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Savanna woody encroachment is widespread across three continents

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1	Savanna woody encroachment is widespread across three continents
2	Running head: Global savanna encroachment
3	
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19 Abstract

20 Tropical savannas are a globally extensive biome prone to rapid vegetation change in response 21 to changing environmental conditions. Via a meta-analysis, we quantified savanna woody vegetation 22 change spanning the last century. We found a global trend of woody encroachment that was 23 established prior the 1980s. However, there is critical regional variation in the magnitude of 24 encroachment. Woody cover is increasing most rapidly in the remaining uncleared savannas of South 25 America, most likely due to fire suppression and land fragmentation. In contrast, Australia has 26 experienced low rates of encroachment. When accounting for land-use, African savannas have a mean 27 rate annual woody cover increase two and a half times that of Australian savannas. In Africa, 28 encroachment occurs across multiple land uses and is accelerating over time. In Africa and Australia, 29 rising atmospheric CO_2 , changing land management and rainfall are likely causes. We argue that the 30 functional traits of each woody flora, specifically the N-fixing ability and architecture of woody plants, 31 are critical to predicting encroachment over the next century and that African savannas are at high 32 risk of widespread vegetation change.

34 Introduction

The savanna biome covers around 20% of the Earth's surface, contributing 30% of terrestrial net primary production (NPP) (Grace *et al.*, 2006), and is increasingly considered critical to the regulation of the terrestrial carbon cycle (Poulter *et al.*, 2014; Liu *et al.*, 2015). However, savannas are also home to 20% of the Earth's population and the majority of rangelands and livestock (Scholes & Archer, 1997). Understanding current trends in vegetation change across the biome is critical to predicting the future of this ecologically, economically and culturally important biome.

41

Global change is increasingly understood to be affecting tropical vegetation at large spatial 42 43 scales. A poorly resolved global driver of vegetation change is the exponential increase of 44 atmospheric CO_2 concentration, henceforth denoted as $[CO_2]$. Rising $[CO_2]$ affects plant growth and vegetation both directly, and indirectly via [CO₂]-driven climate change (Leakey et al., 2009). 45 46 Concurrently, humans have caused extensive change in the spatial processes of fire and mega-47 herbivory that maintain savanna ecosystem function, via land clearing for agriculture, building of 48 roads and fences, hunting, and deliberate lighting or suppression of fire (Lamprey & Reid, 2004; Archibald et al., 2013). Combined, the degree of human-induced land use and land cover change, 49 50 increases in [CO₂] and future climate change mean change in savanna vegetation structure and 51 function must be anticipated. However, estimates of future savanna vegetation change vary 52 considerably; some modelling studies predict that elevated [CO₂] will drive large increases in 53 savanna woody cover (Higgins & Scheiter, 2012; Donohue et al., 2013), while other models that only 54 consider climate changes produce variable predictions ranging from savannas remaining stable 55 (Bergengren et al., 2011) to savannas invading forests (Anadon et al., 2014) and forests invading savannas (Zelazowski et al., 2011). 56

57 Woody encroachment, the increase in woody biomass, stem densities or woody cover, is a 58 symptom of an alteration in ecosystem processes; it has been documented across continents and

biomes (Myers-Smith *et al.*, 2011; Murphy *et al.*, 2014; O'Connor *et al.*, 2014). However, as an opencanopied tropical biome, savannas appear particularly vulnerable to woody encroachment (Parr *et al.*, 2014), where encroachment can have negative consequences for soil carbon storage (Berthrong *et al.*, 2012), ground water recharge, tourism (Gray & Bond, 2013), grazing potential (Angassa &
Baars, 2000) and biodiversity (Ratajczak *et al.*, 2012).

