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Decadal changes in fire frequencies shift tree communities

and functional traits

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Global change has resulted in chronic shifts in fire regimes. Variability in the sensitivity of 1 2 tree communities to multi-decadal changes in fire regimes is critical to anticipating shifts in ecosystem structure and function yet remains poorly understood. Here, we address the 3 overall effects of fire on tree communities and the factors controlling their sensitivity in 29 4 sites that experienced multi-decadal alterations in fire frequencies in savanna and forest 5 6 ecosystems across tropical and temperate regions. Fire had a strong overall effect on tree communities, with an average fire frequency (1 fire every 3 years) reducing stem density by 7 48% and basal area by 53% after 50 years, relative to unburned plots. The largest changes 8 9 occurred in savanna ecosystems and in sites with strong wet seasons or strong dry seasons, pointing to fire characteristics and species composition as important. Analyses of functional 10 traits highlighted the impact of fire-driven changes in soil nutrients because frequent 11 12 burning favored trees with low biomass nitrogen and phosphorus content and with more efficient nitrogen acquisition through ectomycorrhizal symbioses. Taken together, the 13 14 response of trees to altered fire frequencies depends both on climatic and vegetation determinants of fire behavior and tree growth, and the coupling between fire-driven 15 16 nutrient losses and plant traits.

Ecosystem resilience to changing fire regimes^{1–3} will be a key determinant of how terrestrial ecosystems respond to global change^{3,4}. Fire is a pervasive disturbance, burning ~5% of global land area each year and releasing carbon stored in plant biomass equivalent to 20% of anthropogenic fossil fuel emissions⁵. Historically, much of this carbon is re-sequestered through time as plants recover and regrow, then lost again in the next fire. However, in many systems, changes in climate and land use have shifted fire frequencies, potentially changing the ability of plants, especially trees, to regrow between fires^{1,6–8}.

Although trees can die or be top-killed in fires, prior studies have presented mixed results for the effect of burning frequency on tree biomass^{9–13}, highlighting the need to understand the factors leading to different fire effects. Climate extremes can moderate fire effects on trees by influencing both tree growth and mortality as well as fire intensity: sites with strong wet seasons have trees lacking physiological adaptations to burning¹⁴, and sites with strong dry seasons can have intense fires and droughts¹⁵. Furthermore, trees in different ecosystems respond to changes in fire frequency differently, partly due to fuel load and composition but also their physiology¹⁶. For example, traits conferring physiological protection from heating during fire and/or the capacity to colonize and regrow rapidly could decrease tree biomass losses under frequent burning^{13,17,18}. Tests of these hypotheses using observational data are limited by the collinearity between fire regime and environmental variables at individual sites^{19,20}; instead, tests require experimental manipulations across broad geographic ranges that are long-term, ideally multidecadal, because fire-driven mortality and top-kill can take decades to impact the tree community^{10,13,21}.

38 In addition to external factors, changes within the tree community can modify the effect of repeated fires through the increasing abundance of species more tolerant of the direct and 39 40 indirect effects of fire. Fire effects may be buffered by the colonization and growth of species with structural traits that avoid heat-induced mortality, but also of those with different nutrient 41 42 use and acquisition traits that help plants optimize for fire-driven changes in nutrient availability^{22,23}. Such shifts in nutrient use and acquisition strategies have ecosystem-level 43 44 implications, as changes in these traits significantly influence soil nutrient cycling^{24,25}. For example, plants that form symbioses with ectomycorrhizal fungi, arbuscular mycorrhizal fungi, 45 or nitrogen-fixing bacteria may be better equipped to access nutrients²⁶ that can be depleted 46 under frequent burning²². The distinction between strategies is key because ectomycorrhizal 47 48 plants tend to slow nutrient cycling and productivity, while arbuscular and nitrogen-fixing species accelerate it $^{24-26}$. Tissue stoichiometry is also relevant, with species containing lower 49 nutrient concentrations better equipped to tolerate low nutrient supply but in turn contributing to 50 slower nutrient turnover and lower nutrient availability^{24,25}. Thus, how fire filters for species 51 with different nutrient use and acquisition strategies can influence long-term site productivity. 52

Here, we evaluate how changes in fire frequency alter tree communities, and how 53 climate, vegetation composition, and soils influence the variability in the sensitivity of trees to 54 55 changing fire regimes. Furthermore, we quantify how fire changes the distribution of functional 56 traits in the tree community to evaluate nutrient use traits that are indicative of tree responses to 57 fire-driven changes in nutrients. We combined published data with new surveys on tree populations from 29 sites and 374 plots in four continents. At 27 of the sites (324 plots), surface 58 59 fire frequency was experimentally manipulated for 16-64 years (mean of 30 years), and at two sites (50 plots), natural variation in crown fire frequency presented a natural experiment. We 60

focused on tree responses because of their importance for carbon storage, ecosystem productivity 61 and nutrient cycling^{23,27,28}. The sites cover locations in North America (coniferous, deciduous, 62 and mixed forests and broadleaf savanna), South America (tropical savanna and temperate 63 shrubland), Africa (broadleaf savanna), and Australia (wet and dry eucalyptus forests) all of 64 which are ecosystems that experience frequent burning (Figure S1, Table S1, Supplemental 65 Information, SI). For our main analysis, we define ecosystems broadly based on dominant fuel 66 type (grass vs. litter) and tree functional composition (angiosperm broadleaf vs. gymnosperm 67 68 needleleaf trees) but test the robustness of our conclusions to other classifications. Each surface fire site contains replicate plots including an unburned treatment and different prescribed burning 69 frequencies (Figure S2), where fire frequencies ranged from approximately one fire every decade 70 to one fire every year (Table S1). We evaluated the effects of fire alone and in combination with 71 72 environmental covariates using model selection (SI) to test the importance of climate, soil, and species composition in modifying the effect of fire. 73

74 There were clear overall effects of fire treatments on tree population size. Stem density (stems per hectare) and basal area were lower in frequently burned plots relative to infrequently 75 or unburned plots (Figure 1a for effects in each site and 1b for the response ratio across sites). A 76 77 comparison between the most extreme fire frequency treatments using response ratios illustrated 78 that density and total basal area were 44±25% and 54±25% lower, respectively, in the most 79 frequently burned plots compared with unburned plots. The differences between fire treatment effects were larger when fire frequencies contrasted more (e.g., effect of unburned vs. high fire 80 frequency was 55% greater than the effect of intermediate vs. high fire frequency for both stem 81 density and basal area; Figure 1b,2b, Table S2, errors are 95% confidence intervals). Duration of 82 exposure to altered frequencies was also significant, as sites with longer durations of altered fire 83 frequencies had larger differences between fire treatments, with the slope between duration and 84 community size growing more negative with more frequent burning (density: $F_{1,280}=8.4$, 85 p=0.004, basal area: F_{1,289}=23.3, p<0.001; Figure 1c,2c). For example, relative to unburned plots, 86 87 plots with a three-year fire-return interval had 26% lower stem density and 27% lower basal area after 30 years; the differences increased to 48% lower stem density and 53% lower basal area 88 after 50 years (Figure 1c,2c, Table S3, see Figure S3 for non-transformed results). In annually 89 burned plots, the most extreme fire frequency, burned plots had 63% lower stem density and 90 91 72% lower basal area than unburned plots after 50 years (Figure 1c,2c, Table S3). Thus, both

92 duration of experiment and prescribed frequency help to reconcile the variable effects of fire
93 across studies. Accounting for these factors illustrates that the effects of changing fire
94 frequencies may take a several decades to become substantial, but the impact of the changes will
95 continue to increase for many decades.

96 Fire type was also important, with frequent crown fires affecting tree populations to a 97 greater degree than frequent surface fires. Comparison of 50 plots in needleleaf forests that experienced natural variability in the frequency of stand-replacing crown fires (i.e., wildfires) 98 99 illustrated that stands with shorter fire-return intervals had significantly lower tree densities, especially when plots with the shortest return intervals were considered (F_{1,26.5}=5.2, p=0.03 and 100 101 $F_{1,21}=10.3$, p=0.004, Figure S4). Experimental manipulation of surface fire frequency (i.e., prescribed fires) in needleleaf forests in the USA showed that stem densities were lower in more 102 103 frequently burned plots, but less so than differences caused by frequent crown fires (F_{1,47,1}=17.2, 104 p=0.001, Figure S4). The large effect of short-interval crown fires on tree communities, supported by studies from other regions^{29,30}, highlights the importance of higher fire intensities 105 having more severe effects. 106

107 Although fire frequency had a large overall effect on trees, there was substantial variability in the sensitivity across sites. Part of the variability was attributable to fire history 108 prior to the establishment of the experiment. In forest sites that burned regularly in the decades 109 110 prior to the onset of the experiment, fire exclusion resulted in basal area being 50% ($\pm 17\%$) 111 higher than treatments that maintained historical burning frequencies (p=0.002, Figure S5, Table 112 S1 for site fire histories). In contrast, the reintroduction of prescribed fire into forests that had not burned for several decades prior to the onset of the experiment had relatively minimal effects 113 (p=0.13, Figure S5). These results likely differ from those commonly observed with wildfires, 114 115 which can have larger effects in forests with a history of fire exclusion due to high fuel accumulation^{31,32} because prescribed surface fires are less intense. In savannas, where the fire 116 experiments were all initiated in landscapes that burned regularly in the decades preceding the 117 118 experiment, fire exclusion resulted in basal area increasing by 41% (±20%), but increasing fire frequency resulted in basal area declining by 48% (±16%), relative to an intermediate interval 119 120 that maintained the pre-experiment frequency (statistics from log response ratios $\pm 95\%$ confidence intervals, p<0.001 for both, Figure S5, SI). Taken together, the largest effects of 121

altered fire frequencies were due to fire exclusion in landscapes, especially savannas, that hadburned regularly for the past few decades.

