




The Role of Forest Elephants in Shaping Tropical Forest–Savanna Coexistence

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ABSTRACT

Forest edges that border savanna are dynamic features of tropical landscapes. Although the role of fire in determining edge dynamics has been relatively well explored, the role of mega-herbivores, specifically elephants, has not received as much attention. We investigated the role of forest elephants in shaping forest edges of the forest–savanna mosaic in Lopé National Park, Gabon. Using forty camera traps, we collected 1.2 million images between May 2016 and June 2017. These images were classified by over 10,000 volunteers

through an online citizen science platform. These data were combined with a 33-year phenology dataset on elephant-favoured fruiting tree species, and field measurements of elephant browsing preferences and damage. Our results showed a strong relationship between forest elephant density at the forest edge and fruit availability. When fruit availability was high, elephant density at the edge reached values nearly double the highest densities ever reported in any other part of the landscape (7.5 elephants km⁻² in this study vs the previous highest estimate of 4 elephants km⁻²). The highest elephant densities occurred at the end of the dry season, but even outside of this high density period elephant density at the forest edge (2.4 elephants km⁻²) was more than double what other studies estimate for forest interiors with low human hunting pressure (1 elephant km⁻²). We found forest elephants to be selective browsers, but their browsing was non-destructive (in contrast to savanna elephants) and had little effect on tree size demography. Elephant paths acted as firebreaks during savanna burning, making them inadvertent protectors of the fire-sensitive forest and contributing to the stabilising feedbacks that

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allow forest and savanna to coexist in tropical landscapes.

Key words: ecosystem engineer; alternative stable states; forest elephants; *Loxodonta cyclotis*; fire; forest–savanna mosaic; Gabon.

HIGHLIGHTS

- Mean forest elephant density at the forest edge is higher than in most of the forest interior areas.
- The forest edge is an important food resource for forest elephants, especially during the dry season.
- Elephant paths at the forest edge act as fire-breaks, protecting the interior forest from burning.

INTRODUCTION

Elephants are the largest living mega-herbivore and have significant impacts on the ecosystems they inhabit. For example, elephants are part of many obligate seed dispersal mutualisms and therefore affect tree recruitment patterns (Campos-Arceiz and Blake 2011).

Elephants also eat and trample established tree seedlings, which impacts species composition and vegetation structure on the local level and ecosystem services such as carbon accumulation and storage at a global scale (Poulsen and others 2013; Terborgh and others 2016). African elephants form two genetically and ecologically distinct groups (Rohland and others 2010; Maisels and others 2013): the savanna elephant, *Loxodonta africana* (Blumenbach, 1797) and the physically smaller forest elephant, *L. cyclotis* (Matschie, 1900). Although both African elephant species have faced drastic population declines, recently the most severe of these has been experienced by the forest elephants of Central Africa (Maisels and others 2013; Poulsen and others 2017). What this loss means for ecosystem functionality is not well understood, as relatively little is known about forest elephants relative to savanna elephants (Poulsen and others 2018). A better understanding of forest elephant ecology could galvanise greater support for their urgent protection.

As their name suggests, forest elephants are a forest-dwelling species, with their current range mostly enclosed within the rainforests of Central

Africa (Maisels and others 2013). However, forest elephant density varies significantly across this range (White 1994a). In the absence of human pressure, forest elephant density is largely determined by forage availability (Blake 2002). Fruit is a large component of the forest elephant's diet (White and others 1993), but because fruit availability can be strongly seasonal (White 1994b) there are times when forest elephants may rely more heavily on browse to fulfil their nutritional needs. Accessible browse can be scarce in the understorey of the forest interior, leading elephants to leave the forest interior to seek alternative foraging sites (Blake 2002).

The forest edge, considered in this study to be where closed-canopy forest transitions into open-canopy savanna, may be an important alternative foraging site for forest elephants. The forest edge not only has an abundance of accessible browse, but is also home to multiple elephant-favoured fruiting species (White and others 1993) that may have complementary phenology to the fruit trees in the forest interior. At the end of the long dry season (June–September) in our study site, fruit availability is low in the forest interior, the proportion of fruit in elephant dung is at a minimum, and forest elephants are often “visibly emaciated with only skin on their bones” (White 1992; White and Abernethy 1997; Momont 2007) (Figure S1). It is in this season that we expect forest elephants to preferentially turn to the forest edge as an alternative foraging site. Although forest elephants have been anecdotally observed at forest edges (Momont 2007; Oliveras and Malhi 2016), their use of this specific habitat type and how this use varies through time is yet to be quantified.

If forest elephants are using forest edges as a forage resource, it is likely they are affecting edge dynamics. Forest elephants are known to trample seedlings and snap saplings in the forest interior (Terborgh and others 2016; Rosin and others 2017), whereas savanna elephants target medium-sized trees and cause widespread mortality within savanna (Morrison and others 2015). If forest elephants act in a similar way at the forest edge, then edge contraction may occur. Conversely, if forest elephants disperse seeds and promote tree recruitment at the forest edge (Blake and others 2009), as observed in the forest interior, edge expansion may occur. Forest elephants may also have an indirect stabilising effect on the forest edge, via their interactions with fire. For example, if forest elephants at the forest edge reduce fuel load by eating grass, and decrease fuel continuity by repeatedly trampling paths, they may create firebreaks. These

firebreaks would prevent fire from spreading from frequently burnt and fire-dependent savanna into the fire-sensitive forest (Hoffmann and others 2003; Mouillot and Field 2005).

How forest elephants affect forest edge dynamics could either mitigate or exacerbate forest encroachment. Forest encroachment, or the loss of savanna area due to expanding forest, is a pressing environmental crisis in tropical ecosystems and is occurring across sub-Saharan Africa (Mitchard and others 2009; Stevens and others 2017). Forest encroachment leads to losses in savanna biodiversity and savanna ecosystem functionality (Parr and others 2014), and preventing it is often a key management objective within protected areas. It has been suggested that savanna elephants could prevent forest encroachment by maintaining or even creating more open landscapes (Asner and Levick 2012). However, whether forest elephants can play a similar structuring role to savanna elephants has not yet been investigated. Gabon, in Central Africa, is a stronghold for forest elephant conservation and is home to most of the remaining global population (Maisels and others 2013), making it the ideal location for such an investigation.

