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Contrasting effects of fire frequency on plant traits of three dominant perennial herbs from Chaco Serrano

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Abstract Fire frequencies are currently increasing in many regions across the world as a result of anthropic activities, affecting ecological processes and plant population dynamics. Fire can generate important changes in soil properties, altering nutrient dynamics and thereby plant growth. Here we analyse fire frequency effects on soil quality and plant traits of three native perennial herbaceous plants (Cologania broussonetii, Desmodium uncinatum and Rhynchosia edulis; Fabaceae) with the capacity for biological N2 fixation that resprouts and are abundant after fire in Chaco Serrano forests. Based on 22-year fire history, we assessed plant traits in sites with low and high fire frequencies along with unburned scenarios. We found significantly lower water content, nitrates and electrical conductivity in frequently burned soils. As a result, the three species showed consistently lower leaf area and specific leaf area in both fire frequencies, implying lower growth rates in comparison to unburned sites. However, total leaf biomass was not affected by fire, leaf phosphorus concentration was higher in R. edulis in high fire frequency and leaf N concentration was twice as large in plants growing in sites of high fire frequency in C. broussonetii and R. edulis. Such an increase in N and phosphorus concentrations is likely a result of both their conservative use of resources and their biological N2 fixation capacity. To our knowledge, this is the first record of such contrasting fire effects observed consistently in three co-occurring species: while plant growth decreased with fire frequency, leaf nutritional traits remain unchanged or increased in frequently burned sites. Quality-depleted and drier soils that result from increased fire frequencies may not only affect trait variation at the intraspecific level but can also drive to a homogenization of the plant community, selecting species with particular combinations of morphological and functional traits.

Key words: biological N2 fixation, fire regime, leaf nutrient, plant growth, specific leaf area.

INTRODUCTION

Fire is a natural and ubiquitous disturbance across terrestrial ecosystems shaping the distribution, structure and dynamics of biomes throughout the world (Bond et al. 2005). The impact of fire on plant communities, and their potential recovery, depends on both the historical fire regime (i.e. the frequency, intensity and seasonal patterns of fire over time) and the life history traits of the plant species adapted to that disturbance regime (Davies 2013). Anthropic activities such as land-use changes, global warming and human population growth can alter fire regimes, changing their frequency and/or intensity, and thus triggering potential irreversible consequences on ecological processes and plant population dynamics (Pausas & Keeley 2009; Keeley et al. 2011). In the face of current fire regime changes, it is crucial to understand how vegetation responds to such increased fire frequency for predicting the consequences on vegetal communities around the world (Pausas et al. 2004).

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depleting soil C and N, but not necessarily P (Certini 2005; Wang et al. 2012; Pellegrini et al. 2015).

Fire effects on soil properties can indirectly act as a selective agent on certain life history traits of plant species (e.g. Keeley et al. 2011). Several studies have evaluated the effects of fire on certain plant functional traits that are closely associated with photosynthesis and growth rates such as leaf area (LA), specific leaf area (SLA), leaf dry matter content (LDMC) as well as N and P leaf concentrations (Paula & Pausas 2006; Anacker et al. 2011; Cianciaruso et al. 2012; Dantas et al. 2013a). Particularly, SLA has been considered a key functional trait because it is an indicator of broad resource-use strategies and ecosystem functioning (Díaz et al. 1998; Anacker et al. 2011). As a resource-acquisition trait, it is modulated by climate, disturbance (including fire) and biotic interactions (Anacker et al. 2011). For example, in areas with soil nutrient limitations caused by fire, plants may show slower growth resulting in lower SLA values (Anacker et al. 2011). Thus, by assessing changes in SLA across contrasting fire frequency conditions in relatively abundant plant species, we may anticipate potential changes in important ecosystem processes such as litter decomposition, productivity or flammability (Díaz et al. 1999; Anacker et al. 2011).

Most research up to now has focused on assessing interspecific plant functional trait responses under natural abiotic environmental variability (e.g. Rosbakh et al. 2015) or disturbances (e.g. Díaz et al. 1999), including fire (Pausas et al. 2004; Saura-Mas & Lloret 2009) to characterize plant communities. Only a handful of studies have assessed the intraspecific variability of plant functional traits associated with either natural (e.g. Albert et al. 2010; Rosbakh et al. 2015) or anthropogenic disturbance regimes (e.g. Prado-Júnior et al. 2015). The few studies assessing fire frequency effects on plant functional traits have found decreased SLA and plant nutrient levels with increased fire frequency (Loiola et al. 2010; Cianciaruso et al. 2012; Dantas et al. 2013a). Increased fire frequency may have an important influence on intraspecific variability at the local scale, affecting nutritional niche breadth of species and system stability (Cianciaruso et al. 2012; Moreira et al. 2012; Carbone et al. 2013; Dantas et al. 2013a). However, because not all plants respond equally to nutritionally stressed environments, studies assessing intraspecific variability in plant functional traits may allow the detection of species-specific responses that otherwise may not be apparent in community-level studies. For example, plant species with the capability of biological N₂ fixation may counterbalance nutrient limitations, especially nitrogen, imposed by high fire frequencies. Species with biological N₂ fixation can play an important role in microenvironmental nutrient cycling dynamics in post-fire ecosystems (Carreira & Niell 1992; Goergen & Chambers 2009).

