



Transplant Experiments Point to Fire Regime as Limiting Savanna Tree Distribution

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Plant species range shifts are predicted to occur in response to climate change. The predictions are often based on the assumption that climate is the primary factor limiting the distribution of species. However the distribution of grassy biomes in Africa cannot be predicted by climate alone, instead interactions between vegetation, climate and disturbance structure the ecosystems. To test if climatic variables, as predicted by an environmental niche model, determine the distribution limits of two common savanna tree species we established a transplant experiment at a range of latitudes and altitudes much broader than the distribution limits of our study species. We planted seedlings of two common savanna trees, Senegalia nigrescens and Colophospermum mopane, at eight paired high and low elevation sites across an 850 km latitudinal gradient in South African savannas. At each site seedlings were planted in both grassy and cleared plots. After 2 years of growth, rainfall, temperature and location inside or outside their distribution range did not explain species success. Grass competition was the only variable that significantly affected plant growth rates across all sites, but grass competition alone could not explain the distribution limit. Species distributions were best predicted when maximum tree growth rates were considered in relation to local fire return intervals. The probability of sapling escape from the fire trap was the most likely determinant of distribution limits of these two species. As trees grew and survived 100 s of kilometers south of their current range limits we conclude that climate alone does not explain the current distribution of these trees, and that climate change adaptation strategies for savanna environments based only on climatic envelope modeling will be inappropriate.

Keywords: species distribution, range limits, savanna, range shifts, transplant, grass competition, Colophospermum mopane, Senegalia nigrescens

INTRODUCTION

Global change is transforming the planet (Ellis, 2011; Scheffers et al., 2016). One of the most pervasive impacts, which has severe economic and ecological impacts, is that of shifting plant and animal ranges (Parmesan and Yohe, 2003; Thomas et al., 2004; Chen et al., 2011). The shifts are most frequently attributed to the warming of the earth, with the expectation that plant and animal distributions will shift toward the cooler poles or higher elevations (Lenoir et al., 2008; Chen et al., 2011). These expectations (Hutchinson, 1957; Chase and Leibold, 2003) are often derived from the

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Hutchinson concept of the fundamental and realized niche. The fundamental niche describes the set of environmental factors that determine the space where a species could occur in the absence of biotic interactions (Hutchinson, 1978; Chase and Leibold, 2003). The realized niche describes where a species actually occurs after accounting for the effects of inter-specific competition. This concept has been applied almost globally as a framework to describe the geographic distribution of species (Gaston, 2003; Thomas, 2010). However this framework is generally simplified to assume that climate shapes a plant's distribution range, and is the basis of attempts to predict the future ranges of plants using the correlative climate envelope approach (Thomas et al., 2004; Hijmans et al., 2005) where the current distribution of a species is mapped in climate space and any future shifts in climate space are expected to produce an accompanying shift in the future species distribution range (Elith and Leathwick, 2009). Although there is debate as to the broad applicability of climate models, (Davis et al., 1998; Araújo and Luoto, 2007; Svenning and Skov, 2007; Araújo and Peterson, 2012), the underlying general assumption that climate alone is the primary determinant of species distribution limits (Holdridge, 1967; Gaston, 2003) is seldom questioned.

Much of the empirical support for these climate based assumptions comes from studies in temperate and boreal northern hemisphere systems (Parmesan and Yohe, 2003; Thomas, 2010) where temperature is often the primary factor that limits plant production (Jolly et al., 2005; Körner and Basler, 2010). There remains an acute absence of detailed studies on distribution limits of plants in the tropics (Blach-Overgaard et al., 2010; Thomas, 2010). Here, authors acknowledge that the determinants of ranges in tropical regions are more complex (Root et al., 2003; Blach-Overgaard et al., 2010; Thomas, 2010). Water availability and biotic interactions are considered increasingly important in these systems (Woodward, 1987; Thomas et al., 2004; Jolly et al., 2005) but the underlying assumption that the distribution ranges can be predicted by climate alone remains common. However, we contend that this underlying principle and approach is not sufficient to predict the current, or future distribution of plants occurring in Africa's grassy biomes.