64 Predicting woody encroachment across savannas is hampered by their ecological complexity 65 as a wide range of drivers can act to release constraints on woody plant growth and recruitment -66 including browsing, grazing, fire, climate, and [CO₂]. Combinations of local (e.g. land use), regional 67 (e.g. changing temperature, rainfall, evaporative demands) and global drivers (e.g. rising [CO₂]) are 68 all recognised as potential causes of increasing woody cover in savannas (Bond & Midgley, 2012). 69 Local land use such as heavy grazing and consequent reductions in fire frequency and intensity 70 promotes encroachment by releasing woody plants from grass competition (O'Connor et al., 2014). 71 Rising [CO₂] can accelerate woody plant growth through increasing available soil water through 72 improving plant water-use efficiencies (Polley et al., 1997; Leakey et al., 2009), potentially extending 73 the growing season, increasing maximum growth rates for individual woody plants while also 74 increasing potential tree cover (where potential tree cover is not limited by other factors, e.g. soil 75 fertility). Increased [CO₂] will also directly increase the rate of carbon uptake by plants, potentially 76 increasing growth rates and the recruitment of woody plants (Bond & Midgley, 2000; Hoffmann et 77 al., 2000; Kgope et al., 2010). The vulnerability of a region to encroachment will therefore depend on 78 both the prevailing land use history, the environmental setting, as well as the functional traits of 79 woody plants that likely govern their responsiveness to both local (e.g. sensitivity to fire) and global 80 drivers (e.g. growth response to elevated $[CO_2]$). However, there are remarkably few experimental 81 data examining the effect of increased [CO₂] on seasonally dry tropical vegetation and the plant 82 functional types within them (but see (Hoffmann et al., 2000; Stokes et al., 2005; Kgope et al., 2010).

83 These gaps in understanding highlight that there has been no systematic review of recent 84 changes in woody cover that have occurred across savannas. While, the number of continental- and 85 global-scale satellite studies of vegetation greenness are increasing, these studies cover only the 86 short time-scale of the satellite record and record measures of greenness (e.g. NDVI) that 87 incorporate the signal of both trees and grasses (Buitenwerf et al., 2015; Liu et al., 2015). We 88 undertake a meta-analysis of studies documenting change in woody cover, with specific attention to 89 studies of woody cover increase, across the tropical and sub-tropical savanna ecosystems of Africa, 90 Australia and Brazil to assess the extent, magnitude and type of encroachment.

91

92 Materials and Methods

93 Data compilation

94 We conducted a meta-analysis of woody vegetation change in natural tropical and sub-tropical C4 grass-dominated savannas in Africa, Australia and South America. We defined savannas as C4 95 96 grasslands with a significant tree component following the definitions provided by Lehmann et al., 97 (2011) and Ratnam et al., (2011). Sites were untransformed by large-scale agriculture, but could have 98 been subject to livestock grazing. Studies documenting land clearing were excluded. We searched the literature using a combination of the following terms: "bush"; "woody", "woody cover" "shrub"; 99 "tree"; plus "thickening"; "encroachment"; "invasion"; "increase"; "change"; "dynamics", "loss", 100 "mortality", "decline", "death", "elephant damage", "elephant impact", "drought", "wind fall", 101 "cyclone" and "storm". Studies were included where woody cover, biomass or tree basal area was 102 103 recorded at two or more points in time. These data were used to derive an annual increment of 104 woody cover change, i.e. (final cover - initial cover) / time in years. Seven studies from the Brazilian 105 cerrado were used where biomass or basal area was recorded. These values were converted to 106 percent cover using a linear regression based on data from (Pinheiro, 2008). In Australia we used

107 converted measures of basal area to canopy cover using data presented by Lehmann *et al.* (2009). In
 108 Africa, similar conversion formulae were not available so only studies considering changes in woody
 109 cover were considered.

110

111	We also compiled studies documenting change in the extent of forest and savanna
112	vegetation. Sites were included if they were untransformed by large-scale agriculture and human-
113	induced afforestation or deforestation. The literature was searched using the terms, "forest",
114	"savanna", "expansion", "shift", "increase", "decrease", "biome switch". Similarly, these studies
115	required two time points. However, across publications there was far less consistency in the metric
116	of change quantifying changes in the extent of savanna. Change was recorded as both change in
117	forest patch size (where the extent of an entire forest patch was measured) or the proportion of
118	ground area covered by forest versus savanna over a given area. We therefore recorded changes as
119	the proportional change of forest relative to savanna but did not use this in the statistical analyses
120	described later.

121

From each study, we compiled site latitude and longitude, and used this information to compile climatic information from WorldClim (Hijmans *et al.*, 2005). Where possible, for each study by site combination we extracted information on the dominant land use at the site, the identity and basic traits of the encroaching woody plant species.

