

# Elephant-mediated habitat modifications and changes in herbivore species assemblages in Sabi Sand, South Africa

Willem F. de Boer<sup>1</sup> · Jordi W. A. Van Oort<sup>1</sup> · Michael Grover<sup>2</sup> · Mike J. S. Peel<sup>3</sup>

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**Abstract** Elephant *Loxodonta africana* conservation might indirectly influence the wider herbivore community structure, as elephants have the ability to significantly modify the savanna habitat. Uncertainty remains as to the consequences of these effects, as elephants might either compete with other species or facilitate foraging especially for grazers and smaller browsing species by increasing the amount of grass or the amount of browse at lower feeding heights. We studied these potential cascading effects of elephants by using 16 years of data (1992–2011) from the Sabi Sand Wildtuin, South Africa, which showed a steady increase in elephant densities from 0.12 to 2.03 elephants/km<sup>2</sup> over this period. We demonstrate that tree densities, and browse availability at feeding heights below 2 m, decreased with increasing elephant densities, and that there was no positive effect of elephants on browse availability. The changes in elephant densities were good predictors ( $R^2_{\text{adj}} > 0.50$ ) in explaining population fluctuations of other herbivore species. The total body mass of grazers increased more than that of the browsers, shifting the community toward a grazer and megaherbivore-dominated community. An increasing density of elephants changes the composition of the herbivore community, as mesobrowsers are unable to benefit from the impact of elephants on trees, but megagrazers show strong positive responses. Hence, changes in elephant

densities as a result of poaching or conservation may trigger cascading community effects. These are neglected but important consequences of (negative or positive) human impacts on elephant numbers, especially in restricted areas such as reserves and national parks.

**Keywords** Cascading effects · Competition · Facilitation · Keystone species · Tree density · *Loxodonta africana*

## Introduction

The impact of African elephants (*Loxodonta africana*) on the occurrence and density of other herbivore species that share their savanna ecosystems is a highly controversial topic. Both negative and positive effects of elephants on other herbivore species have been claimed, as a consequence of the impact of elephants on the vegetation structure, biomass, and species composition. Elephants are known to push over, debark, and break trees, and various studies have found a decline in tree densities in the presence of elephants (Cumming et al. 1997; Shannon et al. 2008; Woolley et al. 2011). Even at comparatively low elephant densities (<0.2 elephants/km<sup>2</sup>), the densities of certain tree species show signs of decline (Cumming et al. 1997; Shannon et al. 2008). By the early 2000s when elephant densities exceeded 0.5 animals/km<sup>2</sup> in the Kruger National Park (KNP), the extent of the woody layer in parts of the Park had shown significant decline (Whyte et al. 2003), and elephants are regarded as the primary agent of treefall, especially of trees in the 5–9-m height class (Asner and Levick 2012). In the proximity of large perennial rivers, current elephant densities in the KNP exceed 1.5 animals/km<sup>2</sup> (Smit and Ferreira 2010). Elephant impacts can also vary spatially and can be site- or context-dependent (Eckhardt et al. 2000; Guldmond and Van Aarde 2008).

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✉ Willem F. de Boer  
fred.deboer@wur.nl

<sup>1</sup> Resource Ecology Group, Wageningen University,  
Droevendaalsesteeg 3a, 6708 PB Wageningen, The Netherlands

<sup>2</sup> Sabi Sand Wildtuin, P.O. Box 105, Skukuza 1350, South Africa

<sup>3</sup> Agricultural Research Council, Animal Production Institute,  
P.O. Box 7063, Nelspruit 1200, South Africa

This impact of elephants on trees in the system has raised the concern of scientists and managers alike (Whyte et al. 1998; Van Aarde et al. 1999; Van Aarde et al. 2006; Kerley et al. 2008), as the decline in woody cover could negatively affect the browsing species in the system. Fritz et al. (2002) concluded that elephants compete with the smaller mesobrowser species for browse resources, as the relative contribution of these species declined with increasing elephant densities. Valeix et al. (2008) also reported a possible negative influence of elephants on other herbivore species. Elephant impact, especially on the larger trees, could also have negative consequences for large browsing species such as giraffe (Fritz et al. 2002).

Despite the fact that such studies have demonstrated that elephants have a detrimental effect on trees in savanna ecosystems, there is, however, also evidence for positive secondary effects of tree utilization by elephants. The preferred feeding height of elephants is <2 m (Stokke and Du Toit 2000), and, in some cases, elephants appear to facilitate smaller browser species by increasing the quality and biomass availability of trees at lower feeding heights (Du Toit et al. 1990; Smallie and O'Connor 2000; Kohi et al. 2011; Kohi 2013). Further, impala (*Aepyceros melampus*) and kudu (*Tragelaphus strepsiceros*) benefitted from the conversion of woodland to shrubland by elephants in Botswana (Rutina et al. 2005; Makhabu et al. 2006). Elephants may even keep trees and shrubs below a certain maximum height by coppicing trees and shrubs, thus creating browsing lawns (Smallie and O'Connor 2000). Moreover, utilization by elephants can activate dormant buds, and as the number of buds is reduced by elephant browsing, the within-tree competition for resources decreases, resulting in fewer but larger shoots (Du Toit et al. 1990; Järemo et al. 1996). Some studies have suggested that some heavily browsed trees might not only produce higher quantities of foliage but also foliage of a higher quality (Du Toit et al. 1990; Smallie and O'Connor 2000; Kohi 2013). Regrowing shoots are often rich in nutrients and have relatively low concentrations of secondary compounds, turning them into high-quality forage resources which are more susceptible to rebrowsing (Price 1991; Kohi et al. 2011). So, once a tree has been browsed, the likelihood of it being rebrowsed increases (Skarpe et al. 2000). Browsed trees also retained their leaves further into the dry season, and hence, elephants may facilitate other browser species (Kohi 2013).

Grasses can benefit from the release of competition with trees (Scholes and Archer 1997; Fritz et al. 2002; van Langevelde et al. 2003), and opening up of woodlands could therefore have positive effects on grazing herbivores (Parker 1983; Kerley et al. 2008). However, a decrease in tree cover can also increase the visibility for prey and predators, and thereby influence the predation risk (Valeix et al. 2011).