The objective of this study was to quantify forest elephant use of the forest edge, assess the drivers of this use and determine the consequences of this use for forest edge dynamics. We unobtrusively monitored forest elephants at forest edges at high space and time resolution using 40 motion- and heat-triggered camera traps. We combined these camera trap data with field measurements of fruit phenology, vegetation structure, and elephant browsing preferences and damage to address the following research questions and hypotheses:

(Q1) Is the forest edge an important forage resource for forest elephants?

H_{1.1}: Forest elephant density at the forest edge will be higher, or seasonally higher, than previous density estimates made in the forest interior (White 1994a).

H_{1.2}: Elephant use of the forest edge has distinct seasonal patterns that correlate with seasonal patterns in fruit and browse availability at the forest edge versus in the forest interior.

(Q2) Which species do forest elephants prefer to browse at the forest edge, and is this reflected in these species' size demography?

H_{2.1}: Forest elephants are selective browsers at the forest edge, showing preferences for certain species and avoiding others.

H_{2.2}: Forest elephants will prefer to browse on smaller size class stems at the forest edge; therefore, the species on which they prefer to browse (H_{2.1}) will have recruited fewer stems into larger size classes compared to species that they avoid browsing on.

(Q3) Is forest elephant activity at the forest edge important for preventing savanna fires from burning into the forest?

H_{3.1}: Forest elephants' paths at the forest edge that act as firebreaks during savanna burning are more heavily used and have less grass around them than paths that do not act as firebreaks.

METHODS

Study Site

Our study site was the forest–savanna mosaic in the northern part of Lopé National Park, Gabon, in central Africa. Here, Middle Ogooué savannas (van de Weghe 2011) interlock with continuous Lower Guineo-Congolian rainforest (White 1983), with sharp transitions or ecotones, between them (Figure S2a). An elephant path is nearly always observed running along the edge of the savanna, parallel to the forest edge (Figure S2b).

Mean annual temperature in LNP is 24 °C (2003–2016) and mean annual humidity is 81% (2003–2016) (Tutin and others 2016). Mean annual rainfall is 1442 mm, with mean monthly rainfall ranging between less than 2 mm in the dry season to more than 150 mm in the wet season (1984–2016) (Tutin and others 2016). LNP has a short (mid-December to mid-February) and a long (mid-June to mid-September) dry season, with prescribed burning occurring in the latter. Savanna burning has been used as a habitat management tool since 1993 (Jeffery and others 2014) and has been present in the landscape for at least the last 5000 years (White 2001).

Forest Elephant Density

We estimated forest elephant density at the forest edge at forty sites using motion- and heat-triggered camera traps (Agressor No Glow, Bushnell, Kansas, USA) (Figure 1). At each site, we attached a camera to a tree at 1–1.5 m above ground. The camera focussed perpendicularly towards the elephant path, so as to be the most likely to capture a passing elephant while also not being in the elephant's way and inciting them to destroy the camera.

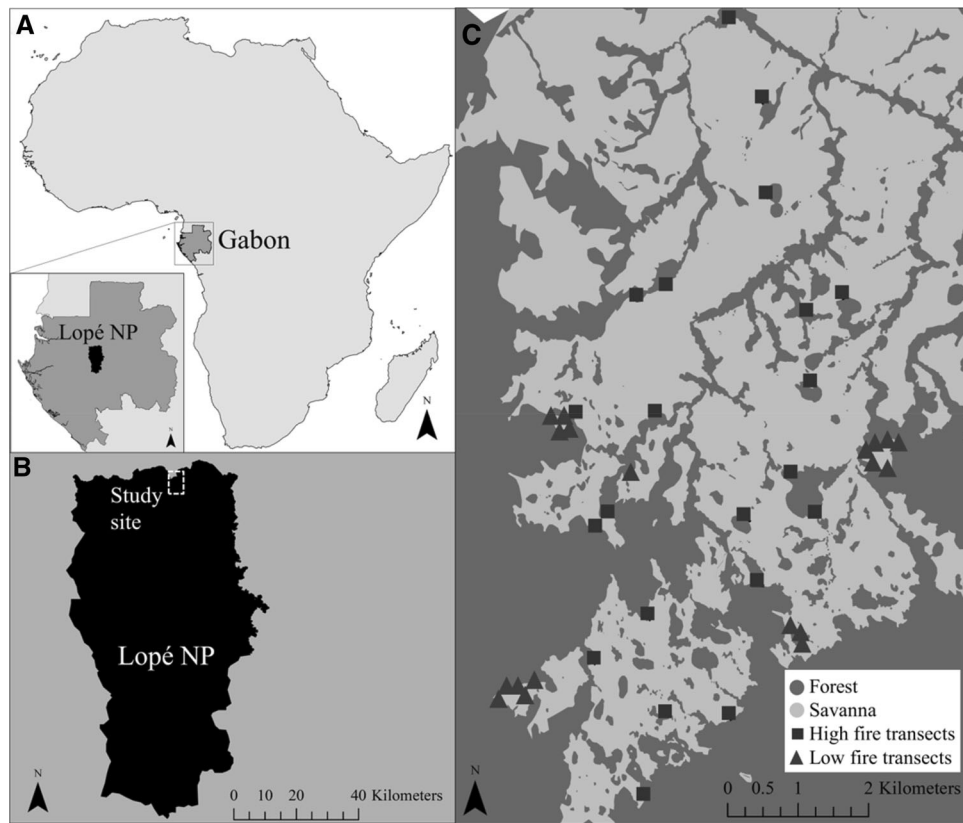


Figure 1. Map showing the location of **A** Lopé National Park in Gabon, **B** the study area within the park and **C** the 40 camera trap locations across high and low fire frequency sites within the forest–savanna mosaic (Agence Nationale des Parcs Nationaux 2006).

