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Tooth wear in captive rhinoceroses (*Diceros*, *Rhinoceros*, *Ceratotherium*: *Perissodactyla*) differs from that of free-ranging conspecifics

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Tooth wear in captive rhinoceroses (*Diceros*, *Rhinoceros*, *Ceratotherium*: Perissodactyla) differs from that of free-ranging conspecifics

Lucy A. Taylor^{1,2}, Dennis W.H. Müller^{3,4}, Christoph Schwitzer¹, Thomas M. Kaiser⁵, Daryl Codron^{3,6}, Ellen Schulz⁵, Marcus Clauss^{3,7}

¹ Bristol Zoological Society, c/o Bristol Zoo Gardens, Clifton, Bristol BS8 3HA, UK

² School of Biological Sciences, University of Bristol, Woodland Road, Bristol BS8 1UG, UK

³ Clinic for Zoo Animals, Exotic Pets and Wildlife, Vetsuisse Faculty, University of Zurich, Winterthurerstr. 260, 8057 Zurich, Switzerland

⁴ National Park 'Bavarian Forest', Freyunger Str. 2, 94481 Grafenau, Germany

⁵ Biocenter Grindel and Zoological Museum, University of Hamburg, Martin-Luther-King-Platz 3, 20146 Hamburg, Germany

⁶ Florisbad Quaternary Research Department, National Museum, Po Box 266, Bloemfontein, 9301, South Africa

⁷ E-mail: mclauss@vetclinics.uzh.ch

Key words: Rhinoceros, mesowear, tooth wear, captivity, nutrition, browser, grazer

Abstract

Tooth wear can affect body condition, reproductive success and life expectancy. Poor dental health is frequently reported in the zoo literature, and abrasion-dominated tooth wear, which is typical for grazers, has been reported in captive browsing ruminants. The aim of this study was to test if a similar effect is evident in captive rhinoceros species. Dental casts of maxillary cheek teeth of museum specimens of captive black (*Diceros bicornis*; browser), greater one-horned (*Rhinoceros unicornis*; intermediate feeder) and white rhinoceroses (*Ceratotherium simum*; grazer) were analysed using the recently developed extended mesowear method for rhinoceroses. Captive *D. bicornis* exhibited significantly more abrasion-dominated tooth wear than their free-ranging conspecifics ($p < 0.001$), whereas captive *C. simum* exhibited significantly less abrasion-dominated tooth wear, particularly in the posterior cusp of the second molar ($p = 0.005$). In *R. unicornis*, fewer differences were exhibited between free-ranging and captive animals, but tooth wear was highly variable in this species. In both free-ranging and captive *D. bicornis*, anterior cusps were significantly more abrasion-dominated than posterior cusps ($p < 0.05$), which indicates morphological differences between cusps that may represent functional adaptations. By contrast, tooth wear gradients between free-ranging and captive animals differed, which indicates ingesta-specific influences responsible for inter-tooth wear differences. Captive *D. bicornis* exhibited more homogenous tooth wear than their free-ranging conspecifics, which may be caused by an increase in the absolute dietary abrasiveness and a decrease in relative environmental abrasiveness compared to their free-ranging conspecifics. The opposite occurred in *C. simum*. The results of this study suggest that diets fed to captive browsers are too abrasive, which could result in the premature loss of tooth functionality, leading to reduced food acquisition and processing ability and, consequently, malnourishment.

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Introduction

Tooth wear experienced by free-ranging ungulates corresponds to feeding type. Browsers have attrition-dominated tooth wear, characterised by an occlusal relief (OR) that is high and a cusp shape (CS) that is sharp, whereas grazers have abrasion-dominated tooth wear, characterised by low OR and blunt CS (Fortelius and Solounias, 2000). To retain tooth functionality on high abrasion diets, grazers have evolved high-crowned (hypsodont) teeth, which are vertically elongated, enabling the tooth to wear for longer (Janis, 1988). Thus,

browsers, which generally have low-crowned teeth, may not be adapted to high abrasion diets (Damuth and Janis, 2011). Excessive tooth wear has been found to influence body condition, reproductive success and longevity in many species (Skogland, 1988; Ozaki *et al.*, 2010), which is attributed to reduced food acquisition and processing ability, leading to negative energy balance (Kojola *et al.*, 1998). In captive wildlife, poor dental health is frequently reported (Martin Jurado *et al.*, 2008), and abrasion-dominated tooth wear has been documented in captive browsing ruminants, such as giraffe (*Giraffa camelopardalis* Linnaeus, 1758) (Clauss *et al.*, 2007). Captive browsing ruminants have a significantly shorter relative life expectancy than mixed feeders and grazers, which suggests there are problems providing appropriate food in captivity (Müller *et al.*, 2011). Conversely, grazing ruminants, such as American bison (*Bison bison* Linnaeus, 1758), experience less abrasion-dominated tooth wear in captivity, and age at a slower rate (Kaiser *et al.*, 2009; Lemaître *et al.*, 2013). Therefore, unnatural tooth wear could be affecting captive animal health, welfare and longevity and, consequently, conservation strategies for captive ungulates.

The Rhinocerotidae, comprising all five extant rhinoceros species, are one of the most threatened families of mammals on Earth. Two monotypic genera and three species of rhinoceros are classified as Critically Endangered on the IUCN Red List, one as Vulnerable and one as Near Threatened. In addition to *in situ* conservation efforts in increasingly small and isolated habitat fragments, conservation breeding programmes in zoos, which exist for three of the five rhinoceros species, are a vital tool for ensuring their survival. Yet, differences in tooth wear between free-ranging and captive rhinoceroses have not been quantified, and their potential impact on the longevity, and hence lifetime reproductive output, is unknown.

Analyses of tooth wear, using the mesowear method, have been used extensively in dietary and habitat reconstruction (Fortelius and Solounias, 2000) and to compare the tooth wear patterns experienced by free-ranging and captive ruminants (Clauss *et al.*, 2007; Kaiser *et al.*, 2009). Recently, an extended mesowear method was developed by Winkler and Kaiser (2011) and adapted for rhinoceroses, facilitating the detection of inter-cusp and inter-tooth wear pattern differences (Taylor *et al.*, 2013). In free-ranging black rhinoceros (*Diceros bicornis* Linnaeus, 1758; browser), the anterior cusps were consistently more abrasion-dominated than the posterior cusps, which may reflect a morphological adaptation to browse. In addition, both *D. bicornis* and the greater one-horned

rhinoceros (*Rhinoceros unicornis* Linnaeus, 1758; mixed-feeder) had a significant tooth wear gradient, with teeth becoming less abrasion-dominated along the tooth row. Taylor *et al.* (2013) suggested that tooth wear gradients may develop due to relative changes in abrasiveness of the ingesta within the oral cavity; in particular, environmental (external) abrasives may affect the anterior teeth more severely before bolus formation. By contrast, high absolute amounts of ingested abrasives may override other signals, leading to homogenous tooth wear in white rhinoceros (*Ceratotherium simum* Gray, 1868; grazer). Comparing the tooth wear experienced by free-ranging and captive rhinoceroses could differentiate factors caused by morphology and ingesta-specific influences, because morphological constraints will remain consistent between the wild and captivity whereas ingesta-specific influences may vary in a systematic way. Therefore, the aim of this study was to quantify the degree of tooth wear observed in free-ranging and captive *D. bicornis*, *R. unicornis* and *C. simum* in order to establish whether different wear patterns are experienced in captivity.