In response to all these reported effects, one might anticipate negative effects of elephants on other browser species in the system at high elephant densities, but a potentially positive effect at low-intermediate densities. The opening up of the canopy and the decrease in tree densities are expected to positively influence the grazers over the entire elephant density gradient. None of the studies mentioned above, however, presented data at appropriate spatial and temporal scales to explore this question meaningfully. Here, we use a long-term data set with measures of tree height and density, as well as concurrent surveys estimating herbivore population sizes. These data were collected at the Sabi Sand Wildtuin-MalaMala complex, South Africa, between 1992 and 2011. We quantify the effect of elephants on the woody layer and test whether elephants were correlated with the population sizes of other herbivore species in the system. The elephant densities in Sabi Sand Wildtuin-MalaMala complex increased from 0.15 animals/km<sup>2</sup> in 1992 to 2.55 animals/km<sup>2</sup> in 2011. These annual monitoring data offer a unique opportunity to analyze the impact of elephants on the vegetation and, through that, on the rest of the herbivore community. It is envisaged that this will increase our understanding of how a natural herbivore community might change under the contrasting impacts of a successful conservation effort or increased poaching (de Boer et al. 2013).

Based on the considerations offered above, we expect that (i) mesobrowser species, such as common duiker (*Sylvicapra grimmia*), bushbuck (*Tragelaphus scriptus*), kudu, and steenbok (*Raphicerus campestris*), should increase at low-intermediate elephant densities, as they could benefit from the large availability of high quality browse at lower feeding heights and decrease at high elephant densities. (ii) The impala, an abundant mixed feeder (Kos et al. 2012) and the nyala (*Tragelaphus angasii*), are expected to track the mesobrowser response, because of their reliance on browse in their diet, although to a lesser extent than the pure browsers. Finally, as above, (iii) the opening up of the canopy and the decrease in tree densities is expected to positively influence the grazers over the entire elephant density gradient.

## Materials and methods

### Study area

The Sabi Sand Wildtuin-MalaMala complex (S 24° 47', E 31° 29'), South Africa, covers an area of some 620 km<sup>2</sup> and shares its northern and north-eastern boundaries with the Manyeleti Game Reserve and southern and eastern borders with the Kruger National Park (KNP). It is dominated by tall shrublands and moderately dense woodlands. The rainy season is from October to April and the region's mean annual precipitation calculated from July in the preceding year to

June in the current year averaged 615 mm (1990–2011, SD=234, three rainfall gauges at Gowrie, Shaws and Newington with monthly data cover the entire study 1991–2011). The mean total dry season rainfall, which has been correlated with elephant population growth and densities of other herbivore species (Ogutu and Owen-Smith 2003; Valeix et al. 2008), was calculated from May to September, averaging 38 mm (SD=24) over the study period. The complex is characterized by an *Acacia* and *Combretum* woodland, and more than 50 % of all trees belong to only 8 species (in order of abundance): *Combretum apiculatum*, *Dalbergia melanoxylon*, *Acacia exuvialis*, *Dichrostachys cinerea*, *Acacia nigrescens*, *Terminalia sericea*, *Ormocarpum trichocarpum*, *Albizia harveyi*. Of the vegetation monitoring sites ( $N=43$ ), 30 % were burned once every 30 to 15 years, 16 % once every 10–8 years, 21 % once every 6–5 years, 12 % once every 4 years, and 21 % once every 3 years, but fire tends to exert a smaller effect on the vegetation structure and heterogeneity than herbivory (Levick et al. 2009).

### Sampling

The woody layer within the reserves was monitored annually in November–December from 1993 to 2011 by the Animal Production Institute of the Agricultural Research Council (ARC-API) using a 100×2-m belt transect in each of 43 permanent plots in which the number of all woody plants within a belt transect over four height classes (0–1.0, 1.1–2.0, 2.1–5.0, >5.1 m; Peel et al. 2005) was recorded, together with the numbers of stems per woody plant (recorded since 1995). The standing crop of the herbaceous layer was recorded from 1997 onward using a disk pasture meter which measures the herbaceous standing biomass in the presence of fire, herbivory, plant senescence, and decomposition (Zambatis et al. 2006).

Annual game counts have been done since 1992 in August–September, using a Bell Jet Ranger helicopter. The team consisted of a pilot and a front observer and two experienced back-seat observers who spot and count game. The front observer also records the data (species, totals, sex, and age where possible) on a computer linked to the helicopter's GPS thus providing a spatial distribution of the animals counted. The counts are carried out during winter months when foliage cover is at its lowest. The survey uses a strip width of 500 m (line lengths up to 15 km) with continuous strips at an altitude of 90 m (Sutherland and Peel 2011). The mammal species present, together with their foraging characteristics, are listed in Table 1.

Fences between the Sabi Sand Wildtuin-MalaMala complex and KNP were dropped in 1993; so, changes in population densities are partly due to animal dispersal between the two protected area, and we were unable to distinguish

between population growth and movements of animals to and from the area.

Since 2004, a total of about 2500 wildebeest (*Connochaetes taurinus*) have also been introduced in several batches in an effort to stabilize the wildebeest population, and the wildebeest population fluctuations were therefore not analyzed in detail. Data on the carnivore species were not available over the entire study period and could therefore not be included in the analysis. Small numbers of elephant, buffalo, and rhinos have also been removed from the area. On average, these represent <3 % of the population of each of the species and fall within the natural year-to-year variation of the species. These data were therefore excluded from the analysis.

Estimated predator population sizes were not available for the entire study period, but sometimes only for the last 4 years, and the impact of predators on prey population was therefore not analyzed further. The estimated population sizes were as follows: lion (*Panthera leo*) 105, leopard (*Panthera pardus*) 74, cheetah (*Acinonyx jubatus*) 15, spotted hyena (*Crocuta crocuta*) 66, and African wild dog (*Lycaon pictus*) 23.

### Data analysis

To test for changes in browse availability over time, a regression that was initially undertaken of the overall stem density per square kilometer was first done. Analyses were then repeated on the four different height classes to determine the temporal trends in woody density and structure. To investigate whether a coppicing effect could be found, we tested if there was a significant increase in the mean number of stems per woody individual over time. We tested whether temporal changes in tree densities in different height classes were related to differences in elephant densities, using a multiple linear regression with woody density as the dependent variable and both elephant density and annual precipitation as independent variables.