Each camera trap could be triggered up to once per second and was monitored from May 2016 to June 2017, resulting in 1.2 million images being collected. Manual analysis by a small research team was unfeasible. Therefore, we created an interactive web interface (“Elephant Expedition”, created in collaboration with Zooniverse.org, available at <https://www.zooniverse.org/projects/anabellecardoso/elephant-expedition>, Figure S3), through which more than 10,000 interested members of the public (citizen scientists) classified our images. Each image was classified based on what animal species was in the image, and how many forest elephants were in the image. Images containing animals were each classified by ten volunteers; thus, each image was associated with up to ten potentially conflicting classifications. To assign each image, a single aggregated classification we applied a plurality algorithm (Swanson and others 2015, script available at https://github.com/zooniverse/Data-digging/tree/master/example_scripts/R_code/survey-task-generalized). We assigned the animal species in the image as the one with the highest number of “votes”, and the number of elephants in the image

as the median count of elephants. Considering that (1) elephants are readily distinguishable from other animals in the study site, (2) images with animals were classified by ten volunteers, (3) the high trigger frequency resulted in multiple images usually being captured during the same elephant visit, we have a high level of confidence in the data obtained from the citizen science classifications (Kosmala and others 2016).

To obtain forest elephant density estimates for each site, we applied a random encounter model (REM) (Rowcliffe and others 2008; Cusack and others 2015) using the formula:

$$D = \left[\left(\frac{y}{t} \cdot \frac{\pi}{v \cdot r(2 + \theta)} \right) \right] * \text{mean group size.}$$

Here D is the density of elephants (elephants km^{-2}). $\frac{y}{t}$ is the trapping rate, or total number of independent capture events (y) divided by the total camera effort (t , h). Forest elephants move in cohesive groups and the camera trap can be triggered up to once per second. To avoid counting the same elephant twice we looked for independent

elephant group capture events. Independent elephant group capture events were peaks in elephant counts separated from other peaks by at least 1800 s before and after the peak (`find_peaks` function, available at https://github.com/stas-g/finDPeaks/blob/master/find_peaks.R). Elephant density was then obtained by multiplying the density of independent group capture events by the mean size of the elephant group (Rowcliffe and others 2008). *Mean group size* was the sum of peaks in elephant count of all independent group capture events divided by the number of independent group capture events. This is a conservative estimate of group size, as the maximum count of elephants in any one image is the minimum group size. Total camera effort is the number of hours for which the camera was functional. v is the average speed of a forest elephant (a constant 0.334 km h^{-1} , Mills and others 2018). r and θ are the radius (km) and angle ($^{\circ}$) of the camera's detection zone (Rowcliffe and others 2008). We calculated r and θ by walking in front of the camera detection beam at varying distances and directions and measuring where the trap was triggered. The value applied to r and θ was then the mean value of a subset of the cameras ($n = 17$, mean $r = 0.00664 \text{ km}$ (max = 0.00360 km , min = 0.01050 km); mean $\theta = 28.8^{\circ}$ (max = 40° , min = 20°)).

Browsing Preferences and Damage to Trees

At each site, we assessed the first 15 trees more than 1 m tall and less than 1 m from the elephant path to the right and the left of the camera for elephant damage ($n = 1201$). This assessment was done during the last weeks of camera deployment. We noted tree species and height class (1–2 m tall, 2–3 m tall, 3–4 m tall, more than 4 m tall), as well as degree of browsing damage (based on Wing and Buss 1970; Ishwaran 1983):

0. No sign of damage
1. Side branches of canopy broken
2. Main stem/trunk broken
3. Entire tree uprooted
4. Bark stripped
5. Elephant tusk marks on trunk

We calculated preference ratios (Petrides 1975) to assess which species and size class of tree forest elephants preferred to browse on using the formula:

$$\text{Preference ratio} = \frac{\% \text{ damage}}{\% \text{ availability}}$$

where:

$$\begin{aligned} \% \text{ damage} \\ = 100 * \frac{\text{number of damaged stems in a species or size class}}{\text{number of damaged stems in all species or size classes}} \end{aligned}$$

and:

$$\begin{aligned} \% \text{ availability} \\ = 100 * \frac{\text{number of stems in species or size class}}{\text{number of stems in all species or size classes}} \end{aligned}$$

Preference ratios above 1 indicated a species or size class was preferred by elephants, whereas ratios below 1 or 0 indicated that a species or size class was somewhat preferred or avoided by elephants.

We evaluated differences between observed and expected frequencies of trees in different size classes using χ^2 tests to assess whether elephants were significantly affecting stem size distribution. Expected frequencies were calculated under the assumption that browsing preferences had no effect on stem size class distribution.

Fruit Phenology

Fruiting phenology data have been collected at the study site over thirty-three years (1986–2018; Tutin and others 2018). The species in this dataset were originally chosen for their importance in ape diet (Tutin and Fernandez 1993). Fruiting species important to elephants were identified from detailed studies on forest elephant diet at the study site and were found to be highly overlapping with those of ape diets (White and others 1994). Thus, for thirty-one of the thirty-eight species most important to elephants, we have access to thirty-three years of data. For the remaining seven species, data collection began in 1996 or 2001. Reliable fruiting seasonality can be established with 6 years of data for most species (Bush and others 2017). All species in this study site reproduce annually, except *Psychotria vogeliana*, *Psidium guineensis* and *Antidesma vogelianum*, which flower and fruit every 6 months (Bush and others 2017).

Each month, individual tree crowns were visually monitored and the presence of ripe fruit in the canopy as well as the proportion (0, 0.25, 0.5, 0.75, or 1) of the canopy covered by fruit was recorded (see Bush and others 2018 for details on the method). We classified tree species as forest edge species if they occurred in our assessment of elephant damage at the forest edge (described above) and are described in the literature as found most commonly on edges and rarely in the forest interior (White and Abernethy 1997). We classified tree

species as forest interior species if they did not occur in our elephant damage assessment and were described in the literature as such (White and Abernethy 1997).

Using these data, a fruit availability score (FAS) for the forest edge (FAS_{edge}) and the forest interior (FAS_{interior}) was calculated for each calendar month. The FAS indicates long-term patterns of fruit availability across the landscape. The FAS assumes that the amount of fruit a tree produces scales isometrically with its crown volume, which scales with the square of trunk radius (Shenkin and others, in review). The FAS also assumes that fruit in the canopy is not just present or absent, but that the proportion of the canopy covered in fruit can vary. By adjusting the formula for the “ripe fruit score” (Mitani and others 2002) to account for these assumptions, we calculated the FAS for each habitat type using the formula:

$$\text{FAS} = \sum_{i=1}^n p_i * c_i * d_i * r_i^2$$

Here i is the tree species. p_i is the mean proportion of individual trees in species i that have ripe fruit in their canopy in any given month. c_i is the mean proportion of each fruiting tree canopy that is covered in fruit. d_i is the mean density of trees of in species i in that habitat type (forest edge or forest interior) (stems ha⁻¹). r_i is the mean radius at breast height (cm) of trees in species i . The mean density and mean radius of each species were determined from existing census data for plots in different habitat types across the study area (Table S1).