Material and methods

Materials

We investigated 49 museum specimens of *Diceros bicornis*, *Rhinoceros unicornis* and *Ceratotherium simum* from 19 zoological museums and collections (Online Appendix S1). *C. simum* was treated as one species, instead of dividing it into *C. simum* and *C. cottoni* as recently suggested (Groves *et al.*, 2010) due to the small sample size for this genus. Measurements on living specimens were not possible due to the invasiveness of the method. Only specimens with known origin from captivity were selected. Dental casts were produced of either the left or the right maxillary (upper) tooth row. A negative mould was made with Provil novo Light C.D. 2 fast set EN ISO 4823, type 3, light and Provil novo Putty regular set EN 24823 (Heraeus Kulzer GmbH, Hanau, Germany) polysiloxane dental moulding putty. One-to-one positive casts were produced by filling the moulds with epoxy resin Injektionsharz EP (Reckli-Chemiewerkstoff, Herne, Germany).

Mesowear scoring

The permanent second, third and fourth premolar (P2, P3 and P4) and the first and second molar (M1 and M2) were analysed using the adjusted rhinoceros mesowear

method (Taylor *et al.*, 2013). Occlusal relief (OR) and cusp shape (CS) were scored separately for the anterior and posterior cusp position due to the asymmetry of rhinoceros teeth. Rhinoceros-adjusted OR categories of 'high-high' (height/length ≥ 0.125), 'high' ($< 0.125 - 0.083$), 'high-low' ($< 0.083 - 0.0417$), 'low' ($< 0.0417 > 0.00$) and 'flat-negative' (≤ 0), were used. CS were scored as 'sharp', 'round-sharp', 'round', 'round-round' or 'blunt' according to the degree of facet development. In addition, the posterior M2 was scored using the 'classical' mesowear method of Fortelius and Solounias (2000), with rhinoceros adjusted OR, for comparisons with previous studies (with OR scored as 'high' (≥ 0.03) or 'low' (< 0.03) and CS as 'sharp', 'round' or 'blunt'). Digital callipers and a triplet hand lens (10x-18mm) were used, when required, to differentiate OR and CS.

As ontogeny can affect mesowear (Rivals *et al.*, 2007), and specimen ages were unknown, the wear stage chart of Taylor *et al.* (2013) was used to ensure all specimens were in the same dental functional stage, excluding young and old specimens. Wear stage 6 was considered for *D. bicornis* and stages 6-7 for *R. unicornis* and *C. simum*. All pathological teeth were excluded.

Both mesowear methods were converted into a mesowear score for analysis. The extended mesowear OR and CS were converted into scores from 0 'high-high'/'sharp' up to 4 'flat-negative'/'blunt'. A mean of OR and CS was then calculated as the mesowear score (Taylor *et al.*, 2013). 'Classical' mesowear results were converted into a combination score of 0 'high' and 'sharp', 1 'high' and 'round', 2 'low' and 'round', 3 'low' and 'sharp' and 4 'low' and 'blunt' (Kaiser *et al.*, 2009).

Statistical analyses

Data from this study were compared to data on free-ranging specimens from Taylor *et al.* (2013). Species differences within each group (free-ranging or captive) were tested using Kruskal-Wallis tests on the posterior M2 for OR, CS and mesowear score.

Overall differences between free-ranging and captive specimens in each species were tested using linear mixed-effects (LME) models with fixed effects of Origin * Tooth position (Cusp position) and random effect of Individual specimen for the premolars (P2-P4) and molars (M1-M2) separately due to tooth wear gradients. Due to broken cusp and wear stage exclusions, sample sizes were unequal. Balanced subsamples were taken separately for the anterior and posterior cusp within each tooth to minimise loss of data. Totally balanced subsamples were tested, but resulted in substan-

tial data loss (*R. unicornis* $n=5$) without changing the results. Random subsamples were taken using the statistical program R version 2.15.2 (R Core Team, 2012). We used restricted maximum likelihood (REML) and calculated denominator degrees of freedom using Satterthwaite's approximation. Differences between free-ranging and captive specimens for each individual cusp were tested on the full dataset using Welch's two sample *t*-tests, which does not assume equal variances and sample sizes (Ruxton, 2006).

Intra-species differences along the tooth row were tested using LME models with fixed effects of Cusp position and Tooth position (nested in Cusp position) and random effect of Individual specimen for the CS, OR and mesowear score for each species and origin using the subsample data. Tooth position differences were subsequently tested separately in the anterior and posterior cusp tooth row using one-way analyses of variance (ANOVA) and Dunnett's T3 post hoc test.

Statistical analyses were conducted in IBM® SPSS® Statistics 19 software (SPSS Inc., Chicago, IL) and R version 2.15.2 (R Core Team, 2012). The significance level was set to $p < 0.05$.

Results

Species-specific wear signatures

Occlusal relief (OR) and mesowear scores in the posterior M2 differed significantly between the three species within both the free-ranging (Kruskal-Wallis: $p < 0.001$) and captive animals ($p < 0.01$). However, in contrast to the free-ranging animals, no significant difference was detected in cusp shape (CS) between captive *D. bicornis*, *R. unicornis* and *C. simum* in the posterior M2 (free-ranging: $p < 0.001$; captive: $p = 0.140$).