In tropical savanna and grassland biomes, interactions between fire, mammal herbivory and climate structure savannas (Bond et al., 2005; Staver et al., 2011; Lehmann et al., 2014). Tropical grassy biomes (TGBs) are therefore considered the most extensive vegetation area not at equilibrium with climate (Bond et al., 2005; Pausas and Ribeiro, 2017). Fire and mammal herbivory are dominant environmental filters that define community assemblages (Staver et al., 2012; Charles-Dominique et al., 2016; Pausas and Ribeiro, 2017), with the relative importance of these processes changing across the rainfall gradient (Archibald and Hempson, 2016). Herbivory can cause local species extinctions, alter competitive interactions, and reduce or enable tree recruitment (Prins and van der Jeugd, 1993; Bond and Loffell, 2001; Augustine and Mcnaughton, 2004; Asner et al., 2009), and fire regimes can alter the community structure and composition (Trapnell, 1959; Higgins et al., 2007; Pausas and Ribeiro, 2017). Whilst these processes can act on woody plants at all demographic stages (Midgley and Bond, 2001), the demographic stages which are most impacted are the seedling establishment and sapling stage where a critical population bottleneck occurs (Higgins et al., 2000). Here, regular disturbance traps saplings in the grass layer, where they are vulnerable to further topkill by fire and herbivory (Higgins et al., 2000; Midgley et al., 2010; Werner and Prior, 2013). Importantly these suppressed saplings are not reproductive and repeated topkill can prevent the recruitment of reproductive individuals, which would eventually lead to local extinction (Higgins et al., 2000). When trees exceed \sim 3 m in height, they will often be sufficiently fire proof to be released from the fire trap (Bond, 2008; Wakeling et al., 2012). Therefore the ability of a plant to escape this trap to reach reproductive maturity within the prevailing climatic and disturbance context is the critical component that will determine the presence of a plant in a region.

To explore potential climatic constraints within a disturbance limited system we tested how well the traditional climate based framework of plant species distribution can explain patterns of species distribution ranges in consumer controlled systems. We established a transplant experiment across a latitudinal gradient across South African savannas inside and outside the distribution ranges of two common savanna tree species; Colophospermum mopane (Kirk ex J.Léonard) and Senegalia nigrescens [(Oliv.) P. J. H. Hurter]. The transplant experiment design was guided by the outputs of species distribution models and literature which describe the climatic determinants for the study species; Colophospermum mopane. Low winter temperatures (below a minimum July average of 8°C), high rainfall (greater than 800 mm), and the presence of frost and increases in rainfall are commonly reported climatic limits to the occurrence of this species (Henning and White, 1974; Burke, 2006; Makhado et al., 2014; Scholtz et al., 2014; Stevens et al., 2014b).

To explore the role of these proposed limits the experiment was established across a climate gradient across a wide range of summer rainfall and temperature conditions, whilst remaining within the limits of the savanna biome. The latitudinal gradient follows a rainfall gradient with the driest savanna areas occurring in the lower latitudes with the northernmost site receiving \sim 350 mm rainfall and the southernmost sites receiving \sim 1,140 mm (**Table 1**). To account for the role of temperature and to extend the range of temperatures that the plants were exposed to, we paired each low altitude transplant site across the 850 km latitudinal gradient with a cooler higher elevation savanna site (above 500 m.a.s.l). Within each site, trees were planted with and without grass competition is grass competition is the strongest form of biotic competition in savannas (Riginos, 2009; February et al., 2013).

If climate is the primary limitation on the distribution of our tree species across a latitudinal gradient, then their distribution limits should coincide with climatic conditions where plant performance declines. In line with the literature, we predict that in the absence of grass competition, plant performance will decline with increasing distance from the range edge (Sexton et al., 2009; Hargreaves et al., 2013; Lee-Yaw et al., 2016). In the presence of grass competition, performance declines should match the current distribution limits and plants will not establish

-	-at/Long	Elev. (m)	MAT (C°)	Mean temp warm. quart. (C°)	Mean temp warm. month (C°)	Mean of cold. month (C°)	Mean min of cold. month (C°)	Mean max warm.month (C°)	MAP (mm)	Grass biomass (kg/ha)	Fire return interval (CI)	S. <i>nig.</i> range	C. <i>mop</i> . range
0.0	95 E, 22.85 S	427	23.2	26.4	27.4	16.3	8.2	38.1	522	96	5-6	Ē	<u> </u>
6	25 E, 22.43 S	631	24.6	28.5	28.0	17.0	8.7	36.3	344	120	40	드	Ч
Ξ.	94 E, 25.10 S	197	23.0	26.6	27.0	16.7	7.4	36.7	625	4056	2.5-3.5	드	Out
ø	.34 E, 25.39 S	1093	18.9	23.7	23.7	11.3	0.68	31.7	605	2026	2.5-3.5	Out	Out
Ξ.	83 E, 27.37 S	221	22.2	25.5	25.9	16.9	9.1	33.8	678	5437	3-4	드	Out
5	.57 E, 27.54 S	551	20.9	24.1	24.5	16.4	10.3	32.4	707	6021	3-4	드	Out
ĕ).68 E, 30.3 S	137	20.5	24.0	24.9	15.9	10.7	29.5	1133	9564	3.5-4.5	Out	Out
2	.37 E, 29.63 S	850	21.9	19.0	22.3	12.31	7.1	30.2	897	10728	3.5-4.5	Out	Out