We calculated fruit availability scores using long-term phenology data (1986–2018) rather than phenology recorded exclusively during the study period (May 2016–June 2017; but see Figure S4 for the comparison between these two datasets). We assume long-term fruit availability patterns will explain elephant density better than those recorded during the study period because we expect elephants to move across the landscape-based largely on memory, as opposed to current smell, sight and sound cues. Previous work supports this assumption, for example savanna elephants use memory to find the closest water hole from as far away as 50 km, a distance that precludes the use of sight, smell or hearing (Polansky and others 2015). Forest elephants have been shown to exhibit similar

abilities in response to *Sacoglottis* fruiting season, during which they are estimated to be attracted to localised fruiting events from a surrounding area as large as 3000 km² (White 1994b).

Elephant Paths as Firebreaks

Twenty-one of the forty camera trap sites were experimentally burnt between July and September 2016 as part of LNP’s normal conservation management protocol (Jeffery and others 2014) (Figure 1). Fires were set in the hottest part of the day to maximise intensity. All burning was conducted following appropriate safety protocols, and details on the fires set can be found in Cardoso and others (2018).

At each site, an elephant path ran along the edge of the savanna, parallel to the forest edge. We graded each path as 1 (lightly used, no clear break in the grass layer but clearly trampled), 2 (intermediately used, some bare ground visible but grass still growing on the path) or 3 (heavily used, bare ground clearly visible and clear break in grass layer). After burning, we noted whether or not the elephant path had acted as a firebreak by examining the burn scar and noting where the fire went out (Figure S2).

We assessed grass biomass around the elephant path using a disc pasture meter calibrated for the site (Cardoso and others 2018). Grass biomass measurements were taken (1) at the forest edge, (2) at 1, 2 and 3 m into the savanna and (3) at 1, 2 and 3 m into the forest. Canopy leaf area index (LAI, canopy leaf area per unit ground area) can determine grass biomass (Hoffmann and others 2012), and so was also measured around the elephant path using hemispherical photographs (Nikkor fish-eye lens 10.5 mm F2.8G AF DX IF ED). Photographs were taken (1) in the savanna, 3 m from the forest edge, (2) at the forest edge and (3) into the forest, 2 m from the forest edge. Photographs were converted into LAI values using post hoc image analysis in Hemisfer (v2.16, Schleppi and others 2007; Thimonier and others 2010).

We tested for significant differences in grass biomass and LAI between paths that had acted as a firebreak and those that had not using non-parametric Kruskal–Wallis tests. Non-parametric tests were used because the data did not fulfil the assumptions (normality and homoscedasticity) of equivalent parametric tests.

We performed all analyses in R (v3.5.0, R Development Core Team 2018). Significance was noted at $p < 0.05$.

RESULTS

Forest Elephant Density at the Forest Edge was Relatively High ($H_{1.1}$) and was Highest When Fruit was Most Abundant Here ($H_{1.2}$)

Mean forest elephant density at the forest edge between May 2016 and June 2017 was 3.1 ± 0.2 elephants km^{-2} (mean \pm standard error) and ranged from 1.3 ± 0.3 elephants km^{-2} (January 2017) to 7.5 ± 1.4 elephants km^{-2} (September 2016) (Figure 2A). Elephant density showed strong seasonal patterns and was highest at the end of the dry season (September to October 2016, 6.7 ± 1.2 elephants km^{-2}) ($H_{1.1}$) (Figure 2A). As expected, even outside of these exceptionally high density months, mean density at the edge remained relatively high (2.4 ± 0.4 elephants km^{-2}) ($H_{1.1}$).

As hypothesised, peaks in forest elephant density coincided with peaks in mean fruit availability at the forest edge, which occurred at the end of the long dry season ($H_{1.2}$) (Figure 2B). Mean forest elephant density at the forest edge had a significant positive relationship with forest edge fruit availability ($H_{1.2}$) (Figure 2C). Contrary to our predictions, no significant links were observed between forest elephant density at the forest edge and fruit availability in the forest interior (Figure 2D). However, one of the lowest mean monthly density estimates of forest elephants at the forest edge (2.2 ± 0.4 elephants km^{-2}) co-occurred (in February 2017) with the forest interior's highest fruit availability score ($H_{1.2}$). Similarly, one of the lowest times for fruit availability in the forest interior co-occurred (in April 2017) with a minor peak in forest elephant density at the forest edge (2.6 ± 0.6 elephants km^{-2}) ($H_{1.2}$).

Forest Elephants are Selective ($H_{2.1}$) but Relatively Non-destructive Browsers, Having No Appreciable Effects on Tree Size Demography ($H_{2.2}$)

Over 1200 individual trees from fifty-four species were sampled for elephant damage at the forest edge. Twenty-five of these species had less than five trees sampled and were consequently excluded from the analysis (211 trees). Of the remaining twenty-nine species (1143 trees), nine showed no elephant damage. Of the twenty species that

showed some elephant damage, eleven were somewhat preferred (preference ratio between 0 and 1) and nine were strongly preferred (preference ratio > 1) by forest elephants (Table 1).

Of the twenty-two species that showed some degree of elephant browsing damage, only six also produced fruit eaten by elephants, and the rest were presumably targeted only for their browse. Of these six species, peaks in elephant density at the forest edge were associated with peaks in fruit availability for *Antidesma vogelianum* and *Sacoglottis gabonensis*, both of which were species strongly preferred for browse by forest elephants (Table 1, Figure S5). Peaks in elephant density were associated to a lesser extent with peaks in fruit availability for the species *Psidium guineense*, *Sarcocephalus latifolius* and *Psychotria vogeliana*, of which the former two were strongly preferred and the latter somewhat preferred by forest elephants.