Tooth wear in free-ranging and captive animals

Overall, the CS, OR and mesowear scores in both premolars and molars were significantly more abrasion-dominated in captive *D. bicornis* (LME: all $p < 0.001$), and significantly less-abrasion dominated in captive *C. simum* (all $p < 0.05$), compared to their respective free-ranging conspecifics (Table 1). In *R. unicornis*, no differences were detected in molar CS ($p = 0.134$), but overall mesowear scores of captive *R. unicornis* were more abrasion-dominated than their free-ranging conspecifics (all $p < 0.05$). There was a subjective impression that captive specimens of all species had more

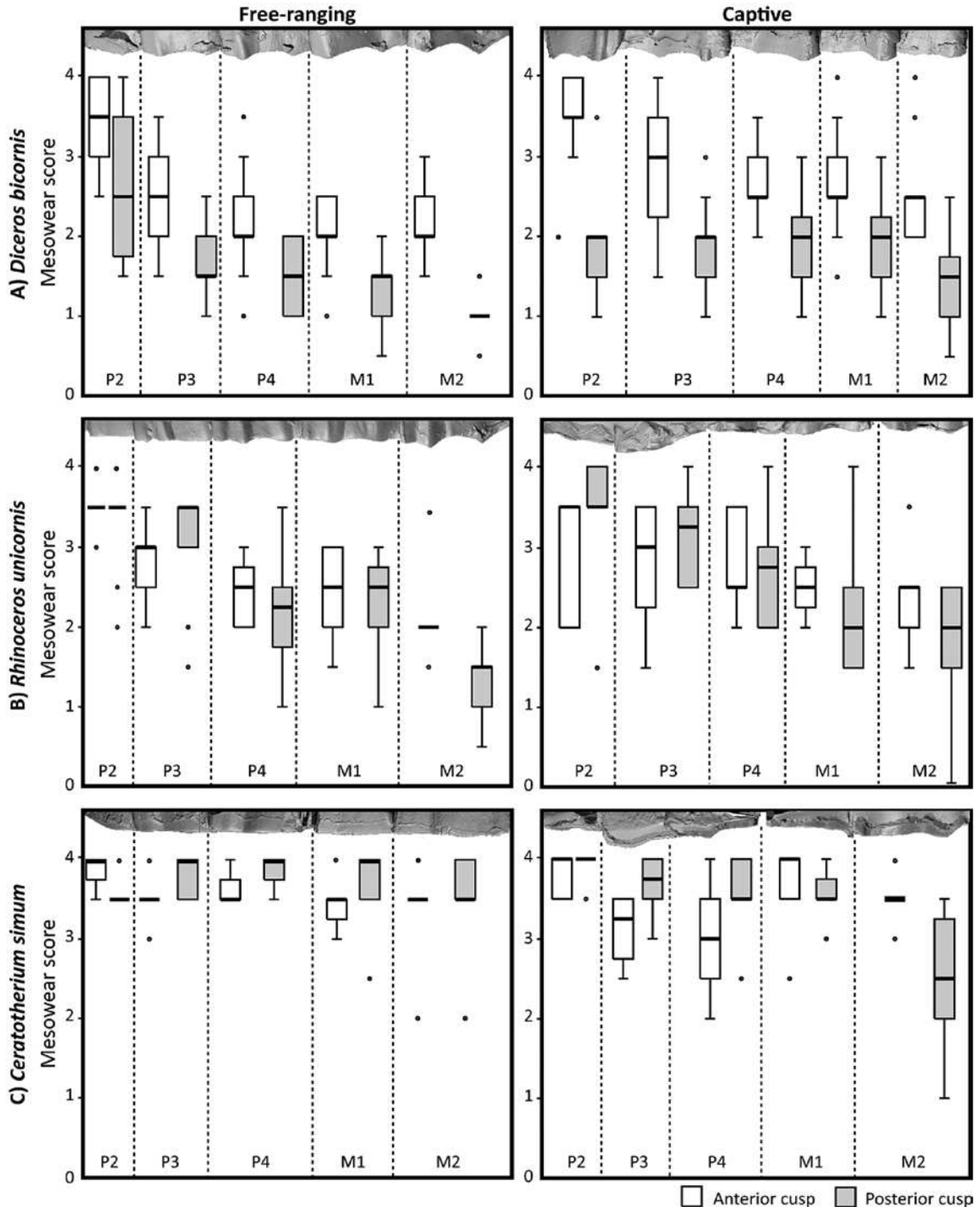


Fig. 1. Comparison of the mesowear scores for each tooth and cusp position of free-ranging and captive (a) *Diceros bicornis*, (b) *Rhinoceros unicornis* and (c) *Ceratotherium simum*. Boxplots denote the median (middle line), the upper and lower quartile (box limits), the outside interquartile range (IQR, whiskers), and outliers (defined as 1.5 times above or below the IQR). 3D scans of typical maxillary tooth rows from free-ranging and captive *D. bicornis*, *R. unicornis* and *C. simum* based on the similarity of their mesowear scores to the sample mean (Specimen identification: L-R: NMB-1021034, NMP-25963, AMNH-54455, ZMH-83585, NHM-752384, NHM-2_03). Note less even tooth wear in the captive specimens.

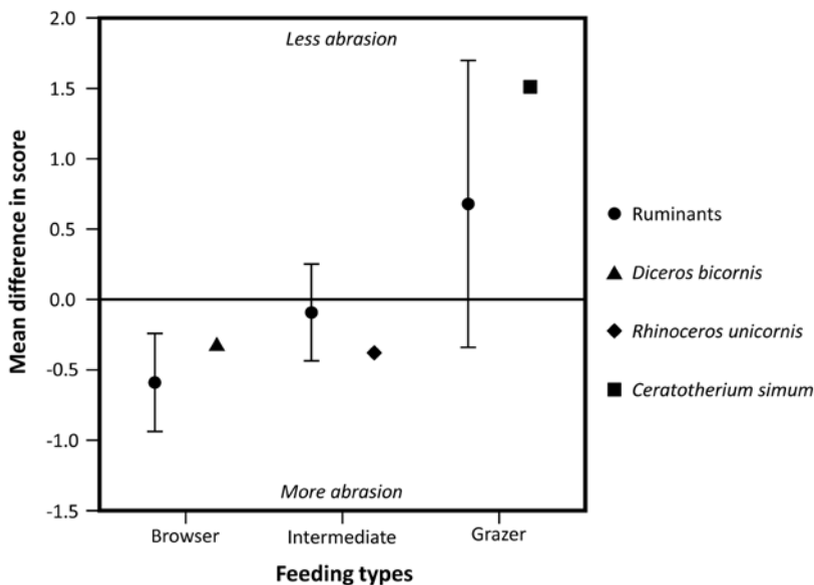


Fig. 2. Comparison of the mean score difference between captive and free-ranging ruminants and rhinoceroses (free-ranging – captive). The ruminant data is from Kaiser *et al.* (2009) and is a score difference for the whole M2 tooth position in the method of Fortelius and Solounias (2000). Rhino data, including the browser *Dicerus bicornis*, mixed feeding *Rhinoceros unicornis* and grazing *Ceratotherium simum* rhinoceros, is from this study and is a score difference for the posterior cusp of M2 in the method of Fortelius and Solounias (2000) with rhinoceros adjusted OR. Error bars indicate standard deviation between ruminant species.