To test whether the increasing elephant densities was correlated with a shift in the total browse and grass consumption of the entire herbivore assemblage, we calculated the combined annual food intake (DM, browse and grass) as  $6.0M-0.191*365*N$  (Owen-Smith 1988), where  $M$  is the species' body mass (Table 1), and  $N$  the number of animals per species. The percentages of browse and grass in the diet for each of the species were obtained from literature estimates (Table 1). To analyze the effect of elephants on the total proportion of browse and grass in the diet of entire herbivore assemblage, a regression was carried out with the proportion as the dependent variable and elephant density as the independent variable. This analysis was done without including the food consumption by elephants themselves to

**Table 1** The surveyed herbivores in the Sabi Sand Wildtuin-MalaMala complex with the percentage browse in their diet and their average weight (based on Codron et al 2007), and a comparison of total herbivore numbers per species in 1992, 1993, and 2011, and their contribution to the total food consumption in 1993 and 2011 (%Food)

Species	Common name	Feeding style	Average weight (kg)	N 1992	N 1993	N 2011	%Food 1993	%Food 2011
<i>Giraffa camelopardalis</i> <sup>†</sup>	Giraffe	Browser	1010	288	303	269	9.15	3.17
<i>Tragelaphus scriptus</i> *	Bushbuck	Browser	30	70	59	132	<0.01	<0.01
<i>Sylvicapra grimmia</i> *	Common duiker	Browser	19	na	103	49	0.12	0.02
<i>Tragelaphus strepsiceros</i> *	Greater kudu	Browser	140	573	862	901	5.26	2.15
<i>Raphicerus campestris</i> *	Steenbok	Browser	10	na	48	42	0.03	0.01
<i>Loxodonta africana</i> <sup>†</sup>	African elephant	Mixed feeder	3750	73	60	1260	5.19	42.57
<i>Tragelaphus angasii</i> *	Nyala	Mixed feeder	73	157	171	418	0.62	0.59
<i>Aepyceros melampus</i> *	Common impala	Mixed feeder	41	14411	17776	10405	40.17	9.18
<i>Syncerus caffer</i> <sup>†</sup>	Buffalo	Grazer	585	45	712	3464	13.82	26.25
<i>Hippopotamus amphibius</i> <sup>†</sup>	Hippo	Grazer	1406	160	31	320	1.22	4.93
<i>Ceratotherium simum</i> <sup>†</sup>	White rhino	Grazer	1850	471	152	293	7.49	5.64
<i>Phacochoerus africanus</i> *	Warthog	Grazer	70	390	437	326	1.52	0.44
<i>Kobus ellipsiprymnus</i> *	Waterbuck	Grazer	220	753	388	439	3.41	1.51
<i>Equus quagga burchellii</i> *	Zebra	Grazer	320	927	816	627	7.05	2.11
<i>Connochaetes taurinus</i> *	Wildebeest	Grazer	215	882	571	423	4.93	1.43

\*Mesoherbivore

†Megaherbivore

better analyze the impact of elephant on the other grazer and browser species in the community.

The effect of elephant density and tree density on the density of the various herbivore species were also analyzed using regression. Changes over time or elephant impact effects might deviate from a linear relationship; so, linear regression, regression on log-transformed data, and exponential regressions were used, and the model with the highest  $R^2$ -adjusted and lowest AIC was selected. Residuals were tested for normality and analyzed for homoscedasticity.

As elephant and tree densities were highly collinear, we were unable to build a multiple regression model that included both elephant and tree densities as predictors to explain the changes of the other browser and grazer species. We therefore carried out a hierarchical partitioning, which is able to calculate the part of variation that is uniquely and in combination with other variables explained by a single explanatory variable (Chevan and Sutherland 1991; MacNally and Walsh 2004). The hierarchical partitioning was carried out for each of the herbivore species separately, using rainfall, year, tree density, and elephant density as predictors, but also included the total biomass consumption by all other grazer and all browser species as two additional predictor variables to represent the competition pressure from these two species groups. These latter two variables were calculated by excluding the species for which the analysis was carried out.

The hierarchical partitioning was carried out in R (R Development Core Team 2013) and the other tests in SPSS (v19).