Model selection illustrated that climate, vegetation type, and continent played significant 124 roles in explaining the variability in the sensitivity of trees to fire. Fire effects were largest in 125 126 areas that received more rainfall in the wet season, less rainfall in the dry season, and had lower 127 mean annual temperatures ($F_{1,292,2}=55.2$, p<0.001, $F_{1,284,7}=9.8$, p=0.002, and $F_{1,283,2}=18.1$, p<0.001, respectively) (Figure 3a-c, Table S4, see Table S5 for stem density). For example, plots 128 129 that experienced more frequent burning (2 fires every 3 years, one standard deviation above mean frequency) had 67% lower tree basal area than unburned plots in sites with high wet season 130 131 precipitation. In sites with average wet season precipitation, the difference between the same treatments was only 22% (Figure 3a,S6, Table S4, see SI for details on calculations). Dry season 132 133 precipitation had the opposite correlation with fire effects: sites with lower precipitation in the 134 dry season experienced twice as large an effect of fire on basal area (46% vs. 22% lower tree 135 basal area in sites with low vs. average dry season precipitation Figure 3c, Table S4). The contrasting response to precipitation in the wet vs. dry season is consistent with our 136 understanding that fires are most intense in areas with stronger wet seasons (leading to more 137 fuel) and more severe dry seasons (lower fuel moisture), thus contributing to potential losses 138 139 with more frequent burning^{33–35}. Rainfall in the dry season likely also influences fire effects by determining the water available for tree growth when fire is excluded. Neither soil texture nor 140 soil carbon explained the sensitivity to changing fire frequencies across sites (Table S4). Thus, 141 climate was a key determinant of fire effects, with more moderately seasonal sites being the least 142 sensitive to changing fire regimes. 143

144 The effect of fire on tree basal area also differed across ecosystems, with frequent burning having a larger effect on tree basal area in savannas relative to broadleaf and needleleaf 145 146 forests (F_{2,279}=14.5, p<0.001, model incorporating climate effects; Figure 3d, Table S4). Relative 147 to the unburned plots, basal area in frequently burned plots was 6% lower in needleleaf forests 148 and 22% lower in broadleaf forests (Figure 3d, burn frequency of two fires every three years, SI). In savannas, frequently burned plots had 70% lower basal area relative to the unburned plots 149 150 (Figure 3d, Table S4). Interestingly, stem density responses to fire frequency were qualitatively different between savannas and forests (Table S5). Stem densities increased with more frequent 151

burning in forests while basal area decreased, potentially due to higher light availability and tree 152 recruitment in the forests. To assess the sensitivity of these findings to our classification of 153 ecosystem type, we also tried using a subdivided classification by partitioning broadleaf forests 154 into oak and eucalypt types and needleleaf forests into those that transitioned between oak and 155 pine dominated (Table S1). When included in the final model, the subdivided vegetation 156 classification still had a significant main effect ($F_{4,19,4}=12.4$, p<0.0001), and a significant 157 interaction with fire frequency (F_{4.276.8}=7.8, p<0.001, Figure S8), with basal area in savannas 158 159 responding the most to changes in fire frequency (Figure S8).

Fire-driven changes in basal area and stem density are important for ecosystem function, but fire can also impact ecosystems by changing the functional composition of trees. To address this, we analyzed functional composition in only the experiments from North America (77 tree species, 16 sites, 181 plots) because trait data were available there to (*i*) categorize species by nutrient-acquisition strategies, and (*ii*) assign wood, leaf, and root traits related to growth, survival, and nutrient-use strategies.

We found that structural traits were important for explaining cross-site variability in the 166 167 sensitivity of tree communities to fire. Across sites, frequent burning impacted basal area more where tree species had thinner bark and denser wood (bark: $F_{1,154,3}=5.7$, p=0.018; wood density: 168 F_{1,154.1}=12.9, p<0.001, Table S6, Figure 4a,b). However, within sites, we found mixed evidence 169 170 that fire filtered for species with different structural trait values. Mean wood density of the tree community tended to be lower in frequently burned plots, potentially because of increasing 171 172 dominance of gymnosperm trees, which tend to have lower wood density. In contrast, we did not observe any effect of fire on the mean bark investment of the tree community (Table S6), 173 demonstrating that bark investment at the community scale does not appear to change in 174 175 response to fire. Nevertheless, bark investment helped to predict basal area loss patterns across 176 broad biogeographic scales.

We found evidence for compensatory responses in the nutrient use and acquisition
strategies of the tree communities under different fire frequencies. On average, plots burned
frequently for longer periods of time were dominated by tree species with low nitrogen (N)
concentrations in green and senesced leaves and roots, and which resorbed a greater proportion
of N before leaf senescence (p<0.001 for all variables, Figure 4d, Table S7). This is consistent

with the hypothesis that fire-driven soil N losses²² filters for species with adaptations to low N
conditions. Tissue phosphorus (P) concentrations also declined with frequent burning in leaves
and litter but not in roots (Figure S10, Table S7). Given that P is less prone to being lost than
N^{22,36,37}, this result was surprising and raises questions about how fire may change the P
economy of trees in an opposite way to changes in soil P availability. Taken together, fire filters
for species with more conservative nutrient-use strategies in both their leaves and fine roots
across several North American savannas and forests.

189 Fire also affected the relative abundance of nutrient-acquisition strategies, evaluated by trees' abilities to form symbioses³⁸. Trees that formed symbioses with ectomycorrhizal (ECM) 190 and arbuscular mycorrhizal (AM) fungi were the most abundant nutrient-acquisition strategies 191 across our plots; ericoid and nitrogen-fixing trees were absent from most sites (Figure 4c; 192 categorization based on tree species' taxonomy³⁸). ECM trees, which associate with fungal 193 symbionts capable of acquiring N from organic matter³⁹, tended to be more successful in 194 195 frequently burned plots. The relative abundance of ECM trees increased from 85% in unburned plots to nearly 100% in annually burned plots (Figures 4c, Table S8). ECM trees were also more 196 common in warmer climates and on soils with low carbon concentrations (Figure S11, Table S8). 197 ECM trees typically have lower concentrations of N and P in leaves, litter, and roots than AM 198 199 trees³⁸ (Figure S12, Table S9), suggesting the turnover in symbiont composition may be driving the shift in stoichiometry of the tree community. Since repeated fire tends to decrease inorganic 200 N and N mineralization²³, it makes sense that frequent fire causes the tree community to shift 201 towards species capable of obtaining N from soil organic matter. 202

The tendency for frequently burned plots to have tree communities dominated by ECM 203 species with low N and P content in leaves, roots, and litter indicates that frequent burning favors 204 205 conservative nutrient use and acquisition strategies. This trend is likely a result of fire-driven soil 206 nutrient losses, which should favor species better equipped to cope with low nutrient environments; however, other factors such as phylogenetic conservatism of traits may be at 207 play³⁸. These trait shifts may themselves reinforce an important fire-nutrient feedback if N losses 208 cause a decline in productivity that limits the ability of trees to regrow in between fire events, 209 210 further decreasing ecosystem N. Our results only unpack one part of this feedback loop (fire

effects on traits), and further study is needed to connect changes in strategies with the observeddifferences in nutrient availability and plant growth.

Our analysis also highlights several areas for future work. For one, an improved 213 representation of fire experiments in different ecosystem types across continents in our dataset 214 (e.g., tropical forests in Africa and savannas in Australia) would help further unpack the 215 216 variability across ecosystems and continents. Second, longitudinal data on how fire effects 217 emerge through time could assist with better understanding how the turnover in tree species composition influences changes in total tree cover within experiments^{10,18,40}. Third, considering 218 other plant groups (e.g., herbaceous plants) will help obtain a more comprehensive picture of 219 220 how shifting fire regimes change the structure of the entire plant community. Finally, the extent of fire effects on plant strategies across temperate ecosystems highlights the need for more 221 222 studies of plant strategies to consider fire. For example, the effect of fire on fungal symbiosis 223 strategies should be integrated into theories seeking to explain their biogeographic distribution, 224 which generally rely on climatic factors alone⁴¹.