Table 1. Linear mixed-effects models comparing the cusp shape (CS) score, occlusal relief (OR) score and mesowear score for cusp and tooth positions, within free-ranging and captive animals, and between these groups, in *Dicerus bicornis*, *Rhinoceros unicornis* and *Ceratotherium simum*. Significant differences are highlighted in **bold**.

Species	Score	Free-ranging		Captive		Origin*Tooth (Cusp)	
		Cusp	Tooth (Cusp)	Cusp	Tooth (Cusp)	P2-P4	M1-M2
Black rhino (<i>D. bicornis</i>)	CS	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
	OR	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
	Mesowear score	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
Greater one-horned rhino (<i>R. unicornis</i>)	CS	0.729	<0.001	0.549	0.007	0.016	0.134
	OR	<0.001	<0.001	0.356	<0.001	0.003	<0.001
	Mesowear score	0.085	<0.001	0.918	<0.001	0.005	<0.001
White rhino (<i>C. simum</i>)	CS	0.503	0.213	0.625	<0.001	<0.001	<0.001
	OR	0.013	0.014	0.264	<0.001	<0.001	0.021
	Mesowear score	0.220	0.081	0.727	<0.001	<0.001	<0.001

uneven and atypical tooth wear (Fig. 1).

In captive *D. bicornis*, mesowear scores were significantly more abrasion-dominated in both anterior and posterior cusps of P4, M1 and M2 compared to free-ranging specimens (all $p < 0.05$) (Table 2). By contrast, captive specimens had significantly less abrasion-dominated mesowear scores in the posterior cusp of P2 ($t_{21,30} = 2.47, p = 0.022$). Differences were caused by a combination of both CS and OR, with significant differences in both parameters. In captive *D. bicornis*, 57% of P2s were excluded due to advanced wear, whereas only 27% had been excluded in free-ranging specimens.

Wear stage exclusions did not affect the mean mesowear score of the P2.

In *R. unicornis*, despite overall significant differences between premolars and molars, no significant differences were found between captive and free-ranging specimens at individual cusp level. Captive specimens generally had blunter CS than free-ranging specimens, but, in both groups, score variance was high. In the posterior cusp of M2, mean OR and mesowear score differences between free-ranging and captive animals were similar to *D. bicornis*.

In captive *C. simum*, CS, OR and mesowear scores

Table 2. Differences in cusp shape score, occlusal relief score and mesowear score between free-ranging and captive *Diceros bicornis*, *Rhinoceros unicornis* and *Ceratotherium simum* rhinoceroses in each tooth and cusp position for the full dataset. Populations are compared by mean \pm standard deviation, the mean score difference (Score diff: free-ranging-captive) and Welch two sample *t*-tests. Significant differences are highlighted in **bold**, with values tending towards significance underlined.