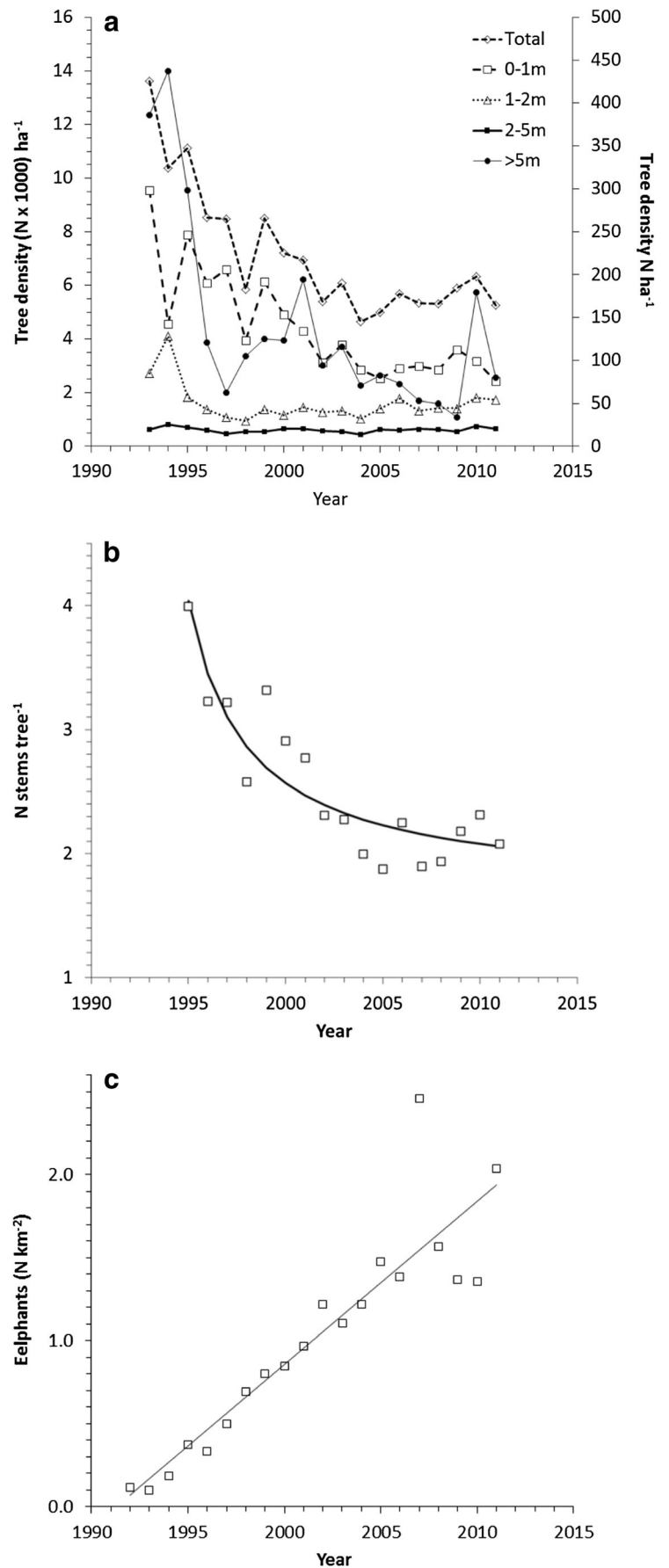
## Results

There was no significant effect of total annual rainfall on the grass standing biomass ( $P>0.05$ ). Because the differences in grass standing biomass were not significantly related to differences in numbers of any of the herbivore species and because it would reduce our analyses considerably, this variable was excluded from further analysis.

### Woody layer

Overall tree density showed a significant decline over time ( $F_{1,18}=100.951$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.84$ ; Fig. 1), and the number of stems per woody individual also declined ( $F_{1,16}=64.715$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.80$ ; Fig. 1). The woody density declined significantly in the 0–1.0, 1.1–2.0, and >5.0 m height classes ( $F_{1,18}>12.416$ ,  $R^2_{\text{adj}}>0.39$ ,  $P<0.003$ ), but no changes were recorded in woody density in the 2.1–5.0 m height class. The density of larger trees (>5.0 m) declined from on average 261 trees/ha in the first 5 years to 79 trees/ha in the last 5 years (Fig. 1).

**Fig. 1** The changes over time in tree density (N/ha) of trees in different height classes: primary y-axis, broken lines: 0–1.0 m, 1.1–2.0 m; secondary y-axis, solid lines: 2.1–5.0 m, >5 m; (a) stems per woody individual (N/tree) and (b) elephant density (N/km<sup>2</sup>), (c) in the Sabi Sand Wildtuin



## Elephants and trees

The elephant density increased over time from 0.12 elephants/km<sup>2</sup> in 1992 to 2.03 elephants/km<sup>2</sup> in 2011 ( $F_{1,18}=88.514$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.82$ , Fig. 1) from the combined effect of natural growth and mortality, and from migration from KNP. Although elephant densities seemed to stabilize in the last 2 years, a linear regression on the untransformed data yielded the best fit, i.e., the lowest AIC. The annual relative growth rate averaged 6.0 % (range 5.3–6.8 %) over the last 5 years of the study. However, this was much larger in the first 5 years of the study, averaging 57 % per year (21.2–140.4 %), due largely to the influx of elephants from Kruger National Park.

The total annual precipitation of the current year alone, the preceding year alone, or the dry season rainfall ( $P>0.10$ ) could not statistically explain the differences in elephant density. Even if we assume that the changes in elephant numbers were caused mainly by the influx of elephants from Kruger National Park in the first years and thus restrict analysis to the data from 2003 onward, none of these three precipitation variables was significant in explaining elephant density.

Density of woody plants declined with increasing elephant density and a forward regression with both elephant density and annual total precipitation as independent variables showed that woody density was negatively correlated with elephant density ( $F_{1,18}=98.550$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=2260.84$ , Fig. 2), but was not correlated with precipitation. Exploring the correlations between woody density in each height class and elephant density, we found that the woody structural classes of 0–1.0, 1.1–2.0, and >5.0 m were all negatively correlated with elephant density (respectively,  $F_{1,18}=43.203$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.70$ ;  $F_{1,18}=13.615$ ,  $P=0.002$ ,  $R^2_{\text{adj}}=0.41$ ;  $F_{1,18}=32.903$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.64$ ), but not the trees between 2.1 and 5.0 m, as the tree density of trees between 2.1 and 5.0 m did not change over time. The number of stems per tree did not increase as was expected under a coppicing response, but in fact decreased with increasing elephant density ( $F_{1,16}=55.093$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.77$ ; Fig. 2). Assuming that there might be a time lag between the effect of elephant and/or the effect of rainfall on the woody density, we also used the elephant density and the annual rainfall in the preceding year as predictors, but this still did not increase the fit of the models.

## Elephants and other herbivore species

In individual regression analyses, none of the browser species showed the expected hump-shaped relationship with highest densities at low to intermediate elephant densities (Fig. 3), i.e., no regression model yielded a significant positive main elephant density term and a negative squared term. Three browser species (duiker, steenbok, and giraffe; Fig. 3, Table 2) and the impala, a mixed feeder, declined with increasing elephant

densities, and also, two grazer species (warthog and zebra) were negatively correlated with increasing elephant densities. However, the three megagrazers in the assemblage, buffalo, hippo, and white rhino also increased after the fence removal and were positively correlated with elephant numbers with a high  $R^2_{\text{adj}}$  (>69 %; Table 2). Nyala and unexpectedly also bushbuck showed a significant positive relation with elephants. As expected, the relationships between the densities of herbivore species and the woody densities showed opposite coefficients to the elephant effect; for instance, the densities of the three megagrazer species were all negatively related to tree density (Table 2).