126

127 Uncleared savannas are dominated by three land uses; conservation and commercial and 128 communal grazing land (predominately cattle grazing) (McAlpine *et al.*, 2009). Conservation lands 129 represent the best control situation where natural disturbances caused by regular fires and 130 moderate herbivory (grazers and browser) are mostly maintained. In some areas, elephants are

131 present, which can interact strongly with woody cover, by extensively damaging woody plants 132 (Guldemond & Aarde, 2008). Commercial grazing is primarily centred around livestock production 133 resulting in a grazer-dominated system (Scholes, 2009). In Africa, the other predominant land use is 134 communally-owned subsistence farming (communal rangelands) (Scholes, 2009). In communal 135 rangelands, communities rely on the land for natural resources and grazing. Where there was 136 sufficient information provided, we assigned the site one of these predominant four land use types: 137 conservation (without elephants); conservation (with elephants); commercial grazing; communal 138 grazing. If the land was not used for any particular purpose it was termed remnant land. Where 139 possible, we documented the main encroaching woody species at each site. If plants belonged to the family Fabaceae we recorded if the plants had the potential to fix atmospheric nitrogen (Sprent & 140 141 Platzmann, 2001) (Appendix S4).

142

143 Statistical Analysis

144 Our database consisted of studies documenting both increases and decreases in woody cover over 145 time (Figure 1). We found that studies that documented woody cover declines were often 146 accompanied by direct evidence of the environmental driver of vegetation change (e.g. drought, 147 elephant damage), and that these changes were generally observed over short time periods. Of the 148 110 sites in the database documenting savanna woody change, 13 sites documented declines in 149 woody cover (Appendix S4). Of those sites, only two did not provide direct evidence for the cause of 150 the decline. Therefore, we performed an analysis which aimed to investigate correlates of woody 151 cover increase.

152

Data were analysed using a generalised linear model (Gaussian error family) framework in R (version 3.1.1) (R Core Team, 2014). We included initial woody cover as a covariate representing site level variation. Initial cover (IC) has an important bearing on the potential for a site to increase in

156 woody cover. We expect that sites low in woody cover are more likely to experience higher rates of 157 woody cover increase simply because these sites are less limited by tree-tree competition, and 158 where suppression of disturbance may facilitate rapid increases in woody cover closer to the 159 potential (Roques et al., 2001; Lehmann et al., 2009). Continent was included as a factor given the 160 observed regional differences in the functional relationships between woody vegetation and climate 161 (Lehmann et al., 2014). Further, based on the literature, it is clear that different human interventions have had more or less impact in different regions. Hence, continent could infer differences in both 162 163 human management and ecological setting. Mean annual precipitation (MAP) was a covariate given 164 that both mean woody cover and potential woody cover increase with MAP (Jolly et al., 2005; 165 Sankaran et al., 2005; Lehmann et al., 2014). Finally, we included the first year of a study to examine 166 whether rates of cover change were constant through time. This could help infer whether global 167 change drivers (i.e. [CO₂]) have relevance in interpretation of these data. We included an interaction 168 between initial cover and continent as patterns of woody cover are also known to vary among 169 continents. Unfortunately, we could not include other site level factors such as fire return time or 170 herbivore abundance as these data were inconsistently recorded across studies. Given the variable 171 time periods over which these studies were undertaken, it would be inappropriate to estimate a 172 mean fire return time based on the more recent remotely sensed data.

173

174 Candidate models to describe rate of woody cover increase were compared using Akaike's 175 information criterion, corrected for small sample sizes (AICc) to rank the models. As models with a 176 Δ AICc < 2 are considered well supported, we used model averaging on the models with a Δ AICc < 2. 177 We used the averaged final parameter estimates, standard errors and confidence intervals to 178 demonstrate the effect size of the different parameters. All analysis was performed using the software 179 R (version 3.1.1) (R Core Team, 2014) with the packages 'stats' and 'AICmodavg' version 2.0.3 for 180 model selection and averaging. Following the full factorial model including continent we performed 181 the same analysis for each continent separately. We did this to test the effect of variables, specifically

182 MAP, without confounding MAP and continent as the available rainfall range varies strongly between
 183 ccontinent (Lehmann *et al.*, 2011).