Less than a fifth (16%) of all stems sampled showed some degree of elephant damage. The majority of damage to trees involved branch breaking, followed by main stem snapping (trunk breaking), with uprooting only observed in three trees, of which only one was taller than 2 m (Figure 3A). Contrary to our predictions that small size trees would experience more damage, forest elephants in this study were found to preferentially damage medium-sized 3–4 m tall trees ($H_{2.2}$) (Figure 3B). Despite this preference, and against our expectations, there were no clear effects of forest elephant browsing preferences in tree size demography, with no significant differences between observed and expected frequencies of trees in each size class or between observed and expected frequencies of trees in each preference category ($H_{2.2}$) (Figure 3C).

The Forest Elephant Paths at the Forest Edge That Acted as Firebreaks Were Those More Heavily Used ($H_{3.1}$)

Of the twenty-one sites that were experimentally burned, sixteen had intermediately (grade 2) and five had heavily (grade 3) used elephant paths. No lightly used paths (grade 1) were present. Ten of these twenty-one elephant paths acted as firebreaks at the edge of the savanna during burning. Heavily used paths were more likely to act as firebreaks than intermediately used ones (80% of the grade 3 paths vs 38% of the grade 2 paths acted as firebreaks) ($H_{3.1}$). Although not statistically significant, mean annual elephant density was marginally higher in sites with heavily used paths (grade 3) than those with intermediately used paths

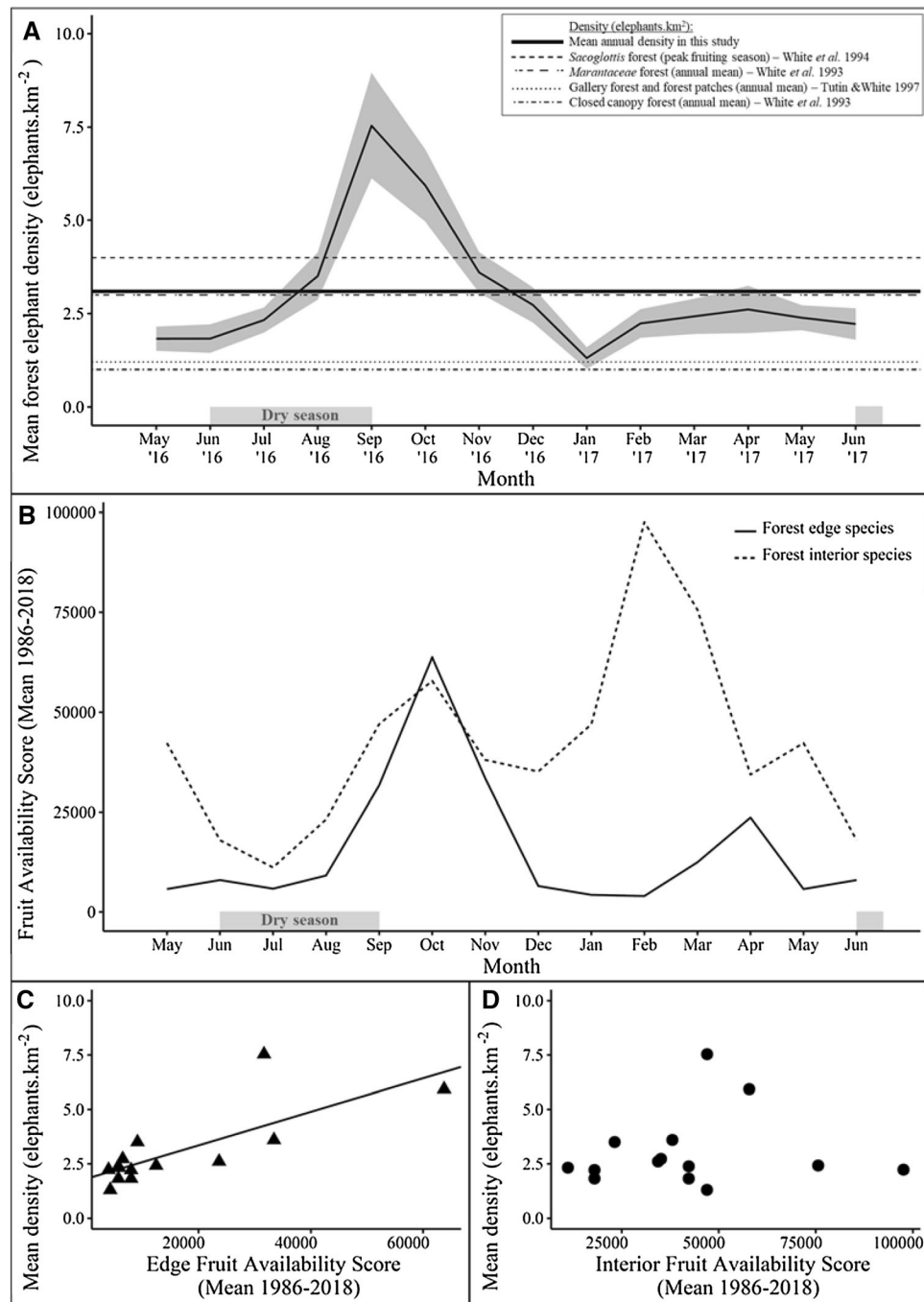


Figure 2. **A** Mean elephant density at the forest edge (May 2016–July 2017) as calculated from camera traps. Shaded area shows standard error around the mean. Horizontal lines show previously estimated forest elephant densities in other vegetation types in the landscape, as well as the mean annual density in this study. Shaded grey blocks show the long dry season that occurs annually. **B** Total fruit availability score (monthly mean from 1986 to 2018) for the forest edge and interior. **C** and **D** Showing the relationship between forest elephant density at the forest edge and fruit availability at the forest edge (**C**) (linear model, slope= 0.000077, intercept= 1.8, adjusted R^2 = 0.55, p < 0.005, df = 12) and the forest interior (**D**) (linear model, slope= 0.000083, intercept= 2.7, adjusted R^2 = - 0.07, p = 0.70, df = 12).

