafue is given by Creel *et al.* (2018) who reported the species of highest densities in northern Kafue National Park (NKNP) as puku, impala, and warthog (*Phacochoerus aethiopicus*). Aerial surveys indicated that for the majority of key wildlife species in GKE, wildlife numbers that had generally declined from the mid to late-1990s increased over the period 2006–2012. Similarly, over the period 2002–2012 in the LV, numbers of most wildlife species were reported to be stable or increasing (DNPW, 2016) (Data S4).

DISCUSSION

Wild carnivores experience increased levels of natural tooth wear (Bodecker, 1925; Van Horn, McElhinny & Holeykamp, 2003; Curtis *et al.*, 2018) and tooth breakage with age (Van Valkenburgh, 1988, 2009). Despite the fact that the age distribution of lions was similar in Luangwa Valley (LV) and Greater Kafue Ecosystem (GKE), we found significantly higher rates of tooth wear and fracture in LV lions. Because the methods of assessing tooth wear differed (photographs vs. specimens in-hand), we were unable to compare tooth wear between modern and historic carnivores. However, the fact that rates of tooth breakage in historic samples of African lion were similar to modern lion in GKE affirms that actual differences in tooth fracture rates exist between modern carnivores inhabiting Zambia's two regions.

A higher proportion of LV lions showed moderate/heavy tooth wear, and the percentage of lions with more than one broken tooth was 18% greater compared with lions from GKE. Tooth breakage in LV lions was higher across all tooth types, but this was more pronounced at the canines (5% in LV vs. 2% in GKE) and incisors (8% in LV vs. 5% in GKE). Previous observations of carnivore feeding behavior documented associations between tooth type employed and specific tissue type being consumed and found that

incisors and canines (to a lesser degree) are used to pull muscle and skin from a carcass whereas posterior teeth, i.e., carnassials, are more often used when consuming skin, or muscle plus bone (Van Valkenburgh, 1996). Tooth breakage rates increase when carcass utilization is higher (Van Valkenburgh et al., 2019). Higher breakage rates among all tooth types in LV lions suggests that they may have been consuming prey more completely and consequently consuming more bone than lions in GKE.

Greater carcass utilization and thus increased tooth wear and breakage has been correlated with declines in the ratio of numbers of prey (elk (*Cervus canadensis*) or moose (*Alces alces*)) to predators (gray wolves) in both North American and Scandinavian ecosystems (Van Valkenburgh et al., 2019). Our study lacked sufficient empirical data to thoroughly examine predator/prey ratios that may have differentially influenced tooth wear and breakage. However, limited prey availability does not appear to explain the higher rates of tooth fracture in LV vs. GKE lions.

A more plausible explanation is that tooth wear and breakage varied as a consequence of the prey species being targeted. Cape buffalo, a preferred prey for lion (Hayward & Kerley, 2005), occurred at far greater density in LV than in GKE, as did elephant and hippopotamus (Chansa & Milanzi, 2010; DNPW, 2016). Moreover, Creel et al. (2018) reported that buffalo recently (2013–2016) made up only 10% of the diet of lions in northern Kafue National Park (NKNP) (significantly less than the 30% reported in GKE by Mitchell, Shenton & Uys (1965) five decades earlier) and that lion diet in NKNP now consisted primarily of small to medium-sized herbivores. This is in stark contrast to lion in LV where 60% of the diet is comprised of large-bodied species, i.e., buffalo, hippopotamus, elephant, and giraffe (Zambian Carnivore Programme unpublished, in Creel et al., 2018). Buffalo are particularly challenging prey with extremely tough skin >1 cm thick (Schaller, 1972) and are among the most dangerous quarry for lion to hunt and kill (Hayward & Kerley, 2005). Canine teeth are critical to the killing bite, and are prone to fracture if subjected to bending loads such as might occur when teeth contact bone (Van Valkenburgh & Ruff, 1987). The largest sharpest cheek teeth (carnassials) are likely critical for opening carcasses (Van Valkenburgh, 1996). Consumption of more tough-skinned, large-boned prey likely contributed to the higher overall rate of tooth wear and breakage found in LV lions.