at sites outside their range, a result that is frequently recorded in similar experiments in temperate systems (Hargreaves et al., 2013; Lee-Yaw et al., 2016). However if climate (either rainfall or temperature) is not the primary process limiting the current range we predict that the plants will grow and survive both inside and outside their distribution ranges in the absence of fire and herbivory. We excluded fire and herbivory at our experimental sites so as to test for climatic limits alone. However as the degree of fire and herbivory also change across rainfall and temperature gradients we used growth data from the experiment to develop models that link growth rates and observed fire return periods to explore how fire regime might interact with growth constraints to limit species range.

Range Limits in Savanna Trees

MATERIALS AND METHODS

Study Species

We transplanted two common savanna trees, Colophospermum mopane (mopane) (Kirk ex Benth.) Kirk ex J.Léon and Senegalia nigrescens (Oliv.). C. mopane is a highly dominant leguminous tree or multi-stemmed shrub, belonging to the detarioideae subfamily of the Fabaceae. Of the 1.5 million km² of the savannas in southern Africa, C. mopane is estimated to almost singly dominate 550,000 km² (25-35%) of savannas (Mapaure, 1994; Timberlake, 1995) where it often forms monospecific stands (Timberlake, 1995) on both nutrient rich and nutrient poor substrates. While dominating many thousands of hectares across southern Africa, its southern-most distribution ends abruptly in South Africa at \sim -22.5S in the west and at \sim 24.0S in the east (Mapaure, 1994) (Figure 1). S. nigrescens (Fabacae) or the Knob thorn is a woody tree that can grow up to 20 m in height, although it usually remains between 8 and 10 m (Figure 6). It is most common and widespread in low-altitude savannas occurring from Tanzania to South Africa. The southern distribution limit for this species is at \sim 28.5 degrees latitude and it is considered frost sensitive, preferring low-lying areas (Smith et al., 2012).

Transplant Experiment

Eight transplant sites were established within the South African savanna biome across an 850 km latitudinal gradient ($\sim 8^{\circ}$ lat.), covering a rainfall gradient from 340 to 1,130 mm (Table 1). Four low elevation sites (<500 m.a.s.l) were paired with four high elevation sites (>500 m.a.s.l) at the same latitude. The paired cooler higher elevation sites were chosen to ensure adequate representation of cold and frost prone sites (Schulze, 2007). The latitudinal and altitudinal gradients included sites inside and outside the distribution ranges of both C. mopane and S. nigrescens (Figure 1). Sites across these gradients were selected if they occurred within a savanna, had a perennial, undisturbed grass layer and a clay content of 20-30% (both on the surface and at 20 cm). At each site, a 30×30 m plot was established and fenced to prevent mammal herbivory. Each plot was divided into 6 sub plots (Figure 2). Three replicated sub-plots were randomly cleared of all grass and forb biomass by the roots using a hoe whilst grass cover in the remaining three was preserved.

S. nigrescens and C. mopane seeds were collected from within their respective distribution ranges. S. nigrescens seeds were

TABLE 1 | Location, climatic and biophysical characteristics of transplant sites



squares & S. nigrescens = white squares). All sites were placed within the savanna biome. The northern most sites ($\sim 22^{\circ}30'S$) are referred to as latitude 1 sites and the southernmost sites ($\sim 30^{\circ}30'S$) as latitude 4.

collected within a \sim 30 km radius of the South African Wildlife College (24.453 S, 31.405 E) and C. mopane seeds were collected within a 30 km radius of All days town (22.67 S, 29.10 E). The collected seeds were mixed and germinated at a nursery in the lowveld savanna at the South African Wildlife College. Seeds were planted in 1 l potting bags in a sand: clay mixture which was inoculated with soil taken from the roots of each species growing in their natural habitat. Following successful germination, the 480 one month old seedlings were transported to the sites and were planted in the early growing season (3 November-21 November 2010). Within each replicate five C. mopane and five S. nigrescens one month old seedlings were planted in random order one meter apart (Figure 3). To reduce losses to transplant stress and to allow the seedlings to establish, an equivalent of a 10 mm rainfall event was supplied every week for 1 month. Recruitment in these species is episodic (Botha, 2006; Stevens et al., 2014a) and by watering at the seedling establishment period