Native plant communities of Chaco Serrano forest in central Argentina are currently under different levels of exploitation, which are increasingly reducing their diversity (Cabido et al. 2010). In the past decades, the fire regime in this region has been modified by anthropogenic activities and climatic conditions (Bravo et al. 2010). Although humans are responsible for most ignitions, climatic variables, such as annual precipitation, annual potential evapotranspiration and temperature seasonality, are also important drivers of fire frequency (Argañaraz et al. 2015a). A recent study conducted in central Argentina showed that Sierras Chicas was the area most affected by fire, with the highest number of fire events, burned area and fire frequency over the 1999–2011 period (Argañaraz et al. 2015b).

Post-fire floristic assessments of Sierras Chicas region have revealed that several native Fabaceae are an important component of post-fire resprouting and also are used as foraging resources for cattle (Fuentes et al. 2011; Carbone 2015). Among the herbaceous species with the highest relative frequency in the community are the N₂-fixing perennials Cologania broussonetii (Balb.) DC., Desmodium uncinatum (Jacq.) DC., and Rhynchosia edulis Griseb. As many long-lived resprouters, they have a conservative use of resources (Saura-Mas & Lloret 2009). These species not only are present in sites with frequent fires, but they are also the most abundant herbaceous Fabaceae of Chaco Serrano forest (Giorgis et al. 2011). Analyzing the intraspecific variability of plant functional traits of these native herbaceous species in frequently burned environments is a crucial first step for understanding their responses to fire.

In this paper, we analyse the effects of fire frequency on soil properties and on intraspecific variability of plant functional traits of C. broussonetii, D. uncinatum and R. edulis, native perennial N₂-fixing herbaceous plant species abundant in post-fire resprouting of Chaco Serrano forest. We hypothesize that increased fire frequency decreases soil moisture and overall soil fertility, which in turn affects intraspecific plant functional traits such as LA, SLA, LDMC and biomass (Fig. 1). Thus, decreased soil quality induced by frequent fires will positively correlate with decreased intraspecific plant functional traits (except for LDMC; Fig. 1). However, because the species have biological N₂ fixation capacity and an efficient and conservative use of resources, leaf nutrients (N and P) would not be affected by low-nutrient soils in frequently burned scenarios. Finally, because the species share ecological and life-history traits, the overall effects of fire frequency on plant traits at the interspecific level will be similar.

METHODS

Study area

The study was conducted in the eastern hillsides of Sierras Chicas of Córdoba, Argentina. The study area corresponds to
the Chaco phytogeographical province, Chaco Serrano district (Cabrera 1971). The predominant vegetation is a xerophytic subtropical forest, characterized by closed and open forests depending on grazing pressures and fire recurrence (Luti et al. 1979) with tree layer (8–15 m high) dominated by Aspidosperma quebracho-blanco Schltdl., Prosopis spp., Zanthoxylum coco Engl. and Lithraea molleoides (Vell.) Engl. (Luti et al. 1979). Elevation ranges from 400 to 1300 m a.s.l. and the climate is characterized as a warm continental type with the average minimum and maximum temperatures ranging between –10 °C and 26 °C. Precipitation ranges between 500 and 1200 mm per year (Cabrera 1971), with an annual average of 881 mm for the 1990–2012 period within the study sites (Malanca Family, personal communication). Rain concentrates in the warm season, between October and March, delimiting a clear period of water deficit during autumn and winter. Wildfires typically coincide with the end of the dry season, and when temperatures begin to increase, fuel moisture is low and winds reach their maximum speeds (Bravo et al. 2001). In the last 30 years there has been a documented increase in wildfire frequency for the Chaco region, mainly associated with an increase of rains at a regional level, causing a greater accumulation of fine flashy (i.e. high surface area-to-volume) fuels (Miglietta 1994; Bravo et al. 2010). Wildfires in the Chaco Serrano forest are currently mainly caused by humans as a result of accidental or negligent ignitions. Within Sierras Chicas, the fire frequency over the period of 1999–2011 was estimated in one fire every 5 years (Argañaraz et al. 2015b).