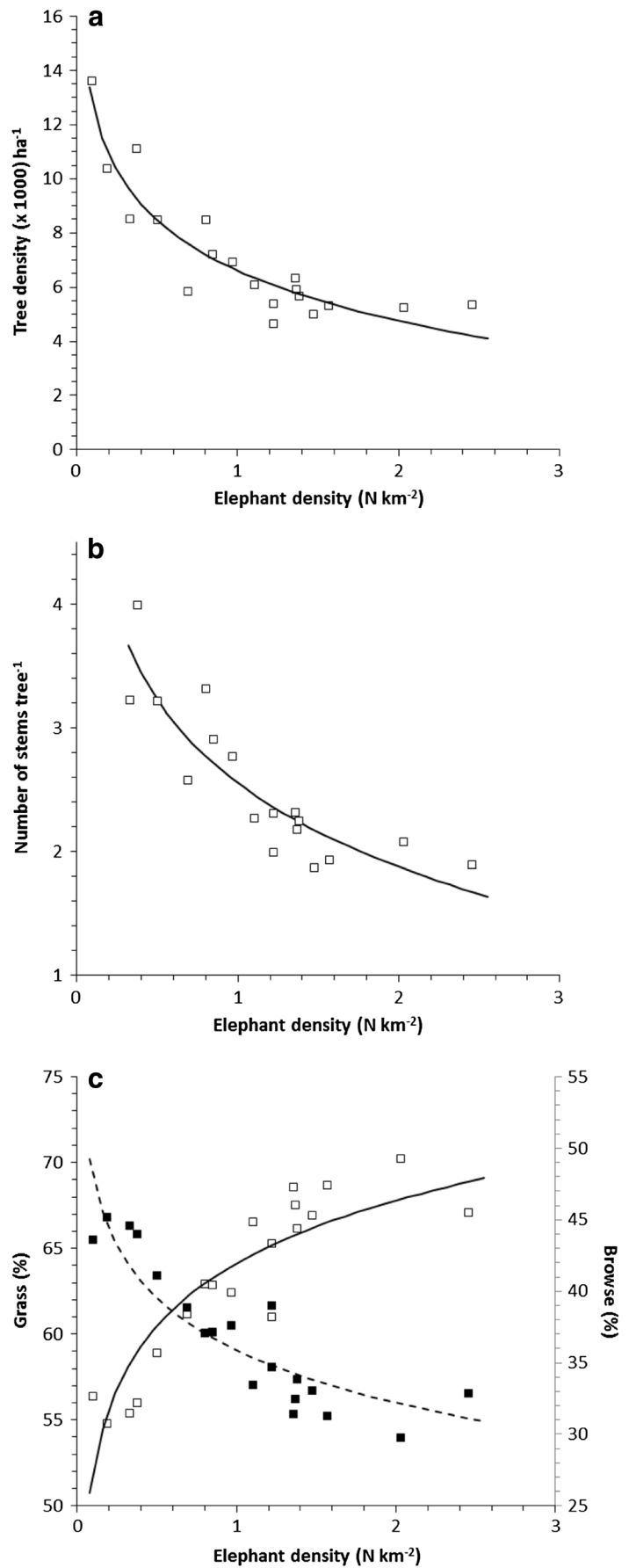
Hierarchical analysis was carried out on each species separately and calculated the percentage of explained variation for each of potential predictor variables and showed that elephant and trees densities were better predictors in explaining the changes in animal densities than year, rainfall, or the competition pressures from other browsing or grazing herbivore species (Fig. 4).

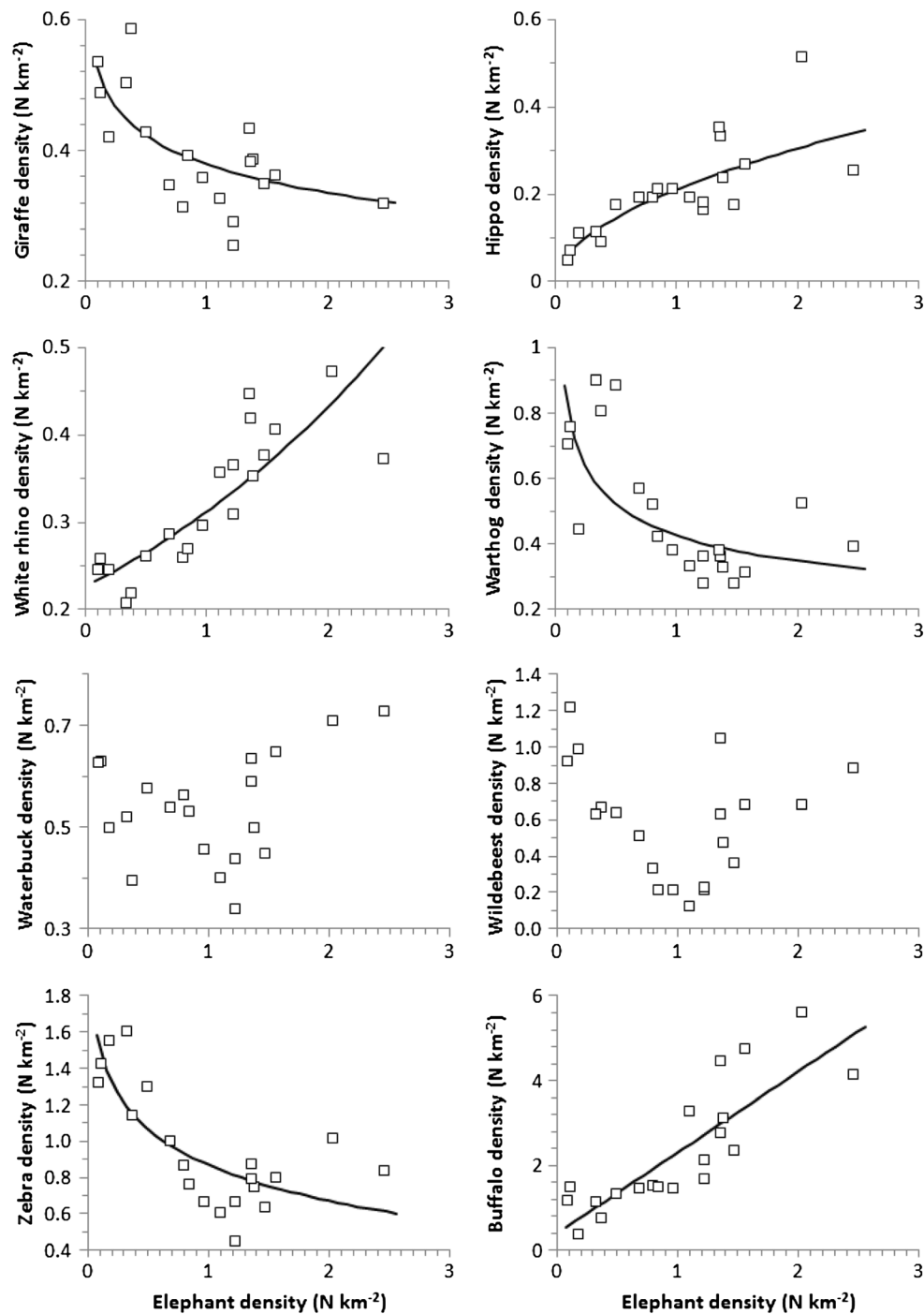
While, overall, total herbivore density decreased by 14 % from 1993 to 2011, the total food consumption by herbivores increased by a factor 2.6, mainly due to the increase of elephants and other megagrazers (Table 1). Surprisingly, the total browse consumption by all herbivore species excluding elephant decreased from 44 to 30 %, and this percentage was strongly negatively correlated with the increasing elephant density (Fig. 2; current year elephant densities:  $F_{1,17}=65.362$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.78$ ; preceding year:  $F_{1,16}=76.733$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.82$ ). This decrease in browse consumption was accompanied by an inverse increase in grass consumption. Elephant density was a better predictor (larger  $R^2_{\text{adj}}$ ) for the decrease in browse consumption than total tree density ( $F_{1,17}=28.059$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.60$ ) or tree density in any of the 4 height classes (0–1.0 m:  $F_{1,17}=29.455$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.61$ ; 1.1–2.0 m: ns; 2.1–5.0 m: ns; >5.0 m:  $F_{1,17}=13.598$ ,  $P<0.002$ ,  $R^2_{\text{adj}}=0.41$ ). However, there may also be a time lag in reaction, and indeed, the relationship with tree density improved considerably if the tree densities of the preceding year was used as the predictor variable in explaining the decrease in browse consumption (overall:  $F_{1,16}=50.681$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.75$ ; 0–1.0 m:  $F_{1,16}=36.506$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.68$ ; 1.1–2.0 m:  $F_{1,16}=5.830$ ,  $P=0.028$ ,  $R^2_{\text{adj}}=0.22$ ; 2.1–5.0 m: ns; >5.0 m:  $F_{1,16}=19.807$ ,  $P<0.001$ ,  $R^2_{\text{adj}}=0.53$ ). However, comparing the AICs of the different models, the elephant density in the preceding year (AIC=27.786) explained the decrease in browse consumption better than any of the other models ( $\Delta\text{AIC}>5$ ).