FIGURE 2 | Experimental layout. Each 30×30 m plot consisted of 6 blocks with three replicated sub-plots of grassy treatments and three replicates subplots of a cleared treatment, where all grass was removed. Five plants of each species were planted in each replicate, at one meter intervals.

we simulated an early growing season with good rainfall that was likely to result in successful establishment. As the requirements for seedling establishment have been documented we replicated these conditions so we could examine the determinants of plant success and growth following early establishment. For successful early seedling establishment the plants require warmth and moisture. These environmental conditions were met at all the savanna sites both inside and outside the distribution range of both species and were therefore not the limiting factor to establishment (Botha, 2006; Stevens et al., 2014a).

Seedling Survival

We monitored plant growth for 2 years. Sites were revisited three times annually at the start, middle and end of the growing season (early November, January, April). A round trip between sites was a distance of 5,000 km; therefore more frequent measurements were not taken. During each measuring period seedling survival was assessed and the standing height and stem diameter of each seedling was measured. At the end of the growing season the grass biomass in the grass treatments was measured using a disc pasture meter (DPM). DPM readings were converted to biomass using the conversion developed by Trollope and Potgieter (1986). i-Buttons were placed at each site at 1.2 m above the ground and at 0.15 m above the ground to assess the thermal environment experienced by the seedlings. I-Button data was used to establish the site specific temperatures reported in Table 1. I-Buttons were suspended inside 4 mm thick PVC tubes (20 \times 40 cm), and remained in the field for the duration of the experiment.

Data Analysis

The data were analyzed in R. 3.5 (R Core Team, 2018). To examine the survival of seedlings during the course of the experiment we used the survfit function to create Kaplan-Meier survival curves for the full set of seedlings distinguishing between treatment (grass: no grass), distribution (inside: outside the species range) and species. Log-rank tests were used to compare



the groups. To examine if there was an effect of transplant

the groups. To examine if there was an effect of transplant shock on seedling survival we ran all the Kaplan-Meier survival curves both with the default alpha, and with an alpha of-10 which reduces the weighting of the first part of the dataset.

We used mixed-effects Cox proportional hazards models using the coxme (version 2.2.10) package (Therneau, 2018) in R to quantify the effects of treatment, distribution, mean annual temperature and mean annual precipitation and species as predictors of seedling survival, as well as the interaction between treatment and species. To account for the ties in the data (due to the limited number of sample dates) we used the "Efron" method for calculating ties. We ran the model for the full dataset, and for a dataset that excludes the transplant period (i.e., only data after day 130 were used in the model). We included both plot and site as random effects (plot nested within site). Candidate models were compared and ranked using Bayesian information criterion (BIC). If the best candidate models differed with a Δ BIC <2 we used model averaging to determine the best model parameters.

In order to determine the relationship between final plant height and climate parameters, mean annual temperature (MAT) and precipitation (MAP) were calculated for each site using temperature data gathered from i-Buttons on site and rainfall records from either long term farm records or the closest SA weather service station. The final mean height differences between sites were analyzed using linear-mixed effects models in the nlme package (Pinheiro et al., 2009). Linear mixed effects modeling fit by maximum likelihood was used to assess the significance of mean annual temperature, mean annual rainfall, grass treatment (grass or no grass) and range (inside or outside of distribution range) on the height of saplings after 2 years of growth. We included mean annual temperature and mean annual precipitation as fixed covariates in the analysis and distribution (inside or outside range) and treatment (grass or no grass) as fixed categorical factors. Site was included as a random factor and plot was nested within site. All possible interactions were considered. The candidate models were compared and ranked using Bayesian information criterion (BIC). The second best candidate model for both species differed with a Δ BIC >2 so model averaging was not used and the first best fit model with the lowest BIC was selected. We present the final parameter estimates, standard errors, and confidence intervals to demonstrate the effect size of the different parameters of the top two models.

Modeling Escape From the Fire Trap

To model the probability of sapling escape from the fire trap we used mean maximum growth rates for the two fastest growing plants of each species from the grass treatment, for each site (Wakeling et al., 2011; Bond et al., 2012). Growth rates were calculated from the beginning of the second growing season to the beginning of the third. We did not include the first season in the growth rate calculations as growth in the first season is characteristic of seedlings which have different allocation patterns and hence different above ground growth rates than that of established saplings (Higgins et al., 2000). We used the growth model from Higgins et al. (2000) to model the height gain of savanna trees, and hence time to reach a fire proof height (\sim 3 m):

$$h = h_{y-1} + \left(1 - \frac{h_{y-1}}{h_{\max}}\right)g_s$$

where g_s is the growth rate of stems (cm year⁻¹) (year 2), h_{max} is the maximum stem height (10 m), and $h_{\nu-1}$ is the stem height

in the previous year. Starting stem heights were set at a starting height of 20 cm, the mean stem height after year one across all the transplant sites. We kept the starting height the same across sites so that responses could be restricted to the sapling stage, not the seedling establishment stage. The model was run for 50 years.