(grade 2) (Kruskal–Wallis, χ^2 = 1.75, df = 1, p = 0.19; mean (\pm standard error) annual elephant: sites with grade 3 path = 4.0 (\pm 0.8) ele-

phants km⁻², sites with grade 2 path = 2.8 (\pm 0.4) elephants km⁻²). Paths that acted as firebreaks had marginally less grass biomass around them than

Table 1. Degree to Which Tree Species At the Forest Edge are Preferred (Preference Ratio > 1), Somewhat Preferred (Preference Ratio < 1) or Avoided (Preference Ratio=0) by Forest Elephants

Species	No. trees sampled	No. trees showing elephant damage	Preference ratio
Strongly preferred			
<i>Holarrhena floribunda</i>	9	4	2.78
<i>Sarcocephalus latifolius</i>	49	18	2.30
<i>Psidium guineense</i>	24	8	2.09
<i>Millettia versicolor</i>	129	38	1.84
<i>Antidesma vogelianum</i>	42	10	1.49
<i>Vitex doniana</i>	13	3	1.44
<i>Crossopteryx febrifuga</i>	106	24	1.42
<i>Sacoglottis gabonensis</i>	84	17	1.27
<i>Paropsia grewoides</i>	18	3	1.04
Somewhat preferred			
<i>Aidia ochroleuca</i>	13	2	0.96
<i>Psychotria vogeliana</i>	208	29	0.87
<i>Xylopia aethiopica</i>	44	6	0.85
<i>Monanthes congolensis</i>	19	2	0.66
<i>Ouratea myrionura</i>	82	8	0.61
<i>Lophira alata</i>	43	4	0.58
<i>Cryptosepalum staudtii</i>	13	1	0.48
<i>Tetracera podotricha</i>	13	1	0.48
<i>Pauridiantha efferata</i>	14	1	0.45
<i>Cnestis ferruginea</i>	54	3	0.35
<i>Barteria fistulosa</i>	53	1	0.12
Avoided			
<i>Anthocleista vogelii</i>	7	0	0
<i>Aucoumea klaineana</i>	11	0	0
<i>Cola lizae</i>	5	0	0
<i>Diospyros dendo</i>	27	0	0
<i>Gomphia flava</i>	11	0	0
<i>Manotes sp.</i>	12	0	0
<i>Maprounea membranacea</i>	21	0	0
<i>Pavetta puberula</i>	5	0	0
<i>Psorospermum tenuifolium</i>	16	0	0

paths that did not act as firebreaks; however, this difference was not statistically significant (Figure 4A) ($H_{3,1}$). This marginal difference in grass biomass was not induced by a higher level of shading, as no significant differences in canopy LAI were found between sites where paths had acted as firebreaks and sites where they had not (Figure 4B).

DISCUSSION

The Forest Edge is an Important Forage Resource for Forest Elephants (Q1)

Mean forest elephant density at the forest edge was exceptionally high (7.5 elephants km^{-2}) at the end of the long dry season (September 2016). At this time, forest elephant density was approximately double the highest estimate ever made in this

landscape (4 elephants km^{-2} in *Sacoglottis gabonensis* forest during fruiting season, White 1994b). Forest elephant density at the forest edge remained relatively high (2.4 elephants km^{-2}) even outside of the exceptionally high density months at the end of the dry season. This relatively high density of elephants is lower than the previously estimated mean annual density in *Marantaceae* and colonising forests (3 elephants km^{-2} , White and others 1993), but is more than double the estimated mean annual density in closed-canopy forests that are relatively unaffected by human hunting pressure (1 elephants km^{-2} , White and others 1993; Maisels and others 2013). Differences in methodology may be responsible for some of the differences between elephant density estimates in this study and estimates made in other studies; however, it is not currently possible to quantify the degree to which this is the case.