Species	Tooth and Cusp position		n	Cusp shape score				Occlusal relief score				Mesowear wear				
				Free-ranging	Captive	Free-ranging	Captive	Score diff	p-value	Free-ranging	Captive	Score diff	p-value	Free-ranging	Captive	Score diff
Black rhinoceros (<i>D. bicornis</i>)	P2	A	20	9	3.7 \pm 0.7	3.7 \pm 0.7	-0.02	0.953	3.2 \pm 0.6	3.2 \pm 0.7	-0.02	0.933	3.4 \pm 0.5	3.4 \pm 0.6	-0.02	0.937
		P	20	10	2.6 \pm 1.1	1.9 \pm 1.0	0.70	<u>0.095</u>	2.6 \pm 0.8	1.8 \pm 0.9	0.75	0.044	2.6 \pm 0.8	1.9 \pm 0.7	0.73	0.022
	P3	A	18	12	2.5 \pm 0.9	2.9 \pm 1.0	-0.42	0.260	2.7 \pm 0.5	2.8 \pm 0.7	-0.11	0.641	2.6 \pm 0.6	2.9 \pm 0.7	-0.26	0.318
		P	18	10	1.3 \pm 0.8	1.9 \pm 0.6	-0.57	0.036	1.8 \pm 0.8	2.0 \pm 0.7	-0.17	0.559	1.6 \pm 0.4	2.0 \pm 0.6	-0.37	<u>0.084</u>
	P4	A	24	16	2.0 \pm 1.1	2.7 \pm 0.9	-0.69	0.032	2.4 \pm 0.5	2.8 \pm 0.4	-0.33	0.035	2.2 \pm 0.6	2.7 \pm 0.5	-0.51	0.007
		P	23	16	1.4 \pm 0.6	2.0 \pm 0.6	-0.57	0.008	1.7 \pm 0.6	2.0 \pm 0.7	-0.35	0.122	1.5 \pm 0.4	2.0 \pm 0.5	-0.46	0.009
	M1	A	22	19	2.0 \pm 0.7	2.6 \pm 0.8	-0.62	0.016	2.1 \pm 0.6	2.7 \pm 0.6	-0.60	0.002	2.0 \pm 0.5	2.7 \pm 0.6	-0.61	<0.001
		P	21	19	1.5 \pm 0.6	2.2 \pm 0.6	-0.68	<0.001	1.2 \pm 0.8	1.7 \pm 0.7	-0.50	0.042	1.4 \pm 0.5	1.9 \pm 0.6	-0.59	0.002
	M2	A	22	19	2.0 \pm 0.5	2.3 \pm 0.7	-0.26	0.171	2.2 \pm 0.5	2.6 \pm 0.6	-0.35	0.057	2.1 \pm 0.4	2.4 \pm 0.5	-0.31	0.044
		P	26	19	1.4 \pm 0.6	1.9 \pm 0.5	-0.56	0.002	0.5 \pm 0.6	1.0 \pm 0.8	-0.46	0.044	1.0 \pm 0.4	1.5 \pm 0.6	-0.51	0.002
Greater one-horned rhinoceros (<i>R. unicornis</i>)	P2	A	9	7	3.9 \pm 0.3	3.0 \pm 1.3	0.89	0.121	3.0 \pm 0.5	2.7 \pm 0.5	0.29	0.271	3.4 \pm 0.3	2.9 \pm 0.8	0.59	0.107
		P	9	7	3.6 \pm 0.9	3.6 \pm 1.1	-0.02	0.976	3.1 \pm 0.6	3.3 \pm 0.8	-0.17	0.626	3.3 \pm 0.7	3.4 \pm 0.9	-0.10	0.817
	P3	A	9	7	2.6 \pm 0.9	2.9 \pm 1.5	-0.30	0.641	2.9 \pm 0.3	2.7 \pm 0.5	0.17	0.436	2.7 \pm 0.5	2.8 \pm 0.8	-0.06	0.860
		P	9	6	3.2 \pm 1.1	3.7 \pm 0.5	-0.44	0.312	2.8 \pm 0.4	2.7 \pm 0.8	0.11	0.769	3.0 \pm 0.8	3.2 \pm 0.6	-0.17	0.644
	P4	A	7	5	2.3 \pm 0.5	2.8 \pm 1.3	-0.51	0.440	2.6 \pm 0.5	2.8 \pm 0.4	-0.23	0.441	2.4 \pm 0.4	2.8 \pm 0.7	-0.37	0.320
		P	8	6	2.5 \pm 0.9	3.2 \pm 1.0	-0.67	0.226	1.9 \pm 0.6	2.3 \pm 0.8	-0.46	0.284	2.2 \pm 0.8	2.8 \pm 0.8	-0.56	0.196
	M1	A	12	8	2.3 \pm 0.6	2.8 \pm 0.7	-0.50	0.127	2.5 \pm 0.7	2.3 \pm 0.5	0.25	0.339	2.4 \pm 0.6	2.5 \pm 0.4	-0.13	0.562
		P	11	9	2.2 \pm 0.9	2.2 \pm 1.2	-0.04	0.934	2.5 \pm 0.5	2.3 \pm 0.9	0.12	0.719	2.3 \pm 0.6	2.3 \pm 0.8	0.04	0.905
	M2	A	11	8	2.0 \pm 0.8	2.4 \pm 0.9	-0.38	0.364	2.1 \pm 0.3	2.4 \pm 0.7	-0.28	0.335	2.0 \pm 0.5	2.4 \pm 0.6	-0.33	0.224
		P	12	9	1.8 \pm 0.6	2.1 \pm 0.9	-0.36	0.331	0.8 \pm 0.7	1.3 \pm 0.9	-0.50	0.179	1.3 \pm 0.6	1.7 \pm 0.8	-0.43	0.207
White rhinoceros (<i>C. simum</i>)	P2	A	7	5	4.0 \pm 0.0	4.0 \pm 0.0	0.00	1.000	3.7 \pm 0.5	3.6 \pm 0.5	0.11	0.719	3.9 \pm 0.2	3.8 \pm 0.3	0.06	0.719
		P	7	5	4.0 \pm 0.0	4.0 \pm 0.0	0.00	1.000	3.1 \pm 0.4	3.8 \pm 0.4	-0.66	0.029	3.6 \pm 0.2	3.9 \pm 0.2	-0.33	0.029
	P3	A	9	8	3.9 \pm 0.3	3.4 \pm 0.7	0.51	0.104	3.2 \pm 0.4	2.9 \pm 0.4	0.35	<u>0.092</u>	3.6 \pm 0.3	3.1 \pm 0.4	0.43	0.039
		P	9	8	4.0 \pm 0.0	4.0 \pm 0.0	0.00	1.000	3.7 \pm 0.5	3.4 \pm 0.7	0.29	0.367	3.8 \pm 0.3	3.7 \pm 0.4	0.15	0.367
	P4	A	8	11	4.0 \pm 0.0	3.3 \pm 0.8	0.73	0.012	3.3 \pm 0.5	2.7 \pm 0.6	0.52	0.056	3.6 \pm 0.2	3.0 \pm 0.7	0.63	0.013
		P	8	10	4.0 \pm 0.0	3.8 \pm 0.6	0.20	0.343	3.8 \pm 0.5	3.3 \pm 0.5	0.45	0.062	3.9 \pm 0.2	3.6 \pm 0.4	0.33	<u>0.063</u>
	M1	A	7	9	3.9 \pm 0.4	3.9 \pm 0.3	-0.03	0.864	3.0 \pm 0.6	3.4 \pm 0.7	-0.44	0.194	3.4 \pm 0.3	3.7 \pm 0.5	-0.24	0.280
		P	9	8	3.6 \pm 0.9	3.9 \pm 0.4	-0.32	0.339	3.6 \pm 0.5	3.3 \pm 0.5	0.31	0.223	3.6 \pm 0.6	3.6 \pm 0.3	-0.01	0.977
	M2	A	9	11	3.8 \pm 0.7	4.0 \pm 0.0	-0.22	0.347	3.0 \pm 0.5	3.1 \pm 0.5	-0.09	0.701	3.4 \pm 0.5	3.5 \pm 0.3	-0.16	0.449
		P	9	12	3.8 \pm 0.7	2.6 \pm 1.2	1.19	0.011	3.3 \pm 0.7	2.5 \pm 0.7	0.83	0.014	3.6 \pm 0.6	2.5 \pm 0.8	1.01	0.005

varied, revealing both significantly more and less abrasion-dominated wear patterns than their free-ranging conspecifics. The posterior cusp of M2 was significantly sharper ($t_{17.53}=2.83, p=0.011$) and higher ($t_{16.91}=2.7262, p=0.014$) in captive than in free-ranging specimens.

In 'classical' mesowear, *D. bicornis* had a mean score difference in the posterior cusps of M2 of -0.33 (free-ranging mean $=0.6\pm0.5$; captive $=0.9\pm0.5$), *R. unicornis* -0.39 (0.8 ± 0.4 ; 1.2 ± 0.7) and *C. simum* 1.5 (3.7 ± 1.0 ; 2.2 ± 1.5) (Fig. 2). Despite a larger difference in *R. unicornis* than *D. bicornis*, no significant difference was detected in *R. unicornis* ($t_{12.04}=-1.56, p=0.144$), whereas significant differences were detected between free-ranging and captive *D. bicornis* ($t_{37.66}=-2.15, p=0.038$) and *C. simum* ($t_{18.89}=2.78, p=0.012$).

Cusp and tooth-specific wear signatures

In both free-ranging and captive *D. bicornis*, anterior cusps were significantly more abrasion-dominated than posterior cusps (LME: all $p<0.001$) (Table 1). By contrast, no significant differences were observed in CS and mesowear scores between cusps in free-ranging and captive *R. unicornis* and *C. simum*. Some OR cusp differences were detected in free-ranging *R. unicornis* and *C. simum*, whereas no differences were detected in captive animals.