As the model produced an output of tree height per year, we could determine the time it took a tree at each site to reach the "escape height" of 3 m, given the maximum growth rate at a site (Wakeling et al., 2012). We used maximum growth rates as they are more likely to be ecologically relevant than mean growth rates as predictors of sapling release and successful recruitment into the adult stage (Wakeling et al., 2011; Bond et al., 2012). We plotted this value against the median fire return interval (FRI) for each experimental site. FRI was calculated based on fire return intervals for low-human density African savanna regions (Archibald, 2016). This was estimated by fitting a Weibull distribution (Johnson and Gutsell, 1994) to fire interval data derived from the MODIS burned area product (see Archibald et al., 2009; Archibald and Hempson, 2016 for methods) for \sim 1,000 points from undisturbed areas.

RESULTS

Site Characteristics

Sites spanned across a latitudinal gradient (Table 1). With increasing latitude mean annual precipitation (MAP) increased from \sim 400 to 1100 mm and mean annual temperature (MAT) decreased by \sim 3°C (**Table 1**).The differences in rainfall between high and low elevation sites were negligible except for the lower latitude northern sites (VN & PM) where the low elevation site received 120 mm more per year. High and low elevation sites differed in their mean annual temperatures with the low elevation sites having higher mean annual temperatures, warmer winters and mild winter minimums. VN, the low latitude higher elevation site was the exception, as it is had the highest MAT, however the mean monthly maximums were higher at the low elevation site. This site is also the driest site receiving a MAP of 344 mm. The wettest site in the low elevation, higher latitude (UK) received a MAP of 1,133 mm. The coldest site was the higher elevation KW site with a MAP of 18.9 and a July mean minimum of 0.68°C (the coldest month). KW was also the only site that received regular frost events during winter.

Seedling Survival

We used Kaplan-Meier survival curves to show how distribution, treatment and species impacted seedling survival during the experiment (**Figure 3**). The presence or absence of grass had the strongest impact on seedling survival with grasses reducing survival probability by ~50% [$X_{(1, 457)=}^2$ 31.0, p < 0.0001]. This effect still remained significant after accounting for transplant induced mortality at the beginning of the experiment [$X_{(1, 457)=}^2$ 28.0, p < 0.0001]. Interestingly there was significantly lower survival inside the distribution range [$X_{(1, 457)=}^2$ 34.0, p < 0.0001], but the bulk of this effect occurred during the initial transplant period (first 130 days). However distribution range was not significant when the early stages of the experiment

were given less weight (alpha-10) $[X_{(1, 457)=}^2 1.3, p = 0.253]$. There were no significant differences in the survival rates between species with both species showing similar responses to distribution range and grass treatment.

For the first dataset (all time steps included) mixed-effects Cox proportional hazard models with the most support (based on BIC and model inference) included grass treatment and distribution range. Although this model had a weight of 0.51 there were two other important models-one that included MAP, and one that included MAT (Table 2A). As there were three models where Δ BIC <2, we averaged these three models. The model averaged co-efficients indicate that treatment, distribution range, MAT and MAP all have explanatory power. The terms with the highest weighting were treatment and distribution with a weighting of 1, whilst MAP and MAT each had a weighting of 0.23. The final averaged model indicates that distribution range (p < 0.01) and treatment (p < 0.001) are the only significant explanatory variables in the model. The log of the co-efficient values show that plots without grass had \sim 2 times the survival rate as plots with grass, and plots inside the distribution range had about half the survival rates as plants outside the range (Table 3A).

To see whether the effect of distribution was limited to the early transplant period, we ran the same model but excluded all data before day 130. In this case the top 4 models all had very similar weights (0.10–0.11) with distribution range, MAP, MAT and treatment appearing as important terms (**Table 2B**). We therefore averaged the model results from models where Δ BIC <2 (**Table 3B**). Treatment was the most important variable and received a weight of 1, followed by MAT (0.73), MAP (0.59), distribution range (0.42), and species (0.5). In the final model the grass treatment was the only significant variable (*p* < 0.0001) and individuals planted with grass were ~three times as likely to die (**Table 3B**).