**Study species**

Cologania broussonetii (Balb.) DC., D. uncinatum (Jacq.) DC. and R. edulis Griseb. (Fabaceae, Faboideae) are subtropical native perennial herbs widely distributed from southern United States to central Argentina. These species are a main component of the herbaceous stratum of Chaco Serrano forests (Giorgis et al. 2011); thus, their relative abundance is similar across all study sites, ranging from 2.5 to 7.5% total cover (Carbone 2015). They represent important foraging resources for native and livestock grazing animals in tropical dry forests (Fortunato 1983; Vanni 2001). Their growing season begins in late spring and lasts throughout the summer, with individuals growing branches from 0.5 to 2.5 m long. In winter, they undergo a period of dormancy that matches with the dry season where they lose their aerial biomass. The three species have root nodules as corroborated in field observations, with the capacity of biological N₂ fixation. Their underground reserve organs are woody rhizomes and xylopodial roots that can reach 2 to 5 cm in diameter and represent a post-fire regeneration strategy allowing them to survive fire and resprout quickly (personal observations).
Site selection

There is a common problem in most fire studies, which implies that rarely all aspects of fire regimes are considered or controlled (Whelan 1995). While acknowledging such limitation, here, we searched for sites sharing fire frequency and the same time elapsed since the last fire. To accomplish this, we compiled the fire history of the last 22 years from the entire study area using different databases. We used fire records registered by M.A. Giorgis together with Civil Defence of Rio Ceballos city for the period of 1991–2011 (Giorgis et al. 2013). We further corroborated the geographic boundaries of the fire events from vectors of fire in coordinates based on Landsat TN and ETM (30 m spatial resolution) satellite imagery for the 1999–2011 period (Argañaraz et al. 2015a). Finally, we conducted interviews with local area residents combined with in situ observations for final corroboration of fire records. We selected nine sites with different fire regime history in the last 22 years: (i) three ‘unburned’ sites; (ii) three ‘low fire frequency’ sites (1–2 fires); and (iii) three ‘high fire frequency’ sites (3–4 fires) (Table 1; Fig. 2). The sites were georeferenced; the elevation and orientation were obtained from a GPS. At each site, we estimated the stocking rate by counting cattle faeces in an area of 500 m². The stocking rates were low and similar among all study sites. The average distance between sites was 1530 m.

Soil properties

In December 2013, we collected four soil samples (0–10 cm depth) taken randomly nearby the studied individuals at each of the nine study sites. The samples were immediately placed in double plastic bags and stored in a portable Styrofoam cooler to avoid soil dehydration in the field. To obtain gravimetric soil moisture, the samples were weighed immediately after collection and placed in a drying oven at 60 °C until the sample reached a constant weight. Soil water content (%) was calculated as the ratio between water weight (g) and dry soil weight (g). Subsequently, we made a single composite sample from the four samples per site, and took them to the Soil and Water Laboratory of the Faculty of Agronomic Sciences, National University of Córdoba. The soil parameters analysed were (i) organic matter (OM, %); (ii) organic C (%); (iii) total N (%); (iv) C:N ratio; (v) NO₃⁻ (ppm); (vi) extractable P (ppm); (vii) sulphates (SO₄²⁻, ppm); (viii) current pH; and (ix) electric conductivity (EC, dS/m). The analyses were carried out according to (i) organic C by the Walldey–Black wet digestion method; (ii) total N by semi-micro Kjeldahl; (iii) NO₃⁻ using ion selective electrodes; (iv) extractable P by the Bray–Kurtz method; (v) pH in water; and (vi) electrical conductivity (EC; 1:1, soil: water ratio). Soil organic carbon data were multiplied by a factor of 1.72 to give soil OM values (Howard & Howard 1990). The soil texture in the study area was characterized as a sandy loam.

Sampling and measurements

The sampling of individuals was also conducted in December 2013. Twelve adult individuals randomly selected from each of the three species in nine sites belonging to three fire scenarios (36 individuals/species/scenario; totalling 108 individuals

Table 1. Topographic characteristics and fire regime (1991–2013 period) of the nine studied sites in three different fire scenarios of Chaco Serrano, Córdoba, Argentina