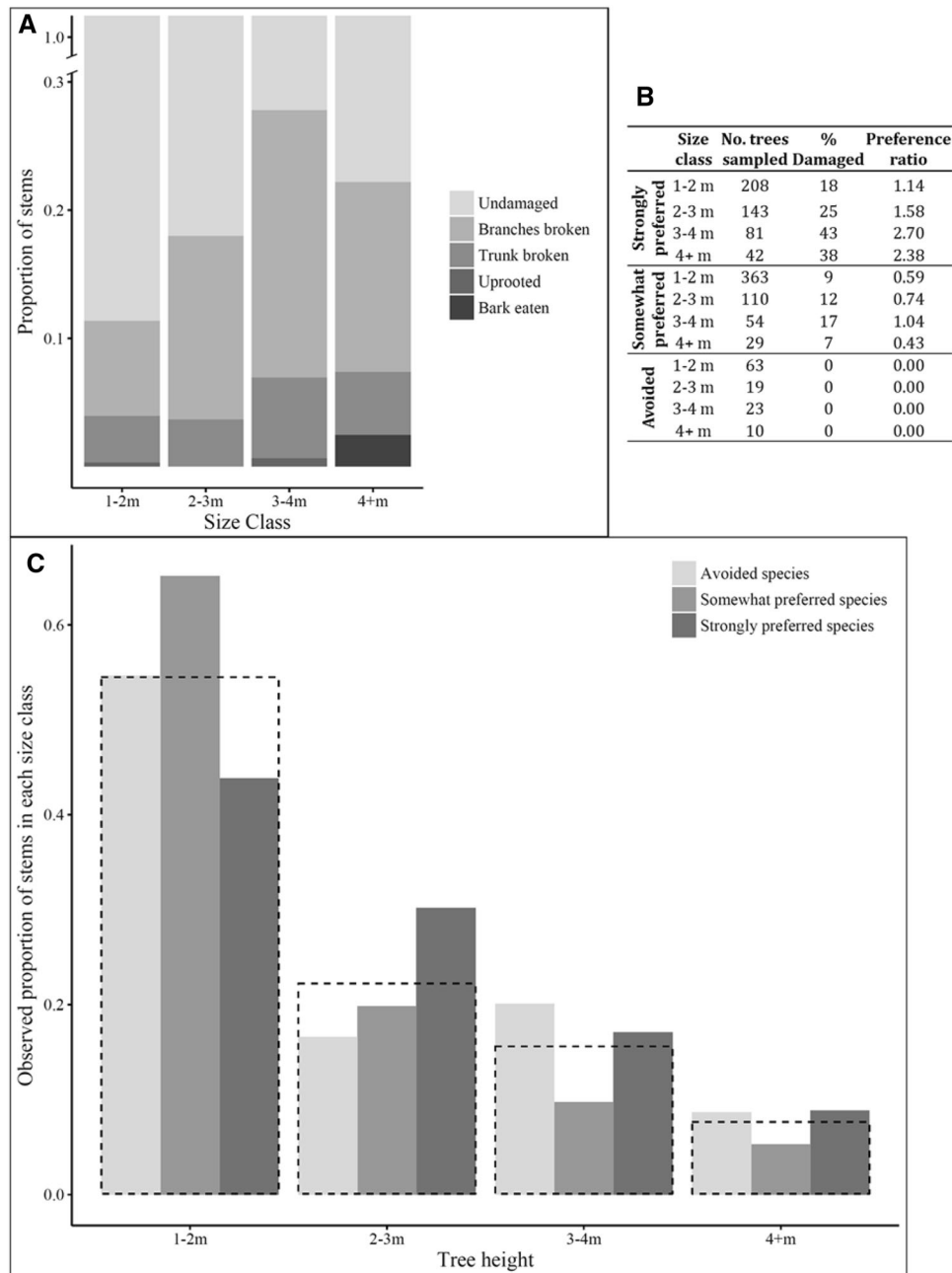


Figure 3. **A** Proportion of trees in each size class incurring different types of forest elephant damage. Note that the y-axis has been compressed for display purposes. **B** Table of which size class trees forest elephants prefer to browse in each preference categories of species. **C** Size distribution of trees for species avoided, somewhat preferred and strongly preferred by forest elephants. No significant differences were found between observed (different shaded bars) and expected (dotted lines) frequencies of stems in each size class, or between frequencies of stems in each preference categories (in χ^2 tests p was always >0.5).

The relatively high density of forest elephants at the forest edge throughout the year provides compelling evidence that the edge is a key resource for these elephants. Due to the abundance of forest edges in forest–savanna mosaics, this finding may also explain why previous work has found forest–

savanna mosaics to be such a heavily utilised part of the forest elephants' range. For example, Momont (2007) showed four collared female forest elephants to prefer the forest–savanna mosaic to the closed-canopy forest. Mills and others (2018) found similar patterns and indicated that conser-

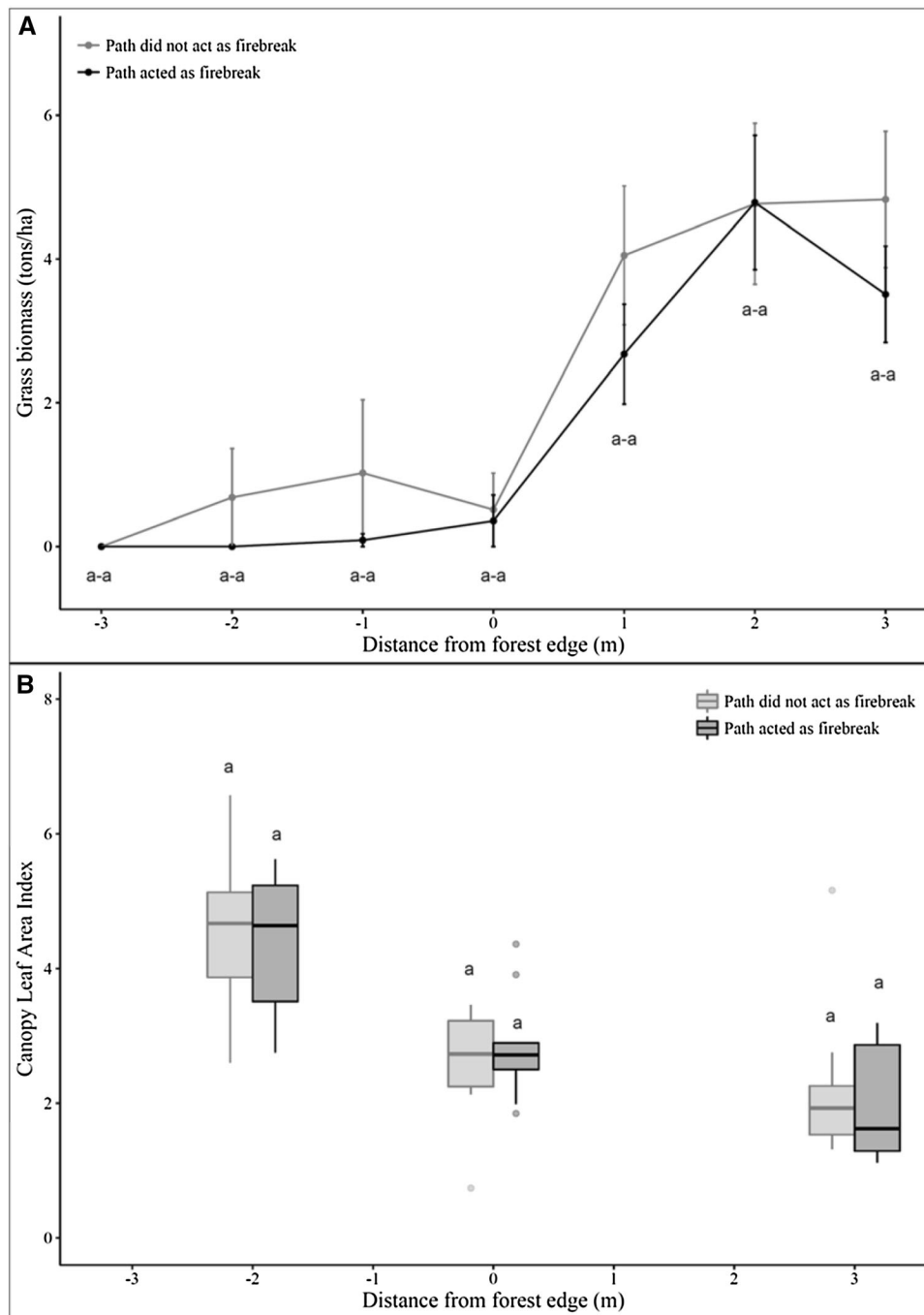


Figure 4. **A** Grass biomass and **B** tree canopy cover (leaf area index, LAI) around the forest elephant path that runs along the edge of the savanna, parallel with the edge of the forest. Positive distances from the forest edge are in the savanna, negative distances in the forest. Different letters indicate significant differences between sites where the path acted as a firebreak versus those in which it did not as determined by Kruskal–Wallis tests ($p < 0.05$).

vation of the forest–savanna mosaic should be a priority for this reason. Furthermore, Maisels and others (2013) found that forest elephant density was higher near the forest–savanna mosaic of our study site than anywhere else in Africa.