184

We tested for differences in the rate of woody encroachment between comparable land uses within Australia and Africa using a one-way ANOVA. To test for differences in the rates of woody cover between different land uses within Africa we used a two-way ANOVA. Post-hoc analysis was performed using a TukeyHSD test in R (version 3.1.1) (R Core Team, 2014).

189

190 Results

191 Global patterns of woody cover change in C4 grassy ecosystems

192 Of 94 studies covering 110 savanna sites, woody encroachment was apparent in 84% of sites (Figures 193 1–2). Woody encroachment within savannas has occurred across the rainfall gradient in all three 194 regions (Figure 2a). Australia had the lowest average rate of woody cover change of 0.1% per year 195 (1% per decade) (Table 1), where only small net increases in woody cover were recorded (Figure 2b). 196 In Australia, 4 sites recording declines in tree cover were consistently attributed to inter-decadal 197 drought (Appendix 1, 4). Brazil had the highest mean rate of woody cover increase of 0.7% per year 198 (7% per decade) (Table 1, Figure 2b) with only 2 sites recording decreases in untransformed 199 savannas (Appendix 1,4). The average rate of increase in African savannas was 0.25% per year (2.5% 200 per decade) (Table 1, Figure 2b). Africa had the highest number of sites (14) documenting declines in 201 woody cover (Figure 2b), which were directly attributed to either elephants or inter-decadal 202 drought.

Twelve studies, covering 82 sites, examined forest expansion into savanna. Forest expansion
 was recorded at 85% of sites examined (Figure 1, Table 1). In Australia, where the majority of sites

and studies occurred, forest expansion occurred at a rate of 0.7% per year (7% increase in forest
area into savanna matrix, per decade) (72 sites). Here, forest expansion into savanna was usually
attributed to fire suppression. There were significantly fewer studies of forest–savanna boundary
shifts in Africa (5 studies) and Brazil (5 studies), forest expansion was still apparent across all sites. In
Africa, a mean annual increase of 0.3% of forest area per year within the savanna matrix was
observed, and in Brazil a larger mean annual increase of 1.1% forest area per year was recorded
(Table 1).

212 We examined the relevance of continent, mean annual precipitation, start year and initial 213 cover on rates of woody cover increase and identified two plausible models (Table 2a). Model 214 averaged coefficients indicated that continent, mean annual precipitation, start year and initial cover 215 had relevance in these data. Rates of woody cover increase were higher in studies which started 216 later in the 1900s and early 2000s. We found a strong interaction between initial cover and 217 continent, particularly in South America (Table 2b). There, annual rates of increase were highest with 218 low initial woody cover, and thereafter the rate of annual increase in woody cover decreased as 219 initial woody cover increased. We repeated this analysis separately for each continent. In Africa the 220 start year was positively correlated with the rate of encroachment. In South America, observed rates 221 increase were positively correlated with increasing the initial cover. Mean annual precipitation was 222 not an important predictor of rates of woody cover increases in any of the continents.

African savannas can be classified into four land management types (Table 1 and methods). When comparing rates of woody cover increase among land management types, significant woody cover increases occurred across all of these management types except in conservation areas where elephants were present (0.1% per year) ($F_{3,49}$ =2.75, p<0.033) (Table 1). A post-hoc Tukey test confirms that the presence of elephants is associated with significantly lower rates of increase in comparison to communal rangelands and grazed lands.

229 *Contrasting patterns between continents*

Cattle grazing is a land use common across Australia and Africa, enabling a comparison of woody
cover changes across these comparable regions. We found that average rates of change in Africa
were two and a half times of Australian savannas (2.5% vs. 1.1% per decade; Table 1), and the
patterns of woody cover increase were significantly higher in African grazing land when compared to
Australian grazing land (F (1, 27) = 16.04, p<0.000).

The biology of the encroaching species differed between continents. In Africa, 94% of sites (from the 71% of studies reporting such information) were characterised by species with the potential to fix nitrogen (Appendix S4). In South American savannas only 10% of sites were recorded as being characterised by N-fixing species (from the 85% of studies reporting such information). In Australia, none of the encroachment was caused by N-fixing species (from the 30% of studies reporting the dominant woody encroaching species).