Tooth position differences were detected in both free-ranging and captive *D. bicornis* (all $p<0.001$) and *R. unicornis* (all $p<0.01$). In contrast to free-ranging black rhinoceroses that exhibited a negative tooth wear gradient with teeth becoming less abrasion-dominated from P2-M2, captive black rhinoceroses only exhibited a significantly blunter CS in the anterior cusp of P2 ($F_{4.37}=4.91, p=0.002$; Dunnett's T3 post-hoc M1-M2: $p<0.01$) and a significantly higher OR in the posterior cusps of M2 ($F_{4.37}=4.97, p=0.001$; P3-P4: $p<0.05$) (Fig. 1). In *C. simum*, captive specimens exhibited larger differences in CS, OR and mesowear score ($p<0.001$) along the tooth row than free-ranging conspecifics (OR: $p=0.014$). Differences in captivity were predominantly due to a significantly sharper M2 (P2-M1: $p<0.05$) and higher posterior cusp of M2 (P2-M1: $p=0.009$).

Discussion

The results indicate that captive *Diceros bicornis* (browsers) exhibit more abrasion-dominated tooth wear than their free-ranging conspecifics, whereas captive

Ceratotherium simum (grazers) exhibit less abrasion-dominated tooth wear, particularly in the posterior cusp of M2. In *Rhinoceros unicornis* (mixed-feeder), fewer differences were exhibited between the free-ranging and captive animals. Overall, differences in tooth wear pattern between the species were less pronounced in captive than in free-ranging rhinoceroses, particularly in cusp shape (CS), which did not significantly differ in the posterior cusp of M2. CS convergence indicates that the abrasiveness and/or physical properties of the diets fed in captivity are similar across all rhinoceros species. Whilst this may not be problematic for high-crowned *C. simum*, high-abrasion diets for low-crowned *D. bicornis* and also *R. unicornis* could result in the premature loss of tooth functionality, leading to reduced food processing ability and, consequently, malnourishment.

Limitations

Although a method for taking dental impressions from live rhinoceroses exists (Wucher, 1994), measurements of living specimens were not possible due to the necessity of immobilising animals for the procedure. Instead, dental casts of museum specimens were used, for which no feeding records were available. Therefore, it cannot be stated with certainty whether our findings are representative of current captive feeding practices. Individual age was indirectly controlled for in this study using functional wear stages; nevertheless, mesowear is affected by ontogeny (Rivals *et al.*, 2007) and age effects cannot be excluded completely. The sample sizes of *R. unicornis* ($n=11$) and *C. simum* ($n=15$) were more limited than that of *D. bicornis* ($n=23$). Mesowear scoring for rhinoceroses uses different OR boundaries than ruminants; thus direct comparisons with other taxa should be limited to qualitative evaluations.

Differences between the wild and captivity

Overall, captive *D. bicornis* had more abrasion-dominated CS, OR and, consequently, more abrasion-dominated mesowear scores than their free-ranging conspecifics. Clauss *et al.* (2007) and Kaiser *et al.* (2009) suggested increased abrasion may be due to a higher intake of abrasives as plant phytoliths (silica) via grass products (such as grass hay, but also others like grain bran), and also via silica-based flow enhancers in pelleted diets. Clauss and Hatt (2006) reported two cases of excessively worn teeth in *D. bicornis* fed on grass hay. The causes of tooth wear are currently debated in the scientific literature, with some arguing

that environmental abrasives have more influence (Dammuth and Janis, 2011; Lucas *et al.*, 2013). The high impact of external abrasives on tooth wear (Healy and Ludwig, 1965) notwithstanding, Schulz *et al.* (2013) recently found that the 3D surface textures of rabbits fed grass or lucerne distinctly differed, indicating that internal abrasives also affect dental tissue. Additionally, Rabenold and Pearson (2011) demonstrated that molar enamel thickness related to the phytolith content of the diet in 12 primate species, which suggests that increased dental durability also evolved as an adaptation to the effect of internal abrasives. Kaiser *et al.* (2009) suggested that an increase in environmental abrasives is unlikely in captivity because of increased feeding hygiene and industrial processing techniques for feeds designed to minimise soil contamination. Castell (2005) reported higher faecal acid insoluble ash (silica) values for captive *D. bicornis* ($3.7 \pm 1.9\%$ dry matter [DM], range 1.2–10.5) than reported for free-ranging *D. bicornis* ($1.7 \pm 0.6\%$ DM; Hummel *et al.*, 2011), but approaching the range measured in free-ranging *C. simum* ($7.5 \pm 1.3\%$ DM), indicating that the intake of abrasives (whether dietary, environmental, or both) is higher in captive than free-ranging browsers. Therefore, diet-specific properties may also play a significant role in tooth wear, and abrasive feeds may be causing excessive tooth wear in captive browsers.

The intrinsic toughness, physical form and chemical properties of the diet may also affect tooth wear. In domestic horses, feeding pelleted compound diets increases the vertical and decreases the lateral excursion of the chewing movement (Bonin *et al.*, 2007). Taylor *et al.* (2013) suggested an increased vertical upstroke could blunt sharp cusp tips. In captive *D. bicornis*, CS was significantly blunter than in free-ranging specimens, which could also be caused by chewing compound feeds. Furthermore, chewing higher proportions of compound feeds, in parallel to similar feeding practices, could explain the lack of difference in CS scores between rhinoceros species in captivity. Imbalances of calcium (Ca), phosphorus (P) and/or vitamin D3 have also been shown to affect tooth growth and wear rates (McRoberts *et al.*, 1965; Harcourt-Brown, 1996). Low serum Ca and imbalanced Ca:P ratios have been reported in captive ruminants, with enamel hypoplasia and urolithiasis in captive giraffes (Miller *et al.*, 2003; Franz-Odenaal, 2004). By contrast, hypophosphatemia has been reported in captive *D. bicornis* (Dennis *et al.*, 2007). The link between mineral imbalances and tooth wear should be explored further, but might not be a substantial contributing factor in captive

rhinoceroses.

In *R. unicornis*, overall differences were detected between free-ranging and captive animals, but differences were not significant at individual cusp level. Taylor *et al.* (2013) suggested variability in scores in free-ranging *R. unicornis* may be due to seasonal variation in diet (53–87% grass; Pradhan *et al.*, 2008). Variation in captivity likely reflects differences in feeding practices between institutions. The CS, OR and mesowear scores of free-ranging *R. unicornis* are closer to scores attained by free-ranging *D. bicornis* rather than free-ranging *C. simum*, which indicates a relatively low ingestion of environmental abrasives (Taylor *et al.*, 2013). Compared to ruminant intermediate feeders studied by Kaiser *et al.* (2009), *R. unicornis* appeared to experience slightly more abrasion in captivity (Fig. 2). In addition to these scores, the generally low crown height of *R. unicornis* (Janis, 1988) indicates that high-abrasion diets may not be suitable for captive individuals of this species.