## Discussion

The question whether the relationship between elephants and other herbivores is facilitative or competitive is highly debated

**Fig. 2** Relationship between the elephant density ( $N/km^2$ ) and the overall tree stem density ( $N/km^2$ ) (a), the number of stems per tree (b) and the estimated total consumption of browse (*filled squares*) and grass (*open squares*) of the relevant herbivores depicted as a percentage of the total consumption excluding elephants in Sabi Sand Wildtuin-MalaMala complex (c)





**Fig. 3** The relationships between elephant densities in the Sabi Sand Wildtuin-MalaMala complex and the densities of other herbivore species ( $N/km^2$ ). Lines indicates best fit of significant models (Table 2)

(Fritz et al. 2002; Kohi et al. 2011; Valeix et al. 2011; Kohi 2013). The impact of elephant may be site- or context-dependent (Guldmond and Van Aarde 2008) and varies spatially within KNP (Eckhardt et al. 2000). Elephant numbers in the Sabi Sand Wildtuin-MalaMala complex were strongly negatively correlated with densities of both small and large trees. However, these correlations were not accompanied by a large coppicing response as predicted by some studies (Kohi

et al. 2011; Smallie and O'Connor 2000), as the number of stems per tree as well as the number of trees  $<1$  m both decreased as tree damage increased. Consequently, the smaller browser species did not benefit from a coppicing response triggered by the large elephant browsing pressure. Interestingly, many previously published studies that suggest a facilitative effect of elephants on mesobrowsers base their findings on a specific (group of) woody species (Lewis 1991;