Final Sapling Height

We examined how the final height was explained by distribution range, MAT, MAP, grass treatment and all possible interaction terms. We used BIC values and model inference to select the best fit model. The best fit final mixed linear model for both *C. mopane* and *S. nigrescens* included only the treatment variable. We did not use model averaging for the results as the next best fit model for both species had a BIC>2. Plants were significantly shorter at the end of the experiment if they were grown in the presence of grass (**Table 4, Figures 4, 5**).

Contrary to predictions, frost did not cause sapling mortality. KW was the only site which experienced frost. Several frosts during the dry season in year 1 caused a decline in height for both species, (**Figures 4**, **5**). Frosted trees of both species were topkilled, but both species resprouted from the base in the following growing season. Resprouts were always multistemmed.

As grass is an important factor in sapling growth we investigated how it impacts sapling growth across the transplant experiment. Grass biomass had a significant effect on tree sapling growth within the sites and across the latitudinal gradient for both *C. mopane* (p < 0.01) and *S. nigrescens* (p < 0.001) (**Figure 6**). Sapling growth in the presence of grass for both species was highest where the grass biomass was lowest, and

		Co-efficient			df	BIC	Δ BIC	BIC Weight
Range	MAP	MAT	Species	Treat				
(A) MODEL	S BASED ON ENT		AL PERIOD					
+				+	18	2346.5	0	0.5410
+	0.000			+	18	2348.2	1.71	0.2300
+		0.01924		+	18	2348.2	1.72	0.2290
(B) MODEL	S EXCLUDING TH	IE FIRST 130 DAYS						
+	0.001	-0.221		+	18	1203.700	0.000	0.130
	0.001	-0.077		+	18	1203.900	0.150	0.12
		-0.142		+	18	1203.900	0.160	0.12
	0.001			+	18	1204.000	0.240	0.115
+		-0.285		+	18	1204.100	0.340	0.109
+	0.001	-0.231	+	+	19	1205.100	1.340	0.066
+		-0.288	+	+	19	1205.100	1.380	0.065
		-0.154	+	+	19	1205.200	1.500	0.061
	0.001	-0.097	+	+	19	1205.300	1.580	0.059
	0.001		+	+	19	1205.500	1.790	0.053
				+	17	1205.600	1.860	0.051
+	0.002			+	17	1205.600	1.880	0.051

TABLE 2 Best models (where BIC <2) assessing the relationship between variables and seedling survival for (A) the entire experimental period and (B) the entire experimental period but with the first 130 days receiving a lower weighting (Alpha = -10).

Models are ranked based on differences in the Bayesian Information Criteria (BIC). BIC weight is the weight of each model. Range, inside or outside distribution range; MAP, mean annual precipitation; MAT, mean annual temperature; Species, species of seedling (S. nigrescens or C. mopane); treat, presence or absence of grass.

TABLE 3 | Final parameter estimates (β), standard errors (SE) and confidence intervals of model averaging based on top models for (A) analysis of entire experimental period and (B) analysis excluding first 130 days to account for effects of tranplant shock.

(A) MODELS BASED ON ENTIRE EX Range -0.913 Treatment 0.840	PERIMENTAL PERIOD	0.005	
Range-0.91Treatment0.840MAD0.000	–1.505	0.005	
Treatment 0.840		-0.325	0.301
NAD 0.000	0.486	1.194	0.181
MAP 0.000	-0.001	0.001	0.000
MAT -0.004	-0.223	0.185	0.050
(B) FIRST 130 DAYS EXCLUDED FRO	M ANALYSIS TO ACCOUNT FOR	TRANSPLANT SHOCK	
Range -0.32	1.83902	0.314	0.518
MAP 0.00 ⁻	-0.001	0.003	0.001
MAT -0.135	-0.527	0.157	0.170
Treatment 1.105	0.596	1.613	0.259
Species (CM) 0.194		4 055	

Significant parameters are shown in bold. Range, inside or outside distribution range; MAP, mean annual precipitation; MAT, mean annual temperature; Species, species of seedling (S. nigrescens or C. mopane); treat, presence or absence of grass.

sapling height progressively declined as grass biomass increased across the sites for both species (**Figure 6**).