241

242 Discussion

We demonstrate that woody encroachment – both within savannas and forest expansion into
savannas – is widespread across the savanna biome. Woody encroachment, albeit with varying
regional magnitude, occurred across regions and land uses. Our meta-analysis of historical records
demonstrates a widespread trend of increasing woody cover, which was already apparent in the
1970s.

We found that initial cover was an important correlate of the rate of encroachment, where a high initial cover was associated with a lower subsequent rate of encroachment. This could potentially be attributed to density dependence (Roques *et al.*, 2001; Lehmann *et al.*, 2009). Mean annual precipitation was not correlated with the rate of encroachment in any region and encroachment occurred across the available rainfall gradient. Water availability is however a key constraint on savanna net primary productivity and maximum woody cover (Sankaran *et al.*, 2005; Staver *et al.*, 2011). It is therefore possible that unquantified site-level variation outweighed the role

255 of rainfall, or that this metric was insensitive to other water-related factors, such as duration of the 256 dry season. Precipitation has been previously found to be an important correlate of encroachment, 257 where in high versus low rainfall comparisons of encroachment, sites at high rainfall are more likely 258 to fluctuate in woody cover over time (Lehmann et al., 2009). Continent was an important correlate 259 of the rate of encroachment where Brazilian savannas have experienced, on average three times the 260 rate of encroachment of African savannas and seven times the rate of Australian savannas. Local 261 differences in fire and browse history, soil nutrient status and physical structure could not be 262 consistently quantified in this analysis as this information was not consistently reported.

263

264 Extensive encroachment across savannas could suggest a uniform global driver of change, 265 specifically elevated [CO₂] that has been variously hypothesised to drive a re-organisation of tropical 266 vegetation. However, our correlative analyses do not preclude regional and local drivers of change 267 generating similar net effects. Therefore, a more compelling approach to test for evidence of a 268 global driver is to examine encroachment across land uses which are expected to have different 269 trends in woody cover over time. In Africa, three land uses are common to untransformed savannas; 270 commercial cattle ranches, communal rangelands and conservation areas. Commercial cattle 271 farming causes a grazer (cattle) dominated system characterised by a moderate but constant 272 stocking rate with an infrequent fire return period (Higgins et al., 1999; Roques et al., 2001). 273 Encroachment is often predicted for this land use type (Wigley et al., 2010). In contrast communal 274 rangelands, subsistence agriculture around rural villages, are predicted to experience declines in 275 woody cover due to wood harvesting for building, energy and income (e.g. charcoal). Rates of woody 276 cover decline are predicted to increase as human densities increase (Banks et al., 1996; Ryan et al., 277 2014). The third land use is conservation, where little change in woody cover is predicted due to 278 regular fires and moderate herbivory (grazers and browsers) maintaining savanna ecosystems. A 279 common distinction within conservation areas is the presence or absence of elephant, the presence 280 of which is predicted to cause declines in woody cover (Guldemond & Aarde, 2008). Our analysis

281 demonstrates that within Africa encroachment has occurred across all these land uses, except in the 282 presence of elephants. This is a powerful indicator that in Africa a global driver is a likely cause of 283 woody encroachment and also highlights the role of mega-herbivory in slowing encroachment. 284 Indeed, the contrast between areas with and without elephants (Table 1) highlights that the removal 285 of elephants is also a likely significant cause of encroachment across Africa (Guldemond & Aarde, 286 2008), as free-roaming elephants have disappeared from many parts of Africa and are now largely 287 confined to conservation areas (Owen-Smith, 1992). In contrast, the mega-herbivore fauna of Australia and Brazil have been extinct for millennia, such that the legacy will not be evident in 288 289 contemporary trends of vegetation change.

290

291 A second indicator for a global process is how rates of encroachment vary with time. We 292 found that in Africa, the later the start year the higher the rate of encroachment. For example, a 10-293 year study starting in 1960 ending in 1970 was likely to have a lower rate of encroachment than a 294 study starting in 1995. Accelerating encroachment rates have been noted in other African studies 295 (Buitenwerf et al., 2012; O'Connor et al., 2014). This is congruent with the encroachment rates 296 responding to an increasing intensity of a global driver, most likely increasing atmospheric [CO₂], 297 increasing land-use intensity and increasing disruption of fire and mega-herbivory (Ramankutty & 298 Foley, 1999; Archibald *et al.*, 2013).