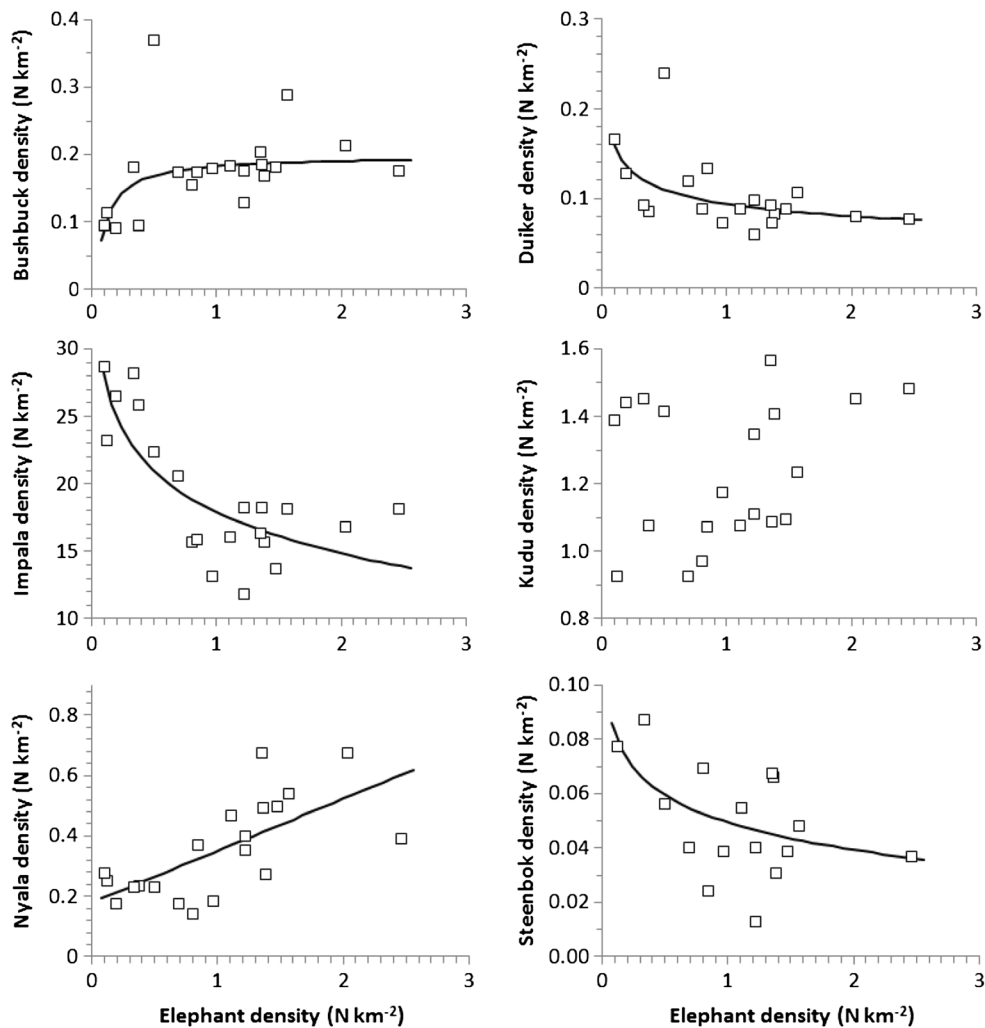


Fig. 3 (continued)

Smallie and O'Connor 2000; Rutina et al. 2005; Makhabu et al. 2006; Kohi et al. 2011). Kerley et al. (2008) suggested that the facilitative effects of elephants largely depend on the growth characteristics and other traits of the utilized species. Hence, the vegetation composition might play a pivotal role in defining the effect of elephants (Levick and Rogers 2008). This might also explain why in some areas, elephants seem to have a facilitative effect whereas in other areas, they show a competitive effect. The decline in the proportion of browsers in Sabi Sand Wildtuin-MalaMala complex might therefore be mediated by the granite lowveld and the *Combretum* spp. vegetation in the area, in which the common woody vegetation does not reach stem densities as high as the stem density in mopane woodlands after being utilized by elephants (Witkowski and O'Connor 1996). The relatively lower stem density and survival rate after elephant utilization probably lead to more open woodlands in the granite lowveld.

The negative correlation between elephant numbers and tree densities in most height classes in our study area, and in other areas (Cumming et al. 1997; Whyte et al. 2003; Shannon

et al. 2008; Woolley et al. 2011), together with the population changes of the browser and grazer species, indicate in this instance that elephant might compete with other browsing species and facilitate grazers, especially megagrazer species, i.e., hippo, white rhino, and buffalo, through a decrease in tree cover and an increase in grass. Also, the proportion of browse consumed by other herbivores, aside from elephant, decreased over time. Therefore, our results suggest that with increasing elephant densities, the herbivore community shifts toward one dominated by grazers and with a smaller contribution of browsers. Hence, successful elephant conservation (an increase in elephant numbers) or poaching (strong decrease in numbers; de Boer et al. 2013) could trigger cascading community effects.

The number of elephant in the area increased 17-fold from 1992 to 2011 (Table 1), whereas the total animal density decreased slightly from 36 to 31 animals/km<sup>2</sup> from 1993, when the fences were dropped, to 2011. Some species did not benefit from the removal of fences between the Kruger National Park and Sabi Sand Wildtuin-Mala Mala complex, and some

**Table 2** Regression results for each of the herbivore species with elephants and tree density, respectively, as the single predictor variable (d.f., sign of the regression coefficient,  $R^2_{adj}$ ,  $F$ , and  $P$  value)

Common name	Elephant					Trees					
	d.f.	+/-	$R^2_{adj}$	$F$	$P$	d.f.	Class	+/-	$R^2_{adj}$	$F$	$P$
African elephant		na				1,17	all	-	0.84	98.550	<0.001
Bushbuck	1,17	+	0.35	11.024	0.004	1,17	>5	-	0.55	22.563	<0.001
Common duiker	1,18	-	0.30	8.603	0.009	1,17	<1	+	0.24	6.524	0.021
Common impala	1,18	-	0.63	32.238	<0.001	1,17	all	+	0.57	24.649	<0.001
Greater kudu	1,18	ns					ns				
Nyala	1,18	+	0.43	15.165	0.001	1,17	<1	-	0.45	15.553	0.001
Steenbok	1,14	-	0.23	5.419	0.035	1,14	<1	+	0.23	5.448	0.035
Giraffe	1,18	-	0.43	15.317	0.001	1,17	all	+	0.37	11.353	0.004
Buffalo	1,18	+	0.69	42.312	<0.001	1,17	<1	-	0.52	20.491	<0.001
Hippo	1,18	+	0.78	69.644	<0.001	1,17	all	-	0.63	31.151	<0.001
White rhino	1,18	+	0.71	46.431	<0.001	1,17	<1	-	0.71	45.755	<0.001
Warthog	1,18	-	0.46	16.835	0.001	1,17	<1	+	0.59	26.825	<0.001
Waterbuck	1,18	ns				1,17	ns				
Wildebeest		na					na				
Zebra	1,18	-	0.60	28.963	<0.001	1,17	all	+	0.54	21.937	<0.001

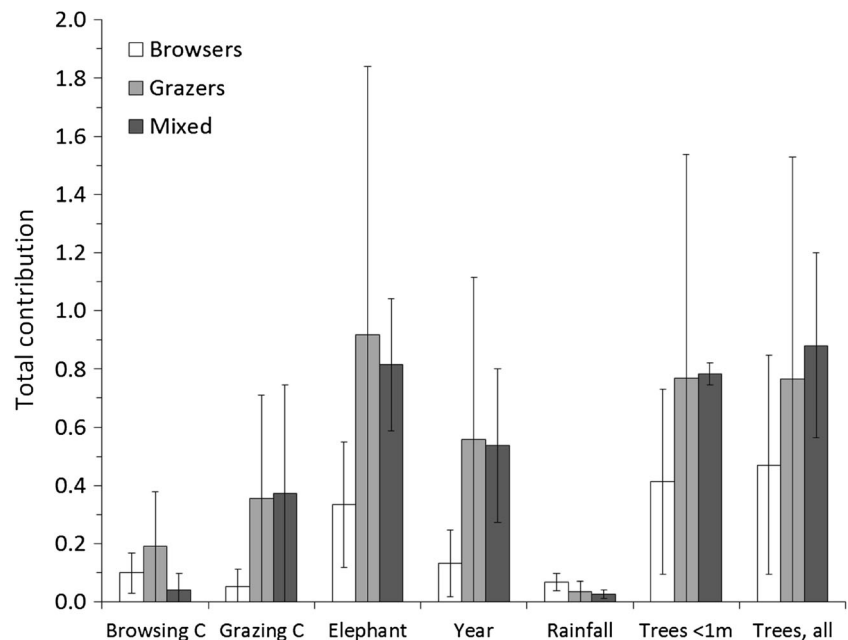
There were three trees classes used as predictors (all: all trees; < 1: 0–1 m; > 5: >5 m); only the regression results with the highest  $R^2_{adj}$  are reported here *na* not applicable, *ns* not significant