In captive *C. simum*, scores varied considerably along the tooth row, which indicates teeth are not wearing evenly. The posterior cusp of M2, the cusp typically used to interpret diet, is significantly higher and sharper than that of their free-ranging conspecifics, indicating the ingesta of captive *C. simum* is less abrasive. Kaiser *et al.* (2009) suggest captive grazers may experience less abrasion-dominated tooth wear due to reduced dietary and environmental abrasives because of the temperate climate, industrial feed processing techniques and increased feeding hygiene. In addition, the lateral excursion while chewing pelleted compound feed may be insufficient to wear the entire occlusal surface, which may facilitate the development of sharp enamel points on the edges of teeth and cause uneven tooth wear (Bonin *et al.*, 2007). Elia *et al.* (2010) also found that domestic horses chewed more when fed hay (43,476 chews/day) than on pelleted diets (10,036 chews/day), which could decrease levels of tooth wear.

Cusp and tooth position differences

In both free-ranging and captive *D. bicornis*, anterior cusps were significantly more abrasion-dominated than posterior cusps. Rhino teeth are asymmetrical, with anterior cusps proportionally smaller than the posterior. Taylor *et al.* (2013) speculated that cusp differences in rhinoceroses are morphological adaptations to browse. Despite more abrasion-dominated scores, captive *D. bicornis* still exhibited significant differences between cusps, suggesting a morphological influence. This

means that mesowear does not only reflect diet but also structural elements, which raises questions about the effects of tooth morphology and occlusion on the development of mesowear patterns and their functional relevance. However, anterior and posterior cusp rows were both significantly different between free-ranging and captive animals, which suggests that both cusps were also affected by ingesta-specific properties.

Free-ranging *D. bicornis* and *R. unicornis* exhibited a negative tooth wear gradient and free-ranging *C. simum* exhibited homogenous tooth wear (Taylor *et al.*, 2013). By contrast, fewer differences were exhibited in tooth wear gradients between captive species. Several intrinsic (jaw biomechanics) and extrinsic factors (ingesta abrasives) might explain tooth wear gradients. Taylor *et al.* (2013) suggested tooth wear gradients may be caused by ingesta abrasiveness, because free-ranging *C. simum* consume high *absolute* amounts of ingesta abrasives, which will override other signals. The results of this study corroborate an ingesta-specific influence because the gradients differed between free-ranging and captive animals. Environmental abrasives probably affect the anterior teeth more severely before being mixed into the bolus. Captive *D. bicornis* may consume less environmental abrasives and more endogenous dietary abrasives. Thus, captive *D. bicornis* may experience less *relative* changes in abrasiveness along the tooth row and an overall increase in the *absolute* ingesta abrasiveness, leading to more homogenous tooth wear. Captive *C. simum* may experience a reduction in the *absolute* abrasiveness of ingesta compared to free-ranging conspecifics, which may lead to more pronounced *relative* changes in wear from P2-M2. The results do not negate a role of jaw biomechanics, as the OR of the posterior cusp of M2 of all captive specimens was higher than more anterior tooth positions.

In free-ranging *D. bicornis*, the P2 was significantly more abrasion-dominated than the other tooth positions, which Taylor *et al.* (2013) suggest may indicate the P2 has a role in food cropping as rhinoceroses do not have occluding incisors. By contrast, captive *D. bicornis* had homogenous tooth wear between P2-M1 in the posterior cusp tooth row. In captivity, animals are mostly fed loose hay and pelleted feeds, which may not require cropping, and may be less contaminated with environmental abrasives, such as dust adherent on browse. Therefore, food presentation, particularly the combination of reduced bite force and reduced environmental abrasives, could also affect tooth wear gradients. However, in captive *D. bicornis*, 57% of P2 teeth were excluded due to advanced wear compared to 27% in the

free-ranging animals. Although age differences cannot be excluded, the P2 wear in captive specimens highlights that despite less abrasion-dominated mesowear scores, total tooth wear (volume loss) was still high (or higher) than in free-ranging specimens, which raises questions about tooth wear rate relative to mesowear pattern.

Implications

The results in rhinoceroses and ruminants (Kaiser *et al.*, 2009) suggest that diets fed to captive browsers are more abrasive than their diet in the wild. Although fewer differences were exhibited in mixed-feeding *R. unicornis*, the low crown height of this species still suggests high-abrasion diets may not be suitable. Excessive tooth wear could have serious implications on captive animal health, welfare and longevity and, consequently, conservation and re-introduction strategies. In the past, captive-raised *D. bicornis* have been re-introduced into protected reservations. Releasing rhinoceroses with excessive tooth wear relative to age could to some extent jeopardise the viability of such programmes. By contrast, captive *C. simum* exhibited significantly less abrasion-dominated tooth wear than free-ranging conspecifics. Decreased tooth wear may result in prolonged tooth functionality, which may have a positive influence on longevity. However, insufficient tooth wear may lead to uneven wear and tooth pathologies, which could negatively impact on captive animal health and may contribute to other dental diseases. Dental health checks should therefore be part of routine animal health checks for rhinoceroses during any veterinary treatment. As mentioned above, Wucher (1994) describes a method of taking dental impressions from rhinoceroses, which could be used to aid in the diagnosis of tooth wear problems. To help prevent tooth wear problems, browsers and intermediate feeders should be fed on a dicot-based (browse) diet that is low in abrasives, including dicot-based pellets, and grazers should be fed on a monocot-based diet, especially grass and grass hay. Pelleted feed quantities should probably be reduced. Further research is required into the abrasiveness of the diets of captive wild animals.