299

300 What is the role of continent?

Encroachment is occurring across all savannas, however the regional context appears linked to observed rates of encroachment. This highlights that regional specific characteristics such as land use and the biology of flora is required to contextualise our findings. For example, human intervention in the contiguity of savannas has been severe in the Brazilian cerrado (Klink & Machado, 2005) and the extraordinary rates of encroachment across Brazil appear a direct consequence of a legislated policy of fire suppression (Klink & Machado, 2005). Fire extent and frequency are further

reduced by landscape fragmentation and transformation with only a few savanna patches larger
than 1000 ha remaining (Durigan *et al.*, 2003; Klink & Machado, 2005; Silva *et al.*, 2008). Therefore,
in Brazil the net impact of local processes is likely causing encroachment at a regional scale across
remaining Brazilian savannas.

311

312 In Africa, savanna encroachment is significantly higher than that observed in Australia. These 313 differences are more pronounced (2.5 times higher) when comparing similar land uses (commercial 314 cattle ranching). Our findings point to a global driver, most likely elevated [CO₂], in African savannas 315 even though this region also has the most records of woody cover declines. Elephant impacts and 316 low soil nutrients, harvesting pressures by humans were instrumental in causing these observed 317 reductions in woody cover particularly in the nutrient poor *Brachystegia* dominated savannas 318 (Mapaure & Campbell, 2002; Ryan et al., 2014). These observations raise the question, what makes 319 Australian savannas relatively resistant to woody encroachment, particularly when no regional land 320 use or policy can explain the changes?

321

322 Australian savannas are characterised by low nutrient soils, highly connected landscapes 323 with little topographic barriers that generate fire regimes of frequent, intense and large fires 324 (Archibald et al., 2013). Frequent fire retards tree growth, biomass and establishment (Murphy et 325 al., 2014) and could buffer encroachment in landscape that has densities of less than 1 person per 326 km²(Australian Burea of Statistics, 2012). Whilst there are regional differences within Australia in 327 both rainfall seasonality and fire regimes (Archibald et al., 2013), Australian savannas generally have 328 extreme seasonality and longer dry seasons than either Africa or Australia (Lehmann et al., 2011; 329 Staver et al., 2011). Although reduced evaporation and improved water use efficiencies from [CO₂] 330 fertilisation could act to increase the duration of the growing season (Donohue et al., 2013), the 331 effect is potentially small, relative to other savanna regions as the dry season remains in excess of six 332 months and will still result in seasonal drought and fire.

334 In contrast with Australia and Brazil, African savannas contain a high abundance of nitrogen-335 fixing woody plant species from the family Fabaceae (Appendix S3). Their dominance could enhance 336 ecosystem level responses to elevated $[CO_2]$ (Leakey et al., 2009) as nitrogen-fixing species can 337 match the elevated rates of photosynthesis with increased nitrogen-fixation (Rogers et al., 2009a) by 338 producing more nitrogen-fixing tissues (Leakey et al., 2009; Rogers et al., 2009a), or a greater 339 nitrogenase activity. Additionally, improved water use efficiencies associated with elevated [CO₂] 340 could lower drought inhibition of nitrogen-fixers (Polley et al., 1997; Rogers et al., 2009b). The role 341 of nitrogen-fixing species in savanna ecology has generally been ignored despite the dominance of 342 these taxa in African savannas. With elevated [CO₂] and/or reduced drought stress, communities 343 with a high proportion of nitrogen-fixing woody species may display rapid increases in biomass over 344 time.

345

346 Regional variation in plant architecture occurs between savannas of Africa, Australia and 347 South America (Dantas & Pausas, 2013; Moncrieff et al., 2014). Regional differences in stem 348 diameter to canopy diameter ratios must change the rate at which tree cover increases per unit of 349 carbon gain. For example, Australian savanna trees are characterised by tall narrow canopies when 350 compared to African savanna trees which have, on average, canopy widths twice that of Australian 351 trees for a given stem diameter (Moncrieff et al., 2014). These architectural differences are likely to 352 result in different rates of encroachment, for example an increase in woody cover from 20% to 40% 353 would require more than twice the carbon gain in Australia as Africa (Moncrieff et al 2014). 354 Therefore, all else being equal, the architectural traits of African savanna trees will promote higher 355 rates of woody cover increase in African than Australian savannas.