species such as duiker, steenbok and giraffe (browsers), and zebra and warthog (grazers) even declined. However, the total metabolic mass of browsers (excluding elephants) increased by a factor of 1.2 between 1992 and 2011, whereas the total mass of grazers increased by 1.7. Thus, while the actual number of animals decreased, all guilds apparently benefitted in terms of total body mass from the removal of the fences. This increase in consumption is particularly caused by the increase

of megaherbivores, since the mesoherbivore densities declined or showed relatively small increases. The grazers benefitted proportionately more from the fence removal and elephant increase, and the community therefore shifted toward a more elephant- and megagrazer-dominated system.

In our correlative approach, we are unable to determine the cause-effect relationships, and changes in elephant densities may covary with other variables. For instance, the decrease in

**Fig. 4** The results of the hierarchical partitioning as depicted by the total contribution (independent and its conjoint contribution  $\pm$ SD) of each of the predictors in a set of regression equations as averaged for all browser ( $N=5$ ; giraffe, steenbok, duiker, bushbuck, kudu), grazer ( $N=6$ ; white rhino, buffalo, hippo, warthog, zebra, waterbuck), and mixed feeder species ( $N=2$ ; nyala, impala), excluding elephants. The used predictor variables were elephant density, year, rainfall, tree density (all trees and only <1 m), and the total biomass consumption by all other browser and grazers species (Browser C, Grazer C) to represent the competition pressure from these two species groups



stem density and the removal of the larger trees in particular might not only increase the frequency and intensity of fires (Govender et al. 2006), but also decrease the facilitative role of trees on grasses by decreasing the role of large trees as nutrient pumps (Treydte et al. 2007). This could be one of the mechanisms that stimulate growth of nutrient poor grasses that can only be used by megagrazers and are of too poor a quality for mesograzers (Olff et al. 2002). Fires certainly play a big role in structuring African savannas (Higgins et al. 2000; Govender et al. 2006) and contribute to the decline in small trees through fire-related mortality. So, the underlying mechanisms explaining the differences in the herbivore community might, besides the direct competition for food or the dispersal after the removal of fences, also be explained by the indirect effects of elephants on the grass layer which indirectly influences the fire regime and food quality.

The changes in herbivore species densities in relation to increasing elephant and decreasing tree densities differed between species. Skarpe et al. (2004) has shown that structural changes due to elephants benefited certain browsing species while others showed a decline in density. The opening up of vegetation by elephants seemed especially beneficial to the megagrazers, with buffalo, hippo, and white rhino numbers being strongly positively correlated with elephant density. Zebras showed a negative correlation with elephant density, suggesting the possibility of resource competition (Young et al. 2005). Bulk feeders do not usually compete with mesograzers such as zebra (Sinclair and Norton-Griffiths 1982), but interspecific competition between mega- and mesograzers could occur during the dry season (Arsenault and Owen-Smith 2002). The differences in response between mega- and mesograzers might also be mediated by fire and a decrease in forage quality, being better tolerated by mega- rather than by mesoherbivores such as zebra. A species that showed a strong negative relation with elephant density and a positive relation with tree density was the giraffe. The decline in giraffe observed in this study was likely to be due to the fact that an increase in elephant numbers was associated with a decline in tall trees. Fritz et al. (2002) suggested that giraffes in particular could suffer from increasing elephant densities as the amount of vegetation at their preferred feeding height decreased, although, as with zebra, predation could also be important in explaining their decline.

In this study, impala density was negatively correlated to the increasing number of elephant. According to Fritz et al. (2002) and Valeix et al. (2008), elephants may compete with mesomixed feeders (such as impala), depending on the abundance and quality of the forage. However, Sinclair et al. (2003) showed that predation is a factor defining herbivore densities on African savannas, particularly for small herbivores (<150 kg). The change in vegetation structure could affect the predation risk of a species; the opening up of the landscape increases the visibility and is expected to decrease

the predation risk (Valeix et al. 2011). The decrease in impala densities in our study is not easily explained by increased visibility as a consequence of the lower tree density, but predation could play a role here through a larger success of cursorial predators, such as the cheetah, in a more open landscape. So, whether the negative relationship between impala and elephant densities is due to direct resource competition with elephants, competition with other herbivore species, changes in predation pressure or other factors, needs to be studied in more detail. Some studies have reported a negative effect of impala on tree recruitment (O’Kane et al. 2012; Prins and Van Der Jeugd 1993) through their impact on seed and seedling survival. However, tree density of small trees (<1 m) was positively correlated with impala density in our study ( $F_{1,17}=17.197$ ,  $P=0.001$ ,  $R^2_{\text{adj}}=0.47$ ).

Elephant populations show large positive and negative changes worldwide as a consequence of successful conservation efforts or increasing poaching activities (de Boer et al. 2013). Our results suggest that these changes can trigger cascading community effects. These effects may not always have been noticed by park managers as they typically take a long period before they become apparent. Future efforts to understand the effects of elephants on herbivores should therefore target and compare specific areas to better understand the impact that elephant have on savanna systems, and the causal mechanisms behind observed changes.

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#### Compliance with ethical standards

**Conflict of interest** No potential conflicts of interest have been identified.

**Human and animal rights and informed consent** This study was carried out without handling animals, using existing survey data.

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