The importance of the forest edge to forest elephants is further illustrated by the exceptionally high elephant density that occurred at the end of the long dry season, when fruit availability at the edge was highest. The end of the long dry season is anecdotally when the elephants are struggling the

most to meet their nutritional needs (White 1994b; White and Abernethy 1997), and as such any fruit available to the elephants during this time is likely to be essential for their wellbeing. The exceptionally high density of elephants during this time also suggests that forest elephants are being attracted to the forest edge from a wide area, indicating the importance of this habitat type for elephant populations across the landscape. This conclusion is supported by previous work by White (1994b), which estimated that forest elephants were attracted to a fruiting event from an area fifteen times the size of the area where the event occurred.

Forest Elephants are Selective Browsers, However This Browsing is Relatively Non-destructive and Not Reflected in Tree Size Demography (Q2)

The forest edge seems to be an attractive resource to forest elephants for browse as well as for fruit, with many species not producing elephant-preferred fruits but still exhibiting signs of elephant browsing damage. Forest elephants did not browse all species of the forest edge equally and showed a strong preference for some species while avoiding others, as is observed for savanna elephants (Owen-Smith and Chafota 2012). At least one of the tree species most favoured by forest elephants, *Millettia versicolor*, seems to be resistant to their browsing, being documented to have a strong coppicing ability (White and Abernethy 1997). A coppicing response to herbivory has been documented in more arid savanna tree species (Staver and others 2012) and, however, has not yet been explored for more mesic forest–savanna transition species. This is thus an interesting topic for future research.

Forest elephants preferred to browse on larger trees (3–4 m tall), similarly to savanna elephants who browse extensively on trees > 2 m tall (Morrison and others 2015). In systems with savanna elephants, this preference leads to large-scale tree mortality (Morrison and others 2015) and a dearth of trees in larger size classes (Asner and others 2009). However, this study showed forest elephants to have no detectable effects on tree size demography, likely because the overall level of damage to trees was relatively low. Damage to larger trees by elephants may be even less severe than estimated here, as we did not take into account that larger trees are usually older and have had more time to accumulate damage, and that some damage may be the result of browsing by other species, specifically large apes.

Previous studies on elephant damage to trees have mostly come from systems dominated by savanna elephants. For example, savanna elephants have been observed to inflict damage on up to 90% of trees taller than 2 m (Morrison and others 2015), while in our study, trees of the same size class were only damaged by forest elephants 28% of the time. The estimates of elephant damage in this study (16% of all stems sampled) are also lower than estimates of forest elephant damage in the forest interior from Terborgh and others (2016) (ca. 50% of all stems 2–6 cm diameter at breast height). However, direct comparison is difficult since Terborgh and others (2016) did not provide the height of the stems sampled and did not sample stems smaller than 2 cm diameter at breast height. Nevertheless, these comparisons highlight the stark differences not only between forest and savanna elephants, but also between forest elephants foraging in different habitats.

Elephants are often assumed to be able to prevent forest encroachment into open-canopied savanna. For example, modelling studies predict how much more open tropical landscapes would be without the widespread loss of mega-fauna (Doughty and others 2015). Specifically, the widespread extirpation of elephants is cited as a key contributor to forest expansion in Africa between the Pleistocene and Holocene (Owen-Smith 1987). The assumption that elephants are important for maintaining and creating open landscapes is based on work done on savanna elephants, who frequently fell canopy-size trees and can be the leading cause of large tree mortality (Asner and Levick 2012; Morrison and others 2015). This study, however, demonstrates that tree-felling may be a uniquely savanna elephant behaviour. That forest and savanna elephants have a non-interchangeable ecological functional lends support to the argument that they should be considered as separate species in conservation policies (for example, by the IUCN (2018)).

Forest Elephant Activity at the Forest Edge is Important for Preventing Savanna Fires from Burning into the Forest (Q3)

The forest interior was protected from burning by the forest elephant path at the forest edge acting as a firebreak in approximately half of the experimental fires. To accomplish this ecosystem function, paths needed to be heavily used, with intermediately used paths being only half as effective. From a conservation perspective, this highlights the importance of maintaining high forest

elephant density within forest–savanna mosaics, as smaller populations are likely incapable of fulfilling this ecosystem function. Although it is possible that other animals contribute to the maintenance of heavily used paths, at our study sites elephants are the mammal species with the greatest cumulative biomass by far (White 1994a) and thus we assume them to play the biggest role in path maintenance. We recommend that the conclusions from this study are tested at forest edges that have no elephant paths or where elephants are locally extinct. Such locations are rare in LNP and were thus not able to be included in this study.

The discontinuity in fuel introduced by a wider, more heavily used, elephant path was a key determinant of whether a path acted as a firebreak or not. In contrast, grass biomass and tree canopy cover were not found to be important factors. However, fuel discontinuity alone is not enough to stop fires, as evidenced by our field observations that heavily used elephant paths in the savanna interior never act as firebreaks (Figure S6). Previous work has shown that fires are significantly less intense nearer the forest edge than in the savanna interior (Cardoso and others 2018). We thus conclude that both reduced intensity and discontinuity in fuel are essential for forming a firebreak at the forest edge.

CONCLUSION

Forest elephants are exceptionally abundant at the forest edge. The forest edge is an important part of the forest elephant's range, providing both fruit and browse resources to them. Forest elephants forage relatively non-destructively compared to their savanna relatives. Thus, the maintenance and creation of open-canopied landscapes in Central Africa remains primarily the role of fire. However, forest elephants' heavily used paths along the edge act as firebreaks and prevent fire from reaching the forest interior. Forest elephants thus provide a valuable ecosystem service by reinforcing the fire-related feedbacks that allow forest and savanna to coexist, and have likely stabilised mosaicked landscapes through time.

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Compliance with Ethical Standards

Conflict of interest The authors declare that they have no conflict of interest.

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