Finally, African savannas are dominated by deciduous woody species (Chidumayo, 2001; Bowman & Prior, 2005). In deciduous savanna systems [CO₂] mediated increases in water use efficiency could extend the growing season by alleviating moisture limitation at the beginning or the

end of the growing season. An extended growing season, where leaves are retained for longer
periods, has already been recorded in some semi-arid African savannas (Buitenwerf *et al.*, 2015). A
delay in leaf fall in the early dry season, when average daily temperatures remain moderate, would
extend the photosynthetically active period for plants (Stevens *et al.*, 2015). In contrast, both
Australian and Brazilian savannas are dominated by evergreen woody plant species and changes may
be relatively less pronounced (Bowman & Prior, 2005; Buitenwerf *et al.*, 2015).

365

366 Gaps and bias in the literature

367 This review highlighted notable gaps in information. Across Africa, the *Brachystegia* dominated

368 savannas cover 2.7 million km² (Pienaar *et al.*, 2015) (similar expanse to all Australian savannas) yet

369 there is a paucity of studies in this region. In South America, we could find only 2 studies

documenting woody cover across Venezuelan savannas. On the other hand, other regions were over

371 represented, generating a spatial bias within the dataset, most notably in South Africa. Additionally a

372 publication bias almost certainly exists where studies which do not document either positive or

373 negative change are less likely to be published.

374 Conclusions

375 Encroachment is occurring across the savanna biome. Our results demonstrate that these patterns

376 and their potential causes are not easily extrapolated across savanna regions and that regional

377 context is key to interpretation of these trends. Savannas regions are dominated by different plant

families with different suites of biological traits (Lehmann et al 2014). We argue that the biology of

379 regional floras will influence the susceptibility of a region to encroachment.

380 While, our study allows us to chart trajectories of woody cover change it does not allow us to

determine the underlying causes of encroachment. However, this comparison enables us to

establish testable hypothesis to explain regional variation in encroachment, highlighting the need for

383 experimentation that to date has been lacking in savanna ecology.

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389	Author contributions
390	NS and CL conceived the study, analysed the data and wrote the paper. NS, CL, GD and BM compiled
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392	
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- **Table 1**: Summary of woody cover change studies across savannas and forest–savanna boundaries
- 551 separated by continent and land use.

	No. data points	Mean duration (years)	Mean starting year	Mean change in woody cover (%)	Mean rate of change (% year ⁻¹)	Mean initial woody cover (%) or % forest cover
African forest–savanna African savanna	5 68	34 68	1962 1948	8.5 10.8	0.3 0.25	14.3 20.1
Subsistence grazing land (communal)	25	62	1941	10.6	0.13	20.4
Conservation (without elephants)	6	46	1953	16.5	0.4	23.4
Conservation (with elephants)	19	33	1962	3.2	0.3	33.6
Grazing	17	58.4	1941	17.2	0.3	13.1
Australian forest- savanna	72	42	1961	27.3	0.7	Not reported
Australian savanna	18	45	1951	4.7	0.1	20.3
Conservation	1	40	1964	4.9	0.1	62.7
Grazing	17	45	1951	4.7	0.1	17.0
S. American forest-	5	10	2002	7.5	1.04	59.8
savanna						
S. American savanna	19	18	1984	16.3	0.74	42.4
Conservation	10	13	1994	4.8	0.43	60.3
Grazed	9	23	1973	29.2	1.1	20.0

Table 2a: Best models (where \triangle AICc <2) assessing the relationship between variables and the annual rate of woody cover increase (% yr⁻¹). Models are ranked based on differences in the corrected Akaike's Information Criterion (\triangle AICc). K is the number of estimated parameters, Akaike weight (AICcWt) is the weight of each model. AC = annual rate of change; CONT = continent; SY= starting year, IC = initial cover; T = duration; MAP = mean annual precipitation.