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References

- Bonin SJ, Clayton HM, Lanovaz JL, Johnston T. 2007. Comparison of mandibular motion in horses chewing hay and pellets. *Equine Veterinary Journal* 39: 258-262.
- Castell JC. 2005. Untersuchungen zu Fütterung und Verdauungsphysiologie am Spitzmaulnashorn (*Diceros bicornis*). Dissertation thesis, University of Munich.
- Clauss M, Hatt J-M. 2006. The feeding of rhinoceros in captivity. *International Zoo Yearbook* 40: 197-209.
- Clauss M, Franz-Odenaal TA, Brasch J, Castell JC, Kaiser TM. 2007. Tooth wear in captive giraffes (*Giraffa camelopardalis*): mesowear analysis classifies free-ranging specimens as browsers but captive ones as grazers. *Journal of Zoo and Wildlife Medicine* 38: 433-445.
- Damuth J, Janis CM. 2011. On the relationship between hypsodonty and feeding ecology in ungulate mammals, and its utility in palaeoecology. *Biological Reviews* 86: 733-758.
- Dennis PM, Funk JA, Rajala-Schultz PJ, Blumer ES, Miller RE, Wittum TW, Saville WJA. 2007. A review of some of the health issues of captive black rhinoceroses (*Diceros bicornis*). *Journal of Zoo and Wildlife Medicine* 38: 509-517.
- Elia JB, Erb HN, Haupt KA. 2010. Motivation for hay: effects of a pelleted diet on behavior and physiology of horses. *Physiology & Behavior* 101: 623-627.
- Fortelius M, Solounias N. 2000. Functional characterisation of ungulate molars using the abrasion-attrition wear gradient: a new method for reconstructing paleodonts. *American Museum Novitates* 3301: 1-36.
- Franz-Odenaal TA. 2004. Enamel hypoplasia provides insights into early systemic stress in wild and captive giraffes (*Giraffa camelopardalis*). *Journal of Zoology* 263: 197-206.
- Groves CP, Fernando P, Robovský J. 2010. The sixth rhino: a taxonomic re-assessment of the critically endangered Northern white Rhinoceros. *PLoS One* 5: e9703.
- Harcourt-Brown F. 1996. Calcium deficiency, diet and dental disease in pet rabbits. *Veterinary Record* 139: 567-571.
- Healy WB, Ludwig TG. 1965. Wear of sheeps teeth. I. The role of ingested soil. *New Zealand Journal of Agricultural Research* 8: 737-752.
- Hummel J, Findeisen E, Südekum K-H, Ruf I, Kaiser TM, Bucher M, Clauss M, Codron D. 2011. Another one bites the dust: faecal silica levels in large herbivores correlate with high-crowned teeth. *Proceedings of the Royal Society B* 278: 1742-1747.
- Janis C. 1988. An estimation of tooth volume and hypsodonty indices in ungulate mammals and the correlation of these factors with dietary preferences. Pp. 367-387 in: Russell DE, Santoro J-P, Signogneau-Russell D, eds, Proceedings of the Teeth revisited Proceedings of the VIIth International Symposium on Dental Morphology Mémoires du Muséum national d'Histoire Naturelle, Paris (serie C).
- Kaiser TM, Brasch J, Castell JC, Schulz E, Clauss M. 2009. Tooth wear in captive wild ruminant species differs from that of free-ranging conspecifics. *Mammalian Biology* 74: 425-437.
- Kojola I, Helle T, Huhta E, Niva A. 1998. Foraging conditions, tooth wear and herbivore body reserves: a study of female reindeer. *Oecologia* 117: 26-30.
- Lemaître J-F, Gaillard J-M, Lackey LB, Clauss M, Müller DWH. 2013. Comparing free-ranging and captive populations reveals intra-specific variation in aging rates in large herbivores. *Experimental Gerontology* 48: 162-167.
- Lucas PW, Omar R, Al-Fadhlah K, Almusallam AS, Henry AG, Michael S, Thai LA, Watzke J, Strait DS, Atkins AG. 2013. Mechanisms and causes of wear in tooth enamel: implications for hominin diets. *Journal of The Royal Society Interface* 10: 20120923.
- Martin Jurado O, Clauss M, Hatt J-M. 2008. Irregular tooth wear and longevity in captive wild ruminants: a pilot survey of necropsy reports. *Journal of Zoo and Wildlife Medicine* 39: 69-75.
- McRoberts MR, Hill R, Dalgarno AC. 1965. The effects of diets deficient in phosphorus, phosphorus and vitamin D, or calcium, on the skeleton and teeth of the growing sheep. *Journal of Agriculture* 65: 1-13.
- Miller M, Weber M, Valdes E, Fontenot D, Neiffer D, Robbins PK, Terrell S, Stetter M. 2003. Hypomagnesemia, hypocalcemia, and ruminitis in unglulates: an under-recognised syndrome. Proceedings of the American Association of Zoo Veterinarians, Minneapolis, USA, pp 15-20.
- Müller DWH, Bingaman Lackey L, Streich WJ, Fickel J, Hatt J-M, Clauss M. 2011. Mating system, feeding type and *ex situ* conservation effort determine life expectancy in captive ruminants. *Proceedings of the Royal Society B* 278: 2076-2080.
- Ozaki M, Kaji K, Matsuda N, Ochiai K, Asada M, Ohba T, Hosoi E, Tado H, Koizumi T, Suwa G, Takatsuki S. 2010. The relationship between food habits, molar wear and life expectancy in wild sika deer populations. *Journal of Zoology* 280: 202-212.
- Pradhan NMB, Wegge P, Moe SR, Shrestha AK. 2008. Feeding ecology of two endangered sympatric megaherbivores: Asian elephant *Elephas maximus* and greater one-horned rhinoceros *Rhinoceros unicornis* in lowland Nepal. *Wildlife Biology* 41: 147-154.
- R Core Team. 2012. R: A language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria.
- Rabenold D, Pearson OM. 2011. Abrasive, silica phytoliths and

- the evolution of thick molar enamel in primates, with implications for the diet of *Paranthropus boisei*. *PLoS One* 6: e28379.
- Rivals F, Muhlbachler MC, Solounias N. 2007. Effect of ontogenetic-age distribution in fossil and modern samples on the interpretation of ungulate paleodiets using the mesowear method. *Journal of Vertebrate Paleontology* 27: 763-767.
- Ruxton GD. 2006. The unequal variance t-test is an underused alternative to Student's t-test and the Mann-Whitney U test. *Behavioral Ecology* 17: 688-690.
- Schulz E, Piotrowski V, Clauss M, Mau M, Merceron G, Kaiser TM. 2013. Dietary abrasiveness is associated with variability of microwear and dental surface texture in rabbits. *PLoS ONE* 8: e56167.
- Skogland T. 1988. Tooth wear by food limitation and its life history consequences in wild reindeer. *Oikos* 51: 238-242.
- Taylor LA, Kaiser TM, Schwitzer C, Müller DHW, Cordon D, Clauss M, Schulz E. 2013. Detecting inter-cusp and inter-tooth wear patterns in Rhinocerotids. *PLoS One* 8: e80921.
- Winkler DE, Kaiser TM. 2011. A case study of seasonal, sexual and ontogenetic divergence in the feeding behaviour of the moose (*Alces alces* Linné, 1758). *Verhandlungen des naturwissenschaftlichen Vereins Hamburg* 46: 331-348.
- Wucher M. 1994. A technique for making dental impressions and casts of immobilised black rhinoceros (*Diceros bicornis*) and white rhinoceros (*Ceratotherium simum*). Proceedings of a Symposium on "Rhinos as Game Ranch Animals", Onderstepoort, pp. 164-167.

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SI. Museum and specimen information.