Modeled Tree Growth and Escape From the Fire Trap

Considering that seedling survival and performance were not well explained by distribution range we then looked at processes preventing sapling recruitment into adult size classes. Grass biomass not only reduces sapling performance (**Figure 6**), but also provides the fuel that determines fire return intervals (Figure 7). High grass productivity, linked to higher rainfall, permits frequent fires. We therefore predicted that a combination of slower growth rates and more frequent fire return intervals might be a significant factor preventing sapling recruitment. Species growing inside their distribution ranges should be more likely to reach fire escape height within the fire free interval common to their area. Our model indicates that *C. mopane* fails to escape from the fire trap when maximum growth rates are low (caused by high grass biomass) (Figures 7, 8), or when fire return interval is short (Table 1, Figure 7). Model results (Figure 8) indicate that within *C. mopane's* distribution range

TABLE 4a | Mixed linear model outputs showing the top two models selected using BIC values for C. mopane.

	Co-effic	sient	BIC	ΔBIC	Weight
	Treatment	MAT			
Model11	-284.72 [-390.21: -179.24]		1788	0	0.74
Model10	-285.18 [-390.46: -179.90]	77.78 [-32.61: 188.18]	1791	2.36	0.23

Bold terms indicate significance. Treatment, the presence or absence of grass, was the only factor that significantly affected tree growth. Confidence intervals are presented in square brackets.

TABLE 4b | Mixed linear model outputs showing the top two models, selected using BIC values for S. nigrescens.

	Co-effic	ient	BIC	∆BIC	Weight
	Treatment	MAT			
Model11	-281.30 [-423.4: -139.56]		1545.81	0	0.85
Model10	-281.26 [-423.2: -139.36]	42.72 [-74.60: 160.03]	1549.75	3.94	0.12

Bold terms indicate significance. Treatment, the presence or absence of grass, was the only factor that significantly affected tree growth. Confidence intervals are presented in square brackets.



sapling growth rates with grass are fast enough to result in trees escaping the fire trap. Outside the distribution range, although actual growth rates can be higher, fires are more frequent, FRI intervals decrease, and *C. mopane* is unlikely to reach escape height before the next fire. Likewise for *S. nigrescens*, the probability of successful escape from the fire trap only occurred within the existing distribution range of this species. At these sites

maximum tree growth rate in the presence of grass can out-pace the fire return interval (**Figure 8**).

DISCUSSION

The performance of two common savanna trees did not decline with increasing distance from the range edge. Instead plants still





survived and grew well hundreds of kilometers outside their distribution ranges. This is contrary to the generally observed pattern that species distribution limits are often niche limits (Lee-Yaw et al., 2016). In our study species, and perhaps more generally in common savanna plants, climate axes of the niche were poor predictors of species performance in and outside range

limits. Plant performance instead was most strongly determined by grass biomass. Whilst saplings still survived and grew within the presence of grass, the poorer performance with high grass productivity reduced the probability of saplings escaping the fire trap between successive fires. Our results and modeling suggest an alternative framework for the determinants of species ranges.



Prediction of ranges for these and potentially other savanna species requires an understanding of the interactions between tree and grass productivity, climate and disturbance (top-kill) which determine the ability of saplings to escape the fire trap at any given site.

As the plants survived and grew well past their southern limits, a classic interpretation would be that both these species were dispersal-limited since suitable, but unoccupied, areas existed beyond their range boundaries (Hargreaves et al., 2013; Lee-Yaw et al., 2016). While dispersal limitation does occur, it is not common (Sexton et al., 2009; Hargreaves et al., 2013), and it is highly unlikely that these two widely distributed, dominant savanna species are dispersal limited. The fruits of C. mopane are well dispersed by wind and water (Styles and Skinner, 1997; Mlambo and Nyathi, 2004), and allozyme frequencies are similar across Southern African populations, indicating that an effective mechanism of gene flow occurs (Villoen et al., 2003). S. nigrescens retains closed pods on the tree until ripe, when gusting winds, associated with the onset of rains, rip the pods off the tree where they open, scattering the seeds widely (Miller, 1994; Grant and Thomas, 2001).

It is often documented that herbaceous competition strongly suppresses sapling and tree growth (Riginos, 2009; February et al., 2013) and likewise the presence of grasses in our study has a strong suppressive effect on seedling survival and sapling growth. It is through this interaction where climate probably has the strongest indirect influence on tree establishment as grass biomass shows a strong positive increase with increasing rainfall (O'connor et al., 2001). In the juvenile stages, tree



FIGURE 8 The modeled time to reach escape height in the presence of grass plotted against the approximate fire return period for the site for (A) *S. nigrescens* and (B) *C. mopane.* Closed circles show a species was planted within its distribution range, open circles show the plant was grown outside its distribution range. The dashed diagonal line indicates the minimum fire return interval required for each time-to-escape-height value. If the points fall above the 1:1 line, into the shaded area, the plants have a high probability of escaping from the fire trap. Both species, if planted within their distribution range, have a reasonable probability of escape the prevailing fire regimes outside their distribution range.