Our observation that changing fire frequencies shift both tree basal area and density as 225 226 well as trait composition is important because it identifies two means by which fire can indirectly alter carbon and nutrient storage in other ecosystem pools, such as soil organic matter. For one, 227 the observed multi-decadal decline in tree populations mirrors multi-decadal shifts in soil carbon 228 and nitrogen^{22,23} both in timescale and across ecosystems. The greater rates of change in tree 229 230 population sizes in savannas and broadleaf forests are consistent with the higher losses of soil carbon and nitrogen reported in those ecosystems relative to needleleaf forests^{22,42}. Furthermore, 231 a shift towards conservative nutrient use and acquisition traits is consistent lower N turnover^{23,42}. 232 Consequently, our findings support hypotheses that fire-driven changes in tree biomass inputs 233 234 and the turnover of plant traits may both contribute to changes soil carbon and nutrient pools^{42–45}.

In conclusion, widespread changes in fire regimes are likely to shift both the population size and functional composition of tree communities, with both factors affecting the storage and cycling of carbon and nutrients. The effects are not homogenous however, with certain ecosystems being especially sensitive, such as savannas experiencing rapid encroachment of tree cover when fire is excluded. Climatic factors were also key, as regions with extreme precipitation amounts in the wet and dry seasons (high and low, respectively) changed more than areas with moderate precipitation. The effects of fire were not limited to tree population sizes,
but also extended to the functional composition of the community. The convergent response of
frequent burning promoting conservative nutrient use strategies indicates that fire not only
impacts nutrient cycling over decadal timescales, but also suggests that fire likely influences the
evolution of these plant strategies. Consequently, climatically sensitive shifts in fire frequency,
even when relatively low intensity, will alter the structure and functioning of ecosystems through
multiple direct and indirect pathways.

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Data availability statement: All data will be made available online following publication orfreely available from the authors upon request.

Author contribution statement: AFAP and RBJ conceived of and designed the overall study.

252 TR, CA, and CT helped with data acquisition and provided feedback on statistical analyses.

253 DGB, AC, WC, CC, JDH, SHE, WAH, JK, TL, WKM, STO, BP, KGP, PBR, CR, MASS, BCS,

TS, GRS, KS, CS, MGT, and JMV provided data and/or assisted with interpreting the field data

from experiments. All authors contributed to the writing on the manuscript.

Code availability: Full code will be made available upon request to the corresponding author.

257 However, most of the analyses are conducted using code described in the statistical software

258 packages cited within our manuscript.





Figure 1: Fire effects on stem density increase with degree of frequency contrast and length of 261 study duration. **a-b**) log response ratios of stem densities and the surrounding 95% confidence 262 intervals. a) comparisons between log_e (burned/unburned) treatments within each individual site 263 colored by broad ecosystem categorization with burned treatment being the most extreme fire 264 frequency treatments (Table S1, ecosystem groups based on broad differences in fire fuel and 265 266 tree composition, SI). b) averages across all sites comparing among the different levels of fire frequency in studies with ≥ 3 levels; less frequent treatment always in denominator, Table S2). c) 267 partial residuals plot from a mixed effects model including fire frequency, the number of years of 268 269 repeated burning, and their interaction for log_e stem density (Table S3); site was used as a 270 random intercept. Panels are centered on cross-section values of one fire every 10 years, 1 every 271 3 years, and 1 every year but encompass a range of fire frequencies within each panel.



Figure 2: Frequent burning decreases tree basal area and compounds with time of 274 exposure to different fire frequencies. a-b) log response ratios of basal area and the 275 276 surrounding 95% confidence intervals. \mathbf{a}) comparisons within each individual site colored by 277 broad ecosystem categorization in most extreme fire frequency treatments log_e(burned/unburned) (Table S1, ecosystem groups based on broad differences in fire behavior and tree composition, 278 SI). b) averages across all sites comparing among the different levels of fire frequency in studies 279 280 with ≥ 3 levels; less frequent treatment always in denominator, Table S2). c) partial residuals plot from a mixed effects model including fire frequency, the number of years of repeated burning, 281 and their interaction for loge basal area (Table S3); site was used as a random intercept. Panels 282 are centered on cross-section values of one fire every 10 years, 1 every 3 years, and 1 every year 283 but encompass a range of fire frequencies within each panel (ranges: unburned to 1 every 5 284 years; 1 every 5 years to 2 every 3 years; 2 every 3 years to every year). 285

287 **Figure 3**



Figure 3: Climate, ecosystem type, and plant traits modify effects of fire frequencies on tree

basal area. Partial residual plots of the mixed-effects model illustrating the interactive effects
between covariates (site as a random intercept). Panels centered on cross-sectional values from

between covariates (site as a random intercept). Panels centered on cross-sectional values from
one standard deviations around the median (-1, 0, 1). MAT: mean annual temperature. Partial

residual predictions account for the values of all other covariates in the model (i.e., accounting

for collinear effects). Comparisons of rainfall scenarios relative to the mean in the text used wet-

season precipitation of +1 standard deviation above the mean (525 vs. 375 mm yr⁻¹) and dry

season precipitation of -1 standard deviation below the mean (25 vs. 133 mm yr⁻¹). The duration

of experiment held at its mean of 28 years. All model fits are p<0.05; statistics are in Table S4.

298









302 communities. Partial regression plots from mixed-effects models with community weighted

means of (a) bark investment scaled to a 10 cm stem size and (b) wood density (WD) as

304 modifying variables. Bark and wood density were included in the same model and were

negatively correlated $\sigma = -0.86$. Statistics are in Table S6. Basal area is log_e transformed. **c-d**)

306 community weighted means of nutrient acquisition (c) and nutrient use (d) strategies. c) relative

basal area (arc sine transformed) of arbuscular vs. ectomycorrhizal trees as a function of fire

frequency. **d**) green leaf and live root N as a function of study duration. Statistics are in Tables

S7-S9. Data on litter N and resorption are in the Tables and phosphorus content data are inFigure S8.

Supplemental Information

313

314 Experimental design and site descriptions

315 The majority of sites sampled are in ecosystems that experience surface fires (from fire 316 manipulation experiments, n=27). Our main analyses are based on the surface fire experiments, 317 but we compare these data with a network of plots across n=2 sites with natural variability in the frequency of stand-replacing crown fires to evaluate the effect of fire regime. We describe the 318 319 sites briefly in Table S1, and present detailed descriptions of site history in Dataset 1. Most of 320 the data were obtained from existing studies, but we complemented these data with unpublished 321 surveys in the Sequoia and Kings Canyon sites. We identified these sites first by using a list of sites from a previous meta-analysis of multi-decadal changes in fire frequencies²². Not all of the 322 sites in the other study contained vegetation surveys and in some cases the authors of the other 323 324 studies did not share data. We then complemented these studies with other long-term fire 325 manipulation experiments using a literature search and conference presentations. Finally, data from some of the sites were collected specifically for this study. 326

The surface fire experiments mostly are experimental prescribed burn plots. The managers generally try to burn in a broad seasonal window (e.g., a spring fire in North America may occur anytime from March-May) to optimize burn timing for the local fire conditions most suitable to their planned fire intensities. Treatments were not applied in a uniform fashion across sites, which is one motivation for using mixed-effects models (see description of our statistical approach below).

The sites contained different land use histories before the establishment of the experiment, which was not always documented in detail, but we describe key factors in Tables S1 and Dataset 1. We describe how we evaluated the potential role of land use history in *Testing role of fire and land use history* below.

Because these experimental sites utilized different survey methods, the classification of a plant as a tree differed. In some cases, such as savannas with relatively small woody plants, all woody plants over a basal diameter of 5 cm were measured (which includes shrubs). In other cases, stems only above 10 cm diameter at breast height were measured. Consequently, the
definition of 'tree' is based on the local knowledge of what is the relevant size threshold for a
particular ecosystem, and in some cases, it includes all woody plants. Generally, surveys of tree
communities were conducted using the standard methods of belt transects, permanent plots, or
random sampling along a grid.

345 The stand-replacing crown fires are from one extensive ecosystem type that accounts for a large amount of forest fire area in North American temperate regions. Specifically, we used 346 data from 50 plots in lodgepole pine forest in the Western United States¹⁷ (n=50) spanning 347 different elevations and plots along a continuum of fire return intervals¹⁷. Because this ecosystem 348 349 experiences stand-replacing fires, time-since-fire is critical for determining tree abundance because it determines the stage of regrowth. We dealt with this by sampling plots that differed in 350 351 their fire return interval over the past several hundred years but shared the same time since last 352 fire because of a large fire that burned forests with different times since the previous fire. Given 353 the previous study found elevation to be significant, we included elevation categories in the model (<2400 m and >2400 m, respectively). 354

355

356 Choice of plots within 27 sites with surface fires

Within each site, we only used one sampling time period for our analysis. Eight sites 357 358 contained time series data: Cedar Creek, Lombard, Sequoia, Kings Canyon, and the four Kruger sites. For Lombard, we used the surveys from 2002, for Cedar Creek we used 2010, for Kruger 359 we used 1996-98 and for Sequoia and Kings Canyon it varied according to the replicate plots 360 because their most recent surveys occurred in different years. For Cedar Creek, more recent 361 surveys exist, but the outbreak of oak wilt has resulted in large amounts of tree mortality not due 362 363 to fire (Reich *personal communication*). For Highland Rim, we used two different sets of data: the first dataset contains plot-level data (thereby allowing us to determine a variance around the 364 mean) but no tree species identities; the second dataset contains no plot-level data but has 365 treatment-level averages within each tree species, which allowed us to analyze composition 366 367 changes. We utilized the plot-level data for analyses of basal area and stem density. For Morton and Okmulgee, there are not always true replicates in each fire treatment. Morton contains two 368

true replicates for the unburned, but no true replicates for the burned plots. Okmulgee containsno true replicates.