Additional support for the concept that LV lions are tackling larger, more perilous prey comes from our cranial injury data. More than half (52%) of the LV lions sampled had suffered blunt trauma to their skull, including 15 lions with major skull fractures that likely resulted from impacts with the hooves or horns of their prey. The combination of elevated canine tooth and skull fracture rates in LV lions may be a result of killing and scavenging large prey. Alternatively or in addition, the higher rates of canine tooth breakage in LV vs. GKE lions may be due to higher lion density (IUCN, 2006) that could have resulted in increased intraspecific conflict (and breakage of canines during fights) in the former. However, lions in both regions exhibit similar rates of cranial bite and scratch mark injuries suggesting no significant difference in levels of lion-lion conflict among regions.

Among leopards, tooth wear and breakage rates showed slight variation by tooth type in LV and GKE, but no significant difference between regions. This finding suggests that prey species of comparable size are taken by leopard in both regions, and is consistent with recent reports of small to medium-sized herbivores representing primary prey of leopards both in NKNP ([Creel et al., 2018](#)) and LV ([Ray, 2011](#)).

In contrast to lions, the incidence of cranial injuries in leopards occurred at roughly the same frequency in LV and GKE. No major cranial fractures were seen among leopards. The lack of empirical data on leopard population sizes in either region, in addition to the number of injuries that could not be confidently assigned to a specific cause precluded further analyses of cranial injuries in leopards.

Consistent with prior work ([Van Valkenburgh, 2009](#)), interspecific differences in rates of tooth fracture corresponded to levels of bone consumption; spotted hyena exhibited the most dental damage, followed by lion and leopard. Notably, the Zambia hyenas exhibited the highest rate of tooth fracture recorded for this species, more than doubling previous estimates from museum specimens. We have yet to find an explanation for these high rates of wear and fracture. Spotted hyena are known to hunt and scavenge a wide variety of prey ([Hayward, 2006](#)), although scavenging of large numbers of elephant carcasses might have been an important factor during this study. In GKE, spotted hyenas returned to chew on the bones of an elephant carcass 35 days after its death ([White & Diedrich, 2012](#)). In Botswana, spotted hyena were found to scavenge on an elephant carcass for as long as 50 days ([Cozzi et al., 2015](#)). Availability of greater numbers of elephant carcasses due to natural mortality, deaths resulting from human-wildlife conflict, and poaching may have resulted in high rates of dental attrition due to the difficulty of slicing elephant hide and/or consumption of larger quantities of bone by hyena, and by lion in LV relative to GKE, although we need comparative data on elephant carcass availability from GKE to draw this conclusion.

Comparisons of our data on tooth fracture in extant populations of lion, leopard, and spotted hyena with previously published data for historical samples yielded mixed results, and should be viewed with some caveats. The historical samples include individuals from multiple locations in Africa and, within a species, tooth wear can vary with location due to substrate (e.g., lions; [Smuts, Anderson & Austin, 1978](#)). Moreover, [Mann, Van Valkenburgh & Hayward's \(2017\)](#) historic samples include individuals collected over a span of decades, and samples that span longer time intervals are likely to produce lower rates of tooth fracture than those that span shorter intervals because tooth fracture rates can vary over time depending on resource availability and longer intervals are likely to include periods of minimal tooth fracture ([Mann, Van Valkenburgh & Hayward, 2017](#)). Consequently, we might have expected that our Zambia samples that span six years would produce higher estimates of tooth fracture frequency than those presented in [Mann, Van Valkenburgh & Hayward \(2017\)](#) collected between 1889 and 1967.

Nevertheless, in most cases, our tooth fracture rates did not differ greatly from those reported in [Mann, Van Valkenburgh & Hayward \(2017\)](#), with two exceptions. Lions from LV had much higher rates of tooth fracture per individual (62% vs. 34%) and somewhat higher per tooth (5% vs. 3.5%) than the historical sample means. The difference at the

individual level is especially striking and suggests that LV lions are unusual in showing such a high prevalence of broken teeth. The difference might reflect the fact that we sampled only males whereas [Mann, Van Valkenburgh & Hayward \(2017\)](#) included both sexes, but this seems an unlikely explanation given that our sample of GKE males had rates of fracture (44%) similar to those of the historical samples. The second exception is the extremely high rates of tooth wear and fracture in our sample of 13 Zambia spotted hyena relative to historical samples (12% vs. 4% on a per tooth basis, and 77% vs. 51% on an individual basis). Examination of a larger sample of skulls from LV and GKE would be of interest to see if the results from this study are truly representative of hyenas from these two regions in Zambia, or if other explanations are needed for the extreme tooth wear and breakage recorded here.