Model	К	ΔΑΙϹϲ	AIC _c Wt	% deviance explained
AC ~ CONT +MAP+SY+ IC + IC:CONT	9	0.00	0.51	17.7
AC ~ CONT+MAP+IC+IC:CONT	8	0.67	0.36	18.32

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Table 2b .Final parameter estimates (β), standard errors (SE) and confidence intervals of model averaging based on top models (see Table 2). Significant parameters i.e. where confidence intervals do not overlap zero, are show in bold. SY = Starting year, AC = annual rate of change; CONT = continent; IC = initial cover; MAP = mean annual precipitation.

Predictors	β	Lower CI (2.5%)	Upper Cl (97.5%)	SE
Intercept	-5.1246	-16.6910	6.4417	5.8590
SY	0.0027	-0.0007	0.0099	0.0030
IC	0.0035	-0.0021	0.0091	0.0028
MAP	0.0004	0.0000	0.0008	0.2639
CONT (Aus)	-0.2190	-0.7443	0.3064	0.3122
CONT (S Am)	0.8764	0.2549	1.4978	0.0002
CONT (Aus): IC	-0.0032	-0.0135	0.0070	0.0051
CONT (S Am):IC	-0.0182	-0.0286	-0.0079	0.0052

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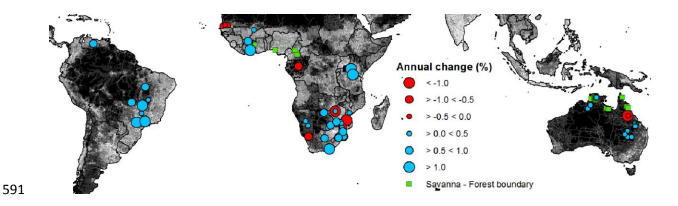
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Table 3 .Final parameter estimates (β) and standard errors (SE) from a GLM performed separately for

575 each continent. Significant parameters are in bold.

	Africa	Australia	South America
Intercent	- 12.83/5.930	-1.703/2.483	23.97/20.42
Intercept			
Start Year	0.006/0.003	0.0009/0.0013 -0.0009/0.0005	-0.0113/0.011 -0.0132/0.0056
Start cover	0.0004/0.0002	0.0009/0.0003	0.0000/0.0012
MAP	-0.000/0.0001	0.0000/0.0001	0.0000/0.0012



592 Figure 1: Sites with reported change in woody cover in tropical C4 grassy ecosystems. Blue circles 593 indicate savanna woody encroachment. Red circles indicate a decline in savanna woody cover. The 594 size of the circle reflects the magnitude of the rate of change in woody cover relative to the duration 595 of the study. Green squares indicate locations where forest expansion into savannas has been 596 reported. Due to the variability in the methods of measurement for change in forest-savanna 597 boundaries, we only denote change without showing the magnitude of change. The background map 598 shading represents the human impact index (HII), where darker shading represents less intervention 599 in the landscape, and lighter shading a higher human impact. The scores provided by the HII range 600 from 0 to 100, and combines socio-economic, roads, land tenure and data on land transformation 601 (Sanderson et al., 2002). Due to the nature of the symbols not all symbols are visible (e.g. forest-602 savanna boundary change in South America). Please see published database for all records (Appendix 603 S1 & S2).

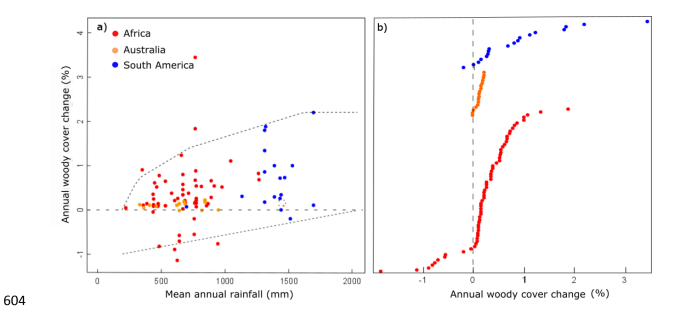


Figure 2a: Annual rate of woody cover change plotted against mean annual rainfall. Upper and lower
dashed lines represent the 5th and 95th percentiles. Positive values indicate woody cover increases
whilst negative values indicate decreases b) Annual rate of woody cover change for each site, shown
in rank order. Points are separated by continent in Africa (red), Australia (orange) and South America
(blue).The dashed vertical line marks the divide between increases and decreases.