In sites that manipulated burn seasons in addition to frequency, we used a single burn 371 372 season treatment in the analysis in an attempt to match seasons of burns within a particular 373 ecosystem type within a particular region. Specifically, we excluded certain treatments because 374 we wanted consistency in the season of burning across frequency levels within a site. For 375 example, a site that had three levels of fire frequency, may have two levels of season of burn in 376 only one treatment (e.g., unburned, triennial winter burns, triennial summer burns, and annual summer burns). We kept only the season of burns that spanned all the fire frequency treatments 377 378 within a site to avoid confounding frequency and intensity within a site. In North America, we 379 standardized fire season to burning conducted in the winter to early spring because not all sites 380 contained fire treatments with summer burns. For Hitchiti, we used the December-March burns, 381 dropping the June burns. For Kisatchie we used March, dropping July and May. For Kruger, we 382 used August, dropping all the other seasons. For Lombard, we used March-May, dropping the 383 June-Aug. Lamto and La Pampa only contained data on the number of tree stems. Consequently, these were incorporated into the stem abundance analysis only. 384

385

386 Soil chemistry data

387 We collected and analyzed soil data using several methods. First, we determined the 388 dominant soil type using either author descriptions or reported soil texture analysis. Second, we used the highest resolution soil data as possible (e.g., soil samples from each replicate plot within 389 390 a fire treatment), but some sites only contained site-level soil properties. Consequently, we analyzed overall effects of fire on all sites without any covariates, followed by a model that uses 391 392 model selection to account for collinearities among variables when testing for factors that modify fire effects. To extend data on soils across plots, we sampled soils (top 0-5 cm of the mineral 393 horizon) in 24 plots across four sites: Kings Canyon, Sequoia, Limestone Flats, and Chimney 394 395 Springs. Each site contained three replicate plots of an unburned treatment and a high fire frequency treatment. We collected n=5 pseudo-replicates within the true replicate plot, analyzed 396 397 the soils for carbon, nitrogen, and texture, and averaged within each plot. For chemical analyses, 398 soils were sieved to <2 mm, dried to constant weight, and ground on a ball mill. Subsets were

then analyzed using combustion on an Elemental Analyzer at Stanford University; duplicates

400 were run every 10 samples (analytical accuracy was >90%). Texture was analyzed using the

401 hydrometer method (adopted from the Kellogg Biological Station, Long-Term Ecological

402 Research center website, https://lter.kbs.msu.edu/protocols) on sieved soils.

403

404 Climate data

To obtain long-term climate averages at each site, we used WorldClim⁴⁶. WorldClim 405 integrates data from 1970-2000 across 9,000-60,000 weather stations and spatially extrapolates 406 407 the values by integrating other covariates from topography maps and satellite data (described in detail in ref. ⁴⁶). The timespan of the climate data overlaps the duration of the experiments 408 409 reasonably well (mean study initiation = 1983; mean survey year = 2006). Managers timed burning to coincide with consistent weather conditions over the course of the experiment, 410 therefore we did not obtain high resolution inter-annual variability in climate. We focused on 411 several climate variables based on ecologically relevant a priori hypotheses: (i) precipitation 412 413 partitioned into the driest and wettest quarters of the year because precipitation influences fuel accumulation (primarily in the wettest quarter) and fire conditions (primarily in the driest 414 quarter) and (ii) mean annual temperature because of its large effect on a variety of 415 biogeochemical processes. Precipitation in wet and dry quarters are not as correlated with one 416 417 another but are highly correlated with mean annual precipitation and temperature (Table S10).

418

419 Calculation of fire effects in different environmental conditions

Several methods exist to calculate variable importance, with no clear optimal method ⁴⁷. 420 We chose to use the regression coefficients in the model to understand the sensitivity of basal 421 422 area and stem density to changes in relative values of each variable. Importantly, the models were fit to re-scaled data by subtracting each value by the mean and dividing by the standard 423 424 deviation of the variable. Consequently, the product between the mean value of a variable and its 425 coefficient is always zero. Thus, we can compare the relative impact of variables by comparing 426 the magnitude of the fitted coefficients because they reflect the potential change in basal area for 427 a one standard deviation change in a variable value.

To perform meaningful comparisons, we use the standard deviations of variables to 428 illustrate the sensitivity of basal area to a change in the value. For example, using the model to 429 estimate the effect of increasing fire by 1 standard deviation from the mean (mean = 0.34, mean 430 $+1\sigma = 0.67$) tells us the sensitivity of basal area to fire, with all other variables held at their 431 means. Interactions can be tested by moving two variables away from their means: for example, 432 433 changing the fire value in conjunction with precipitation in the wet quarter. Because the model is fit to re-scaled data, the intercept of the model is not representative of the unburned fire 434 treatment, which is calculated by re-scaling the fire frequency data $(0-\mu)/\sigma$, which gives a value 435 of -1.081, making the unburned calculation of 25.6 m^2 ha⁻¹ when all other variables are held at 436 their means. 437

Here are the different levels of comparisons we used in the results and the correspondingfigures.

Wet season precipitation (Figure 3a): wet season precipitation varied one standard deviation
above the mean vs. at the mean (525 vs. 375 mm yr⁻¹, respectively). Fire frequency varied from
unburned to one standard deviation above the mean (2 fires every 3 years).

Dry season precipitation (Figure 3c): dry-season precipitation was one standard deviation below
the mean vs. at the mean (25 vs. 133 mm yr⁻¹, respectively). Fire frequency varied from
unburned to one standard deviation above the mean (2 fires every 3 years).

446 *Vegetation type* (Figure 3d): fire frequency effects were made using two levels of comparisons.

447 Unburned plots vs. burning at the mean frequency (1 fire every 3 years) and unburned plots vs.

burning at one standard deviation above the mean frequency (2 fires every 3 years).

449

450 **Testing overall fire effects**

We first tested the overall effects of the fire treatments across sites with log response ratios using techniques employed meta-analyses^{48,49}. First, we calculated the log response ratio between the different fire frequency categories (low, medium, and high) for basal area and stem density averaged within each category, with the lowest fire frequency in the comparison always in the denominator. Next, we determined the variance based on the number of true replicates within each treatment in a site and the standard deviations within the fire frequency category.
These values across sites were then used to determine the effects of fire treatments on tree basal
area and stem density.

We first evaluated the overall effect of fire frequency and duration that frequency was 459 altered on tree basal area and stem density without considering any potential modifying role of 460 461 covariates. To accomplish this we analyzed (i) a mixed-effects model containing fire frequency, fire period, and their interaction, and (ii) log response ratios of stem density and basal area 462 463 relativized within each site. We excluded the 50 crown fire plots for this initial analysis. We fit the mixed-effects models with site as a random intercept. The statistical design is nested because 464 465 each site has several replicate plots receiving different fire treatments. As a result of this design, the responses to fire at the plot level are likely more related within sites than between sites, 466 467 necessitating a random intercept. Although our design is not balanced (sites differ in their number of replicate plots), models are generally robust to unbalanced designs unless sample 468 sizes are low and/or a random slope is being estimated ⁴⁷, neither of which are applicable here. 469 Models were constructed based on our *a priori* hypotheses of how fire would influence tree 470 population sizes and the potential to interact with covariates. In all cases of mixed-effects 471 models, we tested for model significance using Satterwaith's approximation for degrees of 472 freedom and a Type III ANOVA ⁵⁰. In the event of an insignificant main effect but significant 473 interaction, we tested whether the main effect could be dropped from the model using a change 474 in Aikake Information Criterion (AIC) with a threshold of two. 475

476

477 Comparison between surface vs. crown fire regimes

To analyze the effect of crown vs. surface fire types, we analyzed stem density data from 50 plots (paired within 25 locations) in the Western USA in a separate model. All plots had the same time since fire of 12 years. For this analysis, we used a mixed-effects model to test the relationship between fire return interval and stem density for all locations across the entire return interval span with location as a random intercept. As a further test of fire return interval effects, we selected the short fire return interval (<100 years) in each paired plot and analyzed the relationship with a linear model.

486

Testing the role of fire and land use history

We partitioned studies into three categories based on their disturbance history. Using knowledge of fire history for several decades prior to the fire experiments, we determined if the fire treatments within a site reflected (i) an increase in fire frequency above a historical mean, (ii) fire exclusion after decades of repeated burning prior to the experiment, and (iii) reintroduced fire after decades of pre-experiment fire exclusion (Table S1); the historical mean was defined based on fire activity data for several decades prior to initialization of the experiment (Dataset 1).