In our introduction, we suggested that studies of cranial and dental injuries in large carnivores could enhance our understanding of the forces that mold their morphology and behavior, beyond what can be seen from field observations of living individuals. Over half of the lions we sampled had experienced one or more cranial injuries which would not have been easily detected in vivo. This substantial incidence of injury suggests that their skulls are certainly not overbuilt for expected loads and may represent a compromise between maximizing strength and limiting mass. Similarly, the prevalence of broken teeth indicates that the evolution of tooth structure, shape, and size is the result of multiple, sometimes opposing demands (e.g., sharpness vs. fracture resistance). Beyond providing us with insights into the long term selective forces that shaped these carnivores, rates of tooth wear and fracture frequency can give us information about current conditions that impact individual fitness. For example, tooth wear and fracture likely increase when food becomes limiting, and bone and carcass utilization rise. In Zambia, the very high levels of dental attrition and breakage in the spotted hyena suggest that these populations may somehow be food limited. Lions of LV suffer more dental fracture and more severe cranial injuries than those of the GKE, which we propose may be due to hunting larger, more dangerous prey, and/or greater carcass utilization. Resolving this would require more detailed data on prey choice and feeding behavior of carnivores in the two regions.

Our findings of injuries in LV lions raise the question: why risk hunting large prey when smaller prey are available? Buffalo occur at much higher abundance in LV than GKE, and the preference of lions for hunting buffalo ([Mitchell, Shenton & Uys, 1965](#); [Schaller, 1972](#)) and other large game is well-documented ([Hayward & Kerley, 2005](#)). This preference is especially pronounced during the dry season when lack of cover likely contributes to a lowered success at stalking the faster, smaller ungulates ([Mitchell, Shenton & Uys, 1965](#); [Hayward & Kerley, 2005](#)). Although hunting large prey carries heightened risks ([Hayward & Kerley, 2005](#)), the benefit is procurement of a substantial amount of meat and is consistent with optimal foraging theory based on energetic return for effort ([Pyke, Pulliam & Charnov, 1977](#)). In addition, cub growth is influenced greatly by nutrition ([Smuts, Robinson & Whyte, 1980](#)), and cubs are more likely to obtain meat from large vs. small kills ([Schaller, 1972](#)). Thus, from an evolutionary standpoint, the

increased risk to a lion of sustaining a cranial injury while securing large-bodied prey is offset by enhanced cub survival as well as energetic return.

CONCLUSIONS

Our survey of craniodental injuries of trophy hunted lion, leopard, and spotted hyena has revealed some intriguing differences among and within species, and underscores the value of utilizing trophy specimens in empirical research. Undoubtedly, our ability to interpret the differences would be enhanced by additional surveys of these species in other localities, especially those with metadata on predator and prey numbers and carnivore feeding preferences. Still, our low-cost, low-tech survey captured significant information from a sample of trophy hunted individuals, a study group that remains an underutilized resource for scientific investigations.

We recommend that collection and archiving of standardized photographs of carnivore teeth and skulls be incorporated into existing trophy monitoring programs to allow for more comprehensive investigations on naturally-occurring injuries and dietary stress in carnivore populations, including comparisons across different regions and time periods. Additional data on the health and condition of wild carnivores are always welcome, and relatively simple surveys of craniodental trauma are an easy add-on to the conservation biology tool box.

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The authors declare that they have no competing interests.

Author Contributions

- Blaire Van Valkenburgh conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Paula A. White conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.

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Data Availability

The following information was supplied regarding data availability:

The raw data are available as [Supplemental Files](#).

Supplemental Information

Supplemental information for this article can be found online at <http://dx.doi.org/10.7717/peerj.11313#supplemental-information>.

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