competition with grasses is likely to be highest as tree and grass roots occupy the same soil layers (Cramer et al., 2007). We therefore hypothesize that the distribution range of both of these tree species is a product of the interaction between sapling growth rates, grass competition, and the fire return interval which depends on grass biomass for fuel. Here, climate plays an indirect role through its interactions with fire return interval and grass biomass (O'connor et al., 2001; Archibald et al., 2009). Grass biomass can set the maximum annual tree growth rate with growth rates decreasing with increases in grass biomass. The probability that any sapling will successfully escape the fire trap is dependent on its maximum growth rates (Wakeling et al., 2011; Bond et al., 2012). The sapling escape time (the time it takes for savanna tree to reach 3 m) needs to occur within fire free intervals. If a fire burns a sapling it will be top-killed and will have to start growth from a resprout. Our model therefore matches the time taken to reach escape height against the prevailing fire frequency of an area. We suggest that the probability of successful establishment is very low when the time to reach escape height is greater than the prevailing fire return interval. The chances of establishment increase as the times become more closely matched. The highest probability of establishment therefore occurs when escape height can be easily reached within the window of the prevailing fire free interval (Higgins et al., 2000).We therefore suggest that disturbance (e.g., fire) is the critical missing component in determining whether a plant can become a successfully recruiting adult, and hence should be explicitly considered when predicting the future ranges of species. Fire regimes can, and are, being modified by land users (Archibald et al., 2013). Whether climate change, or fire (and herbivore) management has greatest effects on changes in species distribution needs careful consideration in savannas.

Whilst we demonstrate our idea using fire return interval this framework applies to any process that causes top-kill in savannas including mammal herbivory (Staver et al., 2012) and frost (Holdo, 2007; Whitecross et al., 2012). In our experiment, frost caused top-kill in the regularly frosted site (KW). It is a climatic factor considered important in limiting the distribution of C. mopane (Henning and White, 1974; O'Connor and Bredenkamp, 1997; Burke, 2006). However we found that frost did not directly kill these seedlings, as often predicted, instead it acted as a powerful top-kill agent with a similar impact to that of fire. Both species experienced severe frost (Figure 6) and experienced complete topkill. In the subsequent growing season multiple shoots resprouted from the base and the plants continued to survive and grow but lost the previous season's height gains. In sites with regular frost, the equivalent metric would be the frost return interval, and saplings would need to reach a height threshold where frost damage is negligible (Whitecross et al., 2012) or rely on frost free intervals to ensure successful establishment (Holdo, 2007; Wakeling et al., 2012; Whitecross et al., 2012), or experience eventual population exclusion within that area (Wakeling et al., 2012). Likewise mammalian grazers can interact with the distribution of species through the removal of grass and the reduction of fire frequencies which in turn will lessen the fire imposed bottleneck (Hempson et al., 2017). Alternatively a high prevalence of browsers in the landscape has the potential to increase sapling top-kill further reducing the likelihood of escape from the fire trap (Staver et al., 2009).

Similar insights have been made at biome levels across continental scales where fire, vegetation and climate are recognized to be fundamental in predicting the distribution of the savanna biomes (Whittaker, 1970; Bond et al., 2005; Lehmann et al., 2014). As these processes interact to shape the distribution of a biome it should not be surprising that the dominant species within these biomes are shaped by similar forces. Several unrelated studies have demonstrated that fire is an important

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environmental filter and determinant of species composition (Staver et al., 2012; Pausas and Ribeiro, 2017). For example, fire exclusion can drive a switch in species composition or a shift in the dominant functional types e.g., evergreen species (Trapnell, 1959; Plas et al., 2013; Charles-Dominique et al., 2017). However, we do not know of other studies showing how fire regime can restrict the regional distribution of savanna tree species. Our results indicate that top-down processes are important in explaining the distribution of these two savanna tree species and processes that cause top-kill like fire, frost and herbivory can act as strong environmental filters and thus be as important in shaping plant distributions as temperature in temperate systems. Climate-based species distribution models are widely used as a tool in predicting the future of African plant species and as a tool to shape climate change policy across the developing economies of Africa (Heubes et al., 2011; Bocksberger et al., 2016). We suggest that their assumptions need much wider testing given the growing evidence for consumer-control of these ecosystems.

AUTHOR CONTRIBUTIONS

NS and WB conceptualized the idea. NS and SA designed the experiment and NS performed field work. SA commented on drafts, provided funding, fire data and advised on statistics, and performed some statistical analysis. WB provided funding, and commented on drafts.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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