493 The fire experiments in the savannas were all initiated in sites that had regularly burned for several decades before the establishment of the experiment. The intermediate fire frequency 494 treatments were reflective of the historical mean, but the most frequently burned plots in those 495 sites were burned at a frequency higher than the historical mean. Consequently, we could use the 496 intermediate frequency plots to evaluate the relative changes due to fire exclusion (unburned vs. 497 498 intermediate) or tree cover declines because of more frequent burning (frequent vs. 499 intermediate). In one savanna site, Marondera, all trees were removed before the onset of the experiment, and consequently we are not able to assume that the difference between the 500 501 intermediate and high frequency treatment is due to declines in trees since the onset of the 502 experiment, rather, it is likely due to a restriction on recovery. Consequently, we omit Marondera 503 from these calculations.

504 The fire experiments in the forests varied in their historical fire frequency and the occurrence of other disturbances. Several sites were in some stage of recovery from previous 505 506 land use (e.g., selective logging, agriculture, etc.), but we focused on the variability in fire 507 history to categorize the sites into fire response categories. We partitioned forests into those that 508 had remained unburned for several decades before the onset of the fire treatments (i.e., 509 reintroduction burns) vs. sites that burned regularly before the experiment. We assume that in the case of the reintroduction burns, changes in tree cover arises from losses due to more frequent 510 511 burning.

512 In the sites that burned regularly prior to the establishment of the experiment, the 513 differences between the unburned plots and those burned at the historical mean was assumed to arise from gains under fire exclusion, and not necessarily increased losses due to frequentburning (although that can clearly occur).

516 We analyze the effect sizes of fire in the comparisons of the unburned vs. intermediate 517 vs. frequent treatments using the same meta-analysis method described above.

518

519 Model selection to determine parsimonious variable combinations

520 For the plots with surface fires, we performed model selection by incorporating covariates of climate, soil, and plant composition into mixed-effects models to test for pairwise 521 522 interactions and possible collinearities (see discussion below of collinearities). Finally, we constructed a full model containing fire, climate, soil, and composition variables based on our 523 524 hypotheses that these factors will interact with fire frequency as well as information gained from the pairwise tests. There were several insignificant effects in the final model, which we tested for 525 526 removal using model selection with a threshold AIC of two. All variables were re-scaled by 527 subtracting the mean and dividing by their standard deviation.

528 Our selection process in the tables illustrates the sensitivity of the final model to the 529 inclusion of additional interactive effects that are not in the final model as well as main effects of 530 the climate, geography, and soil variables. We do not present the exhaustive comparisons 531 because they are not guided by our *a priori* hypotheses of factors modifying fire effects. Soil 532 type was not reported for one location with stem density measurements in South America, so we 533 just use soil carbon content in the model selection analysis.

534

535 Evaluating assumptions of aggregating ecosystem types

The vegetation composition at each site differs substantially, ranging from diverse tropical savannas with dozens of tree species (e.g., Kruger sites) to monodominant coniferous forests (e.g., Limestone Flats and Chimney Springs). Classifying the sites into broad categories was done methodologically, by balancing the need to maintain parsimony (and thus statistical power) with accurately capturing how plant composition may modify fire effects. Consequently, we performed two levels of classification: (i) a coarse categorization separating savannas vs. 542 forests, and within forests treating broadleaf and needleleaf forests separately, which we refer to

as a vegetation type; and (ii) accounting for variability within forest types by partitioning

544 broadleaf forests into Myrtaceae (eucalypt) vs. Fagaceae (oak) dominated, and needleleaf forests

545 into forests that are near completely dominated by needleleaf trees vs. a mixed forest containing

546 both needleleaf and broadleaf trees, which we refer to as a sub-vegetation type.

547

548 Collinearity among climate variables:

Climate variables can be highly collinear, which can inflate the risk of error in statistical 549 inference. To evaluate collinearity, we first determined the Pearson correlation coefficients 550 551 between the main climate variables (precipitation in wet quarter, precipitation in dry quarter, mean annual temperature, mean annual precipitation, and aridity). We excluded variable 552 combinations with a correlation >0.70. Most climate variables relating to water availability were 553 not correlated with mean annual temperature. For water availability, we used precipitation in the 554 555 driest quarter and the wettest quarter because their correlation coefficient was relatively low and 556 they are ecologically more relevant than annual means because they determine the potential productivity in the wet season when most growth occurs but also potential water stress and fire 557 558 conditions in the dry season (Table S10). In contrast, mean annual precipitation and aridity were tightly correlated with one another, as well as with the precipitation values in the separate 559 560 quarters. Consequently, we used precipitation in the different quarters (because they are more relevant for fire dynamics and water limitation of tree growth) and mean annual temperature. 561

562

563 Species classifications and functional traits

Bark thickness data were collected from a dataset in the Fire and Fuels Extension of the Forest Vegetation Simulator. <u>https://www.fs.fed.us/fmsc/ftp/fvs/docs/gtr/FFEaddendum.pdf</u>. Although broad syntheses of bark investment exist for many tree species in North America, not all species contained data from empirical measurements, and thus we used the data from the Fire and Fuels Extension. Bark thickness was assumed to scale linearly with stem diameter, which is generally valid for smaller stems, but it is known bark saturates with increasing stem diameter ⁵¹. The ability of bark investment to predict fire effects will likely improve with better consideration of the non-linear relationship between bark and stem diameter. We evaluate the relative bark
investment, and not absolute bark thickness, which is based on bark investment as well as stem
size.

Wood density was compiled from the literature using a global wood density database ⁵²,
supplemented with additional data ^{53,54}. We assigned a genus-level average for 19 species lacking
data.

Plant tissue stoichiometry and mycorrhizal type were determined using both trait data as
well as phylogenetic trait estimates calibrated to trait data used in a previous global analysis of
plant mycorrhizal traits ³⁸. Full data selection criteria are presented in ³⁸, but we describe them
briefly below.

The plant phylogeny contained >49,000 plant species ⁵⁵. Plant species were added to this phylogeny as needed using the *congeneric.merge* method ⁵⁶. This method uses congeners to add species missing genetic data to the phylogeny, conservatively replacing genera with polytomies where more than one member of the genus is present in the analysis. Our North American dataset used for this trait analysis contained 78 species.

We next generated a species-level phylogenetically estimated trait value for each species 586 587 and trait by fitting models to all data for a particular trait as a function of phylogenetic distance, leaving out each species one at a time using the *phyEstimate* function within the picante package 588 589 for R statistical software⁵⁷. This way, each species trait estimate is based on its own phylogenetic position and a phylogenetic model of evolution (Brownian motion) parameterized without that 590 specific species trait observation. For species without trait data, we estimated trait values based on 591 a model fit to all available trait data. We did not assign trait values to individuals not identified to 592 593 species or for species in genera lacking any stoichiometry data.

594

595 Testing the interactions between species composition and fire

596 To test for fire effects on the relative abundance of symbiotic strategies, we calculated the 597 relative basal area of the different strategies (ectomycorrhizal, arbuscular mycorrhizal and the 598 less abundant ericoid mycorrhizal, non-mycorrhizal, and nitrogen-fixing tree species). Given the 599 low occurrences of ericoid, non-mycorrhizal, and nitrogen-fixing species, we analyzed the 600 relative abundance of arbuscular mycorrhizal and ectomycorrhizal species only. We then fit 601 mixed-effects models with relative basal area as the dependent variable and fire, climate, broad 602 vegetation type (broadleaf, needleleaf, savanna), and soil conditions as the independent 603 variables, each modified by a symbiont term. Relative basal area was arcsine transformed. This 604 analysis was conducted in the North American plots.

To test how functional traits correlated with the effects of fire frequency and duration of experiment, we calculated community trait means in plot j by averaging the traits of each species i by their relative basal area (BA) in a plot. Bark thickness (Bark) for example:

$$\mu_j = \sum_{i=1}^{S} BA_i Bark_i$$

We calculated community weighted means (CWM) for wood density, bark thickness, live andsenesced leaf nitrogen (N) and phosphorus (P) and live root N and P.

We also calculated retranslocation of N and P from a live leaf before senescence using
the data from live and senesced leaf N and P (i.e., not directly measured). Calculations using N
as an example:

614
$$Retranslocation = \frac{N_{green} - N_{senesced}}{N_{green}}$$

Bark thickness was calculated as a scaling coefficient relative to stem diameter (β)

616 Absolute bark thickness =
$$\beta$$
 * stem diameter + α

To test the potential for traits to predict the response of trees to fire, we fit linear mixed-effects

models with the CWM modifying fire effects but allowing for main effects of fire. For example,

$$\begin{array}{ll} 619 & (plot \ basal \ area) \sim FirePeriod + FireFreq + FireFreq: WD_{CWM} + FirePeriod: WD_{CWM} \\ 620 & + FireFreq: BT_{CWD} + FirePeriod: BT_{CWM} + (1|Site) \end{array}$$

621

622To test how fire influenced the trait composition of the community we fit mixed-effects623models to test the effect of both fire as well as environmental factors in explaining the

624 community weighted mean trait values.

We do not include an independent effect of either wood density or bark thickness because we are primarily concerned with how they may modify fire effects.

We also tested for whether the symbiotic strategies differed in their traits. To do so, we
assigned symbiotic strategies and the dominant ecosystem in which they occurred to different
species. We then analyzed linear models incorporating symbiotic strategy and ecosystem type as
additive effects.

631

- Table S1: List of sites with key meta-data. Cont=continent (AU=Australia, NA=North America,
- 634 SA=South America, AF=Africa). Vegetation type presented in broad categories (NL=needleleaf,
- BL=broadleaf) and the families of the dominant tree species. Sites with a pine-dominated ecosystem that
- can change from pine to oak depending on fire regime are noted. Number of plots is the total within the
- entire site. Duration is the number of years over which fire frequencies have differed across plots.
- Frequency is in # fires yr⁻¹. Prior conditions describe the ecosystem type at the beginning of the
- 639 experiment, whether the site experienced regular burning prior to the experiment and if not, how long it
- had remained unburned (reintroduction burns). The asterisks indicate that the sites included data (1, 2)
- 641 previously unpublished in the peer-reviewed literature (n=7).

Site	Cont	Vegetation type	# plots	Duration	Freq.	Ref	Prior conditions
Alapaha	NA	NL Forest, Pinaceae	8	38	0.55	58	Woodland, 2-4yr FRI
	AU	BL Forest,	18	44	0.89	59	Forest, selective log
Bauple*		Myrtaceae					2yr
	NA	Savanna, Fagaceae	16	48	0.82	10	Savanna, No fire
Cedar Creek							40yrs
Chimney Spring	NA	NL Forest, Pinaceae	21	31	1	60	Forest, No fire 60yrs
	AU	BL Forest,	6	25	0.45	61	Forest, selective log
Dry Peachester		Myrtaceae					15yr
Escambia	NA	NL Forest, Pinaceae	12	23	0.50	Unpub.	Woodland, 2-4yr FRI
Highland Rim*	NA	BL Forest, Fagaceae	9	53	1	62	Forest, 2-10yr FRI
Hitchiti	NA	NL Forest, Pinaceae	12	23	0.50	63	Forest, No fire 50yrs
	SA	Savanna,	7	50	0.40	64	Woodland 2-5vr FRI
IBGE		Vochysiaceae					woodiand, 2-5 yr i Ki
	NA	NL Forest,	5	37	0.14	65	Forest No fire >70vrs
Kings Canyon*		Pinaceae/ Fagaceae					1 01030, 110 me - 70y13
	NA	NL Forest,	8	37	0.53	66	Woodland 1-3vr FRI
Kisatchie		Pinaceae/ Fagaceae					woodland, 1 5yr 1 fer
	AF	Savanna,	16	42	1	67	Savanna 5-8vr FRI
Kruger Mopani		Leguminosae					Savanna, 5-6yr i Ki
	AF	Savanna,	16	42	1	67	Savanna 3-Avr FRI
Kruger Pretoriuskop		Combretaceae					Savanna, 5-4yr i Ki
	AF	Savanna,	16	42	1	67	Savanna 5-7vr FRI
Kruger Satara		Combretaceae					Savanna, 5-7 yr T KI
	AF	Savanna,	16	42	1	67	Savanna 3-5vr FRI
Kruger Skukuza		Leguminosae					

	AF	Savanna, Rubiaceae	8	36	1		Savanna, 1yr FRI
Lamto*						68	savanna, Forests No
							fire >50yrs
	SA	Savanna,	6	18	0.28	69	Grass/shrubland, 10yr
La Pampa		Leguminosae					FRI
Limestone	NA	NL Forest, Pinaceae	21	31	1	60	Forest, No fire 60yrs
	NA	BL Forest,	17	16	1	70	Forest, Pest 2yr, No
Lombard*		Fagaceae/ Pinaceae					fire >50yrs
	AF	Savanna,	21	54	1	71	Savanna, clearcut
Marondera		Leguminosae					10yrs
Morton	NA	BL Forest, Fagaceae	4	21	1	72	Forest, No fire ~50yrs
Okmulgee	NA	BL Forest, Fagaceae	8	19	0.53	73	Forest, Unknown
	NA	NL Forest,	6	32	0.10	65	Equat No fine >70
Sequoia*		Cupraceae					rolest, no life =/0yls
	NA	NL Forest,	8	52	0.50	74	Woodland Jyr FDI
Tall Timbers		Pinaceae/ Fagaceae					woodiand, Tyl PKI
University Missouri	NA	BL Forest, Fagaceae	12	64	1	75	Forest, No fire ~20yrs
	AU	BL Forest,	12	39	0.39	61	Forest, selective log
Wet Peachester		Myrtaceae					15yr
Wharton*	NA	BL Forest, Fagaceae	11	50	0.17	76	Forest, 1-10yr FRI
	NA	NL Forest, Pinaceae	25	varied	varied	17	Forest, 7-250yr FRI
YellowstoneLow							and 7-107yr
	NA	NL Forest, Pinaceae	25	varied	varied	17	Forest, 7-395yr and
YellowstoneHigh							7-57yr FRI

644 Table S2: Meta-analysis statistics. The sample size indicates true replicates. The top section analyzes

645 basal area, the bottom analyzes stem abundance.

Variable	Fire	Response	Standard	Zvalua	n value	Lower CL	Upper CI	I
measured	comparison	ratio	error	L value	p value	Lower CI	Opper CI	
Basal area	Low vs. high							
	n=22	-0.78	0.22	-3.53	0.0004	-1.22	-0.35	
	Low vs. mid							
	n=16	-0.40	0.12	-3.41	0.0006	-0.63	-0.17	
	Mid vs. high							
	n=16	-0.43	0.23	-1.86	0.0632	-0.88	0.02	
Stem	Low vs. high							
abundance	n=23	-0.58	0.19	-3.13	0.002	-0.94	-0.22	
	Low vs. mid							
	n=17	-0.25	0.12	-2.12	0.034	-0.48	-0.02	
	Mid vs. high							
	n=18	-0.33	0.20	-1.65	0.0985	-0.73	0.06	

646

647

649 Table S3: Results from mixed-effects model fit to log basal area and stem density (ANOVA for

significance of terms, and then fitted model coefficients) testing the effect of fire frequency (FireFreq),

the length of time plots were exposed to different frequencies (Duration) and their interaction

(FireFreq:Duration). The means and standard deviations used to re-scale the data were: Basal area: fire

653 frequency, mean=0.34, standard deviation=0.32; duration of experiment, mean=28, standard

deviation=19. Stem density: fire frequency, mean=0.35, standard deviation=0.33; duration of experiment,

655 mean=29, standard deviation=19. Units for frequency are fires per-year and duration are years. The main

effect of fire frequency was dropped from the top model based on the AIC being lower. df=degrees offreedom.

Basal area	df		F value	p value	del AIC
FireFreq					-FireFreq=9.4
Duration	1	290.5	94.3	< 0.0001	
FireFreq:Duration	1	288.6	23.3	< 0.0001	
Stem density	df		F value	p value	
FireFreq					-FireFreq=3.8
Duration	1	281.6	47.3	< 0.0001	
FireFreq:Duration	1	279.9	8.4	0.004	
Model coefficients					
	Intercept	FireFreq	Duration	FireFreq:Duration	
Log basal area	2.7408		-0.4268	-0.2204	
	Intercept	FireFreq	Duration	FireFreq:Duration	
Log stem density	6.3765		-0.3427	-0.1515	

658

659

660

661

Table S4: Results from mixed-effects model fit to log basal area a) ANOVA for significance of terms, b)

fitted model coefficients, and c) change in the model AIC with altered additions and removals. All

analyses performed on mean centered and standard deviation scaled data for continuous variables with

site as a random intercept. ANOVA uses Satterthwaite's method to estimate degrees of freedom. Colon denotes interactions. Variable abbreviations are: FireFreq= fire frequency (fires vr⁻¹), Veg=vegetation

668 type (needleleaf forest, broadleaf forest, savanna), MAT=mean annual temperature (°C),

669 PWQ=precipitation in wet quarter (mm), PDQ=precipitation in dry quarter (mm), Duration=length of

time plots have experienced the repeated burning regime (years). For the fitted model coefficients, the

- 671 intercept gives the value for broadleaf forest (so to calculate the basal area in a savanna, you would
- exponentiate the sum of the coefficient of "VegSavanna" and the intercept). See Figures 3, S7 for the
- effects. Independent effects of PWQ, PDQ, and Continent were not included in the model because the
- models did not pass the criterion that an improved model needed to have a >2 AIC difference. C)
 Sensitivity of model to changes in terms illustrates what happens when the model only includes
- 675 sensitivity of model to changes in terms musuates what happens when the model only includes676 interactions and the effect of adding or removing independent effects, as well as the interactions between
- 677 fire and soil. df=degrees of freedom.

a) Type III ANOVA	df		F value	P value	Mean	SD
FireFreq	1	287.5	29.0	< 0.001	0.34	0.32
Veg	2	17.8	1.9	0.172		
MAT	1	18.6	2.7	0.115		
Continent	3	19.2	4.2	0.020	14	4
FireFreq:Veg	2	279.7	14.0	< 0.001		
FireFreq:PWQ	1	285.1	50.3	< 0.001	375	149
FireFreq:PDQ	1	285.6	7.5	0.007	133	108
FireFreq:MAT	1	280.8	16.8	< 0.001	14	4
FirePeriod:Continent	4	283.4	7.2	< 0.001	28	19

b) Fitted model coefficients					
Variable	Estimate	SE	df	t value	P value
(Intercept)	1.8	0.6	17.2	2.9	0.011
FireFreq	-0.1	0.1	284.7	-1.5	0.135
VegetationNeedleleaf	0.2	0.3	18.5	0.8	0.451
VegetationSavgrass	-0.9	0.6	17.1	-1.5	0.142
MAT	-0.2	0.1	18.6	-1.7	0.115
ContinentAustralia	1.6	0.7	17.2	2.4	0.025
ContinentNorthAmer	1.1	0.6	17.1	1.9	0.078
ContinentSouthAmer	1.7	0.6	23.2	2.8	0.010

FireFreq:VegetationNeedleleaf	0.1	0.1	278.8	1.0	0.310	
FireFreq:VegetationSavgrass	-0.5	0.1	281.1	-3.5	< 0.001	
FireFreq:PWQ	-0.4	0.1	285.1	-7.1	< 0.001	
FireFreq:PDQ	0.2	0.1	285.6	2.7	0.007	
FireFreq:MAT	0.2	0.0	280.8	4.1	0.000	
FirePeriod:ContinentAfrica	-0.2	0.1	285.1	-2.1	0.039	
FirePeriod:ContinentAustralia	0.3	0.1	283.2	1.9	0.059	
FirePeriod:ContinentNorthAmer	-0.2	0.1	291.3	-4.3	< 0.001	
FirePeriod:ContinentSouthAmer	-0.2	0.2	274.7	-0.8	0.418	

**Intercept using broadleaf for vegetation

c) Sensitivity of model terms			
Model	df	AIC	ΔΑΙΟ
$\sim\!\!FireFreq\!+\!Veg\!+\!MAT\!+\!Veg\!:\!FireFreq\!+\!PWQ\!:\!FireFreq\!+\!PDQ\!:\!FireFreq$			Тор
+MAT:FireFreq+Continent:Duration+Continent	19	556.3	model
Only interactions	13	582.0	25.76
+PWQ	17	561.0	4.70
+PDQ	17	566.4	10.18
-MAT	16	567.3	11.03
-VegType	15	589.1	32.83
-Continent	16	562.8	6.5394
+SoilType	26	559.3	3.0709
+%C	20	561.4	5.1847
+SoilType:FireFreq	23	581.3	25.06
+%C:FireFreq	17	569.1	25.76

Table S5: Results from mixed-effects model fit to log stem density a) ANOVA for significance of terms,

b) fitted model coefficients, and c) sensitivity of model terms. All analyses performed on mean centered

and standard deviation scaled data for continuous variables with site as a random intercept. ANOVA uses

684 Satterthwaite's method to estimate degrees of freedom. Colons denote interactions. Variable abbreviations 685 are: FireFreq= fire frequency (fires yr^{-1}), Veg=vegetation type (needleleaf forest, broadleaf forest,

savanna), MAT=mean annual temperature (°C), PWQ=precipitation in wet quarter (mm),

687 Duration=length of time plots have experienced the repeated burning regime (years). For the fitted model

688 coefficients, the intercept gives the value for broadleaf forest (so to calculate the basal area in a savanna,

exponentiate the sum of the coefficient of "VegSavanna" and the intercept). See Figures S7, S9 for

690 effects. Model terms are presented relative to the top model (given in the first row), the only interactions

691 refers to all main effects removed. df=degrees of freedom.

a) Type III ANOVA		df	F value	P value		Mean	SD
FireFreq	1	271.8	14.3	< 0.001		0.35	0.33
FireFreq:Duration	1	272.8	5.0	0.026		29	19
FireFreq:Vegetation	2	270.4	32.9	< 0.001			
FireFreq:PWQ	1	272.2	35.5	< 0.001		389	165
FireFreq:MAT	1	270.2	51.5	< 0.001		15	6
Duration:Continent	3	275.7	14.0	< 0.001			
b) Fitted model coefficients							
Variable	Estimate	SE	df	t value	P value		
(Intercept)	6.30	0.28	23.9	22.3	< 2e-16		
FireFreq	0.31	0.08	271.8	3.8	< 0.001		
FireFreq:Duration	-0.10	0.05	272.8	-2.2	0.026		
FireFreq:VegNeedleleaf	-0.19	0.10	270.3	-2.0	0.051		
FireFreq:VegSavanna	-0.85	0.11	270.8	-8.0	< 0.001		
FireFreq:PWQ	-0.21	0.04	272.2	-6.0	< 0.001		
FireFreq:MAT	0.36	0.05	270.2	7.2	< 0.001		
Duration:ContinentAustralia	-0.73	0.18	271.0	-4.1	< 0.001		
Duration:ContinentNorthAmer	-0.42	0.07	275.5	-6.0	< 0.001		
Duration:ContinentSouthAmer	-0.07	0.21	281.6	-0.4	0.719		
Intercept using broadleaf for veget	ation						
c) Sensitivity of model terms							
Model			Df	AIC	ΔAIC		
~FireFreq+ FireFreq:Veg+ FireFre Duration:Continent+Veg+Continer	q:PWQ+FireFre nt	eq:MAT+	17	630.6	0		
Only interactions			12	639.6	9.0		
+PDQ			18	632.8	2.2		
+PWQ			18	631.3	0.6		
+MAT			18	632.5	1.9		

+%C	18	635.3	4.6	
-Veg	15	630.6	0.0	
-Continent	14	637.0	6.3	
+PDQ:FireFreq	18	636.0	5.4	
+%C:FireFreq	18	633.0	2.4	

697 Table S6: Results from mixed-effects models testing for the interaction between traits and fire frequency

treatments. This is split into two levels: (i) testing whether the relationship between fire frequency and

tree basal area across plots were influenced by the community weighted mean trait value in that plot, and

(ii) testing whether fire frequency changed the community weighted mean trait value within plots.Whether the site was a savanna, broadleaf forest, or needleleaf forest was included in the model because

of the large difference in wood traits between needleleaf forests and the other ecosystems. df=degrees of

702 of the large unreferee in wood trans between needlelear forests and the other ecosystems. di-degree 703 freedom.

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Traits modifying fire effects	df		F value	P value
FireFreq	1	154.4	12.9	< 0.001
FireFreq:Bark	1	154.3	5.7	0.018
FireFreq:WD	1	154.1	12.9	< 0.001
Fire modifying trait values				
Bark				
Duration	1	157.3	0.4	0.520
FireFreq	1	156.3	0.2	0.624
Vegetation	2	12.9	7.9	0.006
Wood density				
Duration	1	156.4	5.4	0.022
Vegetation	2	13.0	9.8	0.003

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Table S7: Results from mixed-effects models testing the effect of fire on nitrogen (N) and phosphorus (P)

concentrations in green and senesced leaves, the proportion of N and P retranslocated before senescence,

and root N and P concentrations. Vegetation type was included as a term given the strong differences in

710 traits between needleleaf vs. broadleaf trees. df=degrees of freedom.

Green N	df		F value	P value	Green P	df		F value	P value
Duration	1	158.9	13.8	< 0.001	Duration	1	157.3	11.8	< 0.001
Vegetation	2	12.7	33.6	< 0.001	Vegetation	2	12.6	12.4	0.001
Senesced	N				Senesced	Р			
Duration	1	158.9	19.7	< 0.001	Duration	1	158.3	21.3	< 0.001
Vegetation	2	12.4	19.1	< 0.001	Vegetation	2	12.5	27.2	< 0.001
Retrans N					Retrans P				
Duration	1	157.8	14.2	< 0.001	Duration	1	159.8	24.1	< 0.001
Vegetation	2	12.3	3.0	0.088	Vegetation	2	12.5	43.3	< 0.001
Root N					Root P				
Duration	1	160.0	14.3	< 0.001	Duration	1	157.2	3.1	0.082
Vegetation	2	12.6	30.7	< 0.001	Vegetation	2	12.4	1.1	0.378
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Table S8: Results from top mixed-effects model on the relative abundance of trees summed within a

symbiotic strategy within a plot with site as a random intercept conducted in North America where

taxonomic resolution was the highest (relative basal area was arcsine transformed). The statistics were

719 only run on ectomycorrhizal and arbuscular mycorrhizal groups because they were sufficiently abundant

across plots, but all other strategies (ericoid, non-mycorrhizal, nitrogen fixer) were included in relative

basal area calculation. df=degrees of freedom.

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Variable	df	df	F value	P value
Symb	1	138.3	49.1	< 0.001
Symb:Soil C	2	34.4	23.4	< 0.001
Symb:FireFreq	2	140.1	7.8	< 0.001
Symb:MAT	2	18.6	12.5	< 0.001
Symb:Family	4	14.0	4.4	0.017
Symb:Duration:FireFreq	2	135.0	4.0	0.021

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725 Table S9: Type III ANOVAs on linear models testing the differences in tissue stoichiometry between the

symbiotic strategy grouped in the different ecosystem vegetation types (broadleaf forest, needleleaf forest,

727 or savanna). Inferred statistics are using phylogenetic relationships to infer trait values for species with
 728 missing data (see supporting information and³⁸) while observed are based on direct trait measurements.

missing data (see supporting information and³⁸) while observed are based on direct trait measurements.
 The inferred vs. observed do not refer to the classification of mycorrhizal type. df=degrees of freedom.

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Inferred				Observed			
N green				N green			
n=61	df	F value	p value	n=48	df	F value	p value
Symb	1	12.4	0.001	Symb	1	4.2	0.046
Veg	2	5.4	0.007	Veg	2	4.0	0.025
N senesced				N senesced			
n=61	df	F value	p value	n=25	df	F value	p value
Symb	1	90.8	0.000	Symb	1	1.1	0.310
Veg	2	4.5	0.016	Veg	2	1.5	0.237
N roots				N roots			
n=61	df	F value	p value	n=31	df	F value	p value
Symb	1	15.8	0.000	Symb	1	0.6	0.427
Veg	2	1.8	0.181	Veg	2	0.7	0.489
P green				P green			
n=61	df	F value	p value	n=38	df	F value	p value
Symb	1	8.5	0.005	Symb	1	3.9	0.054
Veg	2	1.8	0.176	Veg	2	5.7	0.007
P senesced				P senesced			
n=61	df	F value	p value	n=21	df	F value	p value
Symb	1	27.8	0.000	Symb	1	6.6	0.018
Veg	2	6.1	0.004	Veg	2	1.6	0.216
P roots				P roots			
n=61	df	F value	p value	n=9	df	F value	p value
Symb	1	11.0	0.002	Symb	1	9.3	0.014
Veg	2	0.6	0.572	Veg	2	1.2	0.346

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- 736 Table S10: Pearson correlation coefficients between climate variables., PWQ=precipitation in wet quarter
- 737 (mm), PDQ=precipitation in dry quarter (mm), MAT=mean annual temperature (°C), MAP=mean annual
- 738 precipitation (mm yr⁻¹). Data derived from WorldClim.
- 739

	PDQ	PWQ	MAT	MAP	Aridity
PDQ		0.28	-0.05	0.82	0.88
PWQ	0.28		0.44	0.77	0.54
MAT	-0.05	0.44		0.19	-0.04
MAP	0.82	0.77	0.19		0.91
Aridity	0.88	0.54	-0.04	0.91	

743 **Figure S1:**



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Figure S1: Distribution of sites. a) map displaying the distribution of sites (dots) with the surface fire 745 sites filled with black and the crown fire sites filled with white. The coloration indicates the average fire 746 747 frequency within a gridcell using ¹. The sample size of plots is written adjacent to the continent. b) distribution of sites in climate space overlying Whittaker's biome distribution ⁷⁷. (1=tundra, 2=boreal 748 forest, 3=woodland/shrubland, 4=temperate grassland/desert, 5=temperate forest, 6=temperate rainforest, 749 7=subtropical desert, 8=tropical forest and savanna, 9=tropical rainforest). Dots colored according to 750 broad vegetation type category. Plots span a mean annual temperate range from 5.2-27.3° C and a mean 751 annual precipitation range from 408-2378 mm yr⁻¹. c) aerial picture of two different fire treatment plots 752 from Cedar Creek, a temperate oak savanna, where different fire frequencies have created a stark biome 753 754 boundary between forests in unburned plots and savannas in biennial burn plots (Pellegrini et al. 2019).

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Figure S2:



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Figure S2: Example of the experimental layout of a fire manipulation experiment taken from Cedar
Creek (a temperate savanna in Minnesota, USA), where fires have been manipulated since 1964. Aerial
imagery (taken in 2017) from the National Agriculture Imagery Program from the Farm Service Agency.
Plots are outlined with a color corresponding to their fire frequencies expressed in terms of number of
fires per year (e.g. 0.33 is one fire every 3 years).



Figure S3: Untransformed data on stem density (a-b) and basal area (c-d) as a function of the duration
that plots have been exposed to burning in the experiment (0=unburned plots). Each dot represents a site
and the dashed lines connect treatments within sites. Columns represent two sets of fire frequency
contrasts comparing unburned vs. the intermediate frequency in a and c, and unburned vs. the high
frequency in b and d (levels defined based on treatments within sites). Dots and bars based on mean and
standard error calculated across the replicate plots within a fire treatment in a site. Note y-axis is on a
log10 scale.







 $F_{1,21}=10.3$, p=0.004) in needleleaf forests with fire expressed in terms of return period (crown fire

plots are all 12 years postfire, data subset to include short-interval burn plots). c) illustrates the

mean response ratios +/- standard error for the fire types with crown fires split into high (>2,400

m) and low (<2,400) elevation sites (Crown 1 and Crown 2, respectively). Analyses were robust to

considering surface fires in only Western US needleleaf forests: $F_{1,47.1}=17.2$, p=0.001. Response ratios were split into long and short fire return interval plots (Crown 1 and 2, respectively), with the justification for definition of interval in ¹⁷.

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Figure S5:



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Figure S5: Log response ratios of basal area for sites categorized based on their different fire histories

prior to the establishment of the experiment. Suppression is calculated for sites where the experiment was

established on historically burned landscapes and is the unburned/historical frequency. Increased

frequency is calculated for sites where the experiment increased fire frequency relative to the historical

average (for forests this was generally the reintroduction of fire into a historically fire suppressed forest).

797 The extremes compare the highest frequency vs. unburned.





Figure S6: partial residual plot displaying the relationship between log_e basal area and precipitation in the

801 wettest quarter cross-sectioned based on fire frequency. This plot is based on the same mixed-effects

model presented in Figure 3 and Table S4, just re-arranged to emphasize how precipitation-basal area

803 relationship changes with more frequent burning.

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Figure S7:



Figure S7: partial residual plot between the length of time plots were exposed to frequent burning and the
log basal area (a) and stem density (b) in the different continents (from the main mixed-effects model
with site as a random intercept in Tables S4-S5).



814 Figure S8: partial residual plot between the length of time plots were exposed to frequent burning and the

815 log basal area in the different sub-vegetation types (from the main mixed-effects model, presented in

816 Table S4 but substituting the broad vegetation effect with the more detailed classification. We found no

817 evidence that accounting for the finer-scale variability in ecosystem classification increased the accuracy

818 of the model or changed our conclusions

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823 Figure S9: Partial residual plots of the mixed-effects model for stem densities illustrating how fire frequency effects changed according to wet-season precipitation, mean annual temperature, and 824 825 ecosystem type. Panels structured by standard deviations around the median to visualize the spread (-1, 826 0,1), PWQ: precipitation in the wet quarter, MAT: mean annual temperature. All model fits are p<0.05 827 and specific results can be found in Table S5. The predictor variables are mean-centered and standard deviations are scaled to facilitate comparisons of variable influence. In needleleaf and broadleaf forests, 828 829 stem densities actually increased with more frequent burning initially, but declined with increasing experiment duration, potentially because of increased light availability initially stimulating recruitment of 830 small trees (Figure S7, Table S5). Stem density in African sites changed little through time (Figure S7). 831 832 The trends in density may reflect the ability of many of the tree species to re-sprout in between fire events78. 833

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Figure S10: Partial residual plots of the phosphorus (P) stoichiometry of community weighted means as a
function of years of repeated burning. Taken from mixed-effects models presented in Table S7. The
models include a vegetation type effect. Tissue P is rescaled by subtracting the mean and dividing by the
standard deviation.



Figure S11: Partial residual (PR) plots of the mixed-effects model between relative basal area different symbiotic groups within a plot (AM=arbuscular mycorrhizal and ECM=ectomycorrhizal on the left- and right-hand side of each panel, respectively). Relative basal area was arcsine transformed. In all panels, the continuous predictor variables were re-scaled by mean centering and dividing by the standard deviation for comparability testing the relationship with fire frequency (a), soil total carbon content (b). mean annual temperature (c). d) illustrates composition across different plant communities grouped based on the family of the dominant tree species (Cup= *Cupressaceae*, Fag=*Fagaceae*, Pin=*Pinaceae*). Dotted lines illustrate median. Statistics are given in Table S8.

858 **Figure S12:**





Figure S12: Box and whiskers plot displaying the tissue stoichiometry for tree species averaged within symbiont strategy (AM= arbuscular mycorrhizal; ECM=ectomycorrhizal) and then grouped according to which overall ecosystem type the species generally occurred in (broadleaf forest, needleleaf forest, or a savanna-grassland). For both N and P, we conducted our comparisons using data that were either based on direct observations ("obs" in the y-axis), or inferred via a phylogenetic relatedness statistical filling ("inf" in the y-axis). Statistics are in Table S9.

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