<table>
<thead>
<tr>
<th>Map code</th>
<th>Name of site</th>
<th>Range of elevation (m)</th>
<th>Coordinates</th>
<th>Dominant slope aspect</th>
<th>Wildfire years</th>
<th>Number of wildfires</th>
<th>Fire scenario</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>La Quebrada</td>
<td>824–845</td>
<td>31°9′11.73″S 64°20′40.35″W</td>
<td>W</td>
<td>–</td>
<td>0</td>
<td>Unburned</td>
</tr>
<tr>
<td>2</td>
<td>Indarte</td>
<td>976–1000</td>
<td>31°6′38.83″S 64°22′18.54″W</td>
<td>Irregular</td>
<td>–</td>
<td>0</td>
<td>Low fire frequency</td>
</tr>
<tr>
<td>3</td>
<td>Salto</td>
<td>952–965</td>
<td>31°6′29.48″S 64°21′22.38″W</td>
<td>NE</td>
<td>–</td>
<td>0</td>
<td>Low fire frequency</td>
</tr>
<tr>
<td>4</td>
<td>Escuela</td>
<td>980–1000</td>
<td>31°6′52.45″S 64°21′54.10″W</td>
<td>NW</td>
<td>2011</td>
<td>1</td>
<td>Low fire frequency</td>
</tr>
<tr>
<td>5</td>
<td>Site I</td>
<td>1132–1180</td>
<td>31°7′27.30″S 64°22′43.33″W</td>
<td>E</td>
<td>1995</td>
<td>2</td>
<td>High fire frequency</td>
</tr>
<tr>
<td>6</td>
<td>Malanca</td>
<td>1011–1040</td>
<td>31°6′23.70″S 64°23′0.51″W</td>
<td>NE</td>
<td>2003</td>
<td>2</td>
<td>High fire frequency</td>
</tr>
<tr>
<td>7</td>
<td>Site F</td>
<td>1017–1036</td>
<td>31°7′8.16″S 64°21′48.50″W</td>
<td>Irregular</td>
<td>1995</td>
<td>3</td>
<td>High fire frequency</td>
</tr>
<tr>
<td>8</td>
<td>Site A</td>
<td>1180–1206</td>
<td>31°7′49.94″S 64°24′10.49″W</td>
<td>N</td>
<td>1991</td>
<td>4</td>
<td>High fire frequency</td>
</tr>
<tr>
<td>9</td>
<td>Candonga</td>
<td>1030–1040</td>
<td>31°5′38.53″S 64°21′5.79″W</td>
<td>N</td>
<td>1995</td>
<td>4</td>
<td>High fire frequency</td>
</tr>
</tbody>
</table>
per species) were marked with aluminium tags. All sampled individuals from each species were in reproductive stage at the time of sampling and were similar in size, and thereby of approximately similar total plant age. The voucher specimens were deposited in ACOR herbarium of the Faculty of Agronomic Sciences, National University of Córdoba. The studied individuals at each site were separated from each other by a minimal distance of 3 m. In each individual, we randomly selected five fully expanded green leaves of similar age (discarding the base and tip leaves of the branches) concurrently in all sites and performed the following measurements: (i) we measured the leaf area (LA, cm²) by scanning the fresh leaves and then using ImageJ 1.47v software (National Institutes of Health, USA); (ii) we measured the SLA (cm²/g) by dividing leaf area and leaf dry mass; (iii) we measured the LDMC (mg/g) by dividing leaf dry mass and its saturated fresh leaf mass (for the measurements of LA, SLA and LDMC, we followed the protocols detailed by Pérez-Harguindeguy et al. 2013); (iv) we estimated the total leaf biomass (TLB; g) per individual by multiplying the total number of leaves and leaf dry mass; and (v) we obtained leaf total N and (vi) P concentration (mg/g) from the same randomly selected fully expanded normal green leaves of similar age leaves previously mentioned of five individuals from each species per site (N=45 individuals per species). These leaves were ground and then digested in a CuSO₄/K₂SO₄/H₂SO₄ solution for 30 min at 160 °C, and then 1 h at 380 °C in an Easy Block Smart Digester (Westco). After digestion, we proceeded to measure the total concentration of N and P with an automated spectrophotometer SmartChem 200 (Westco).

Data analysis
Analyses were performed in R (R Core Team 2015) and were conducted separately for each plant species. We used linear mixed-effect models (lme function from the nlmef package; Pinheiro et al. 2009) for the quantitative response variables (LA, SLA, LDMC, TLB, N and P) and generalized the linear mixed-effect models (glmer function from the lme4 package; Bates et al. 2014) for soil water content, due to its binomial error distribution. We used fire frequency as the main fixed effect with three levels (unburned, low fire frequency and high fire frequency), and site as a random factor (N=9) nested within fire frequency scenario. The significance of fixed effect was assessed with F-statistics in linear mixed-effect models and with Wald-Z in generalized linear mixed-effect models. For soil properties (except soil water content), which were obtained at the site level, we tested for differences between the three fire frequency scenarios using generalized linear models (glm function from the stats package) for variables with binomial error distribution (OM, C and N) and with F-statistics for the rest of the variables that had normal error distribution. Model parameters were estimated with restricted maximum likelihood and maximum likelihood methods for lme and glme functions respectively. For each species, we tested for relationships between soil properties and leaf traits across all sites with Pearson correlation.

Fig. 2. Location of study sites in the Chaco Serrano forest (Córdoba province) of Central Argentina. Fire frequency scenarios: unburned (sites 1, 2 and 3; squares), low fire frequency (sites 4, 5 and 6; triangle) and high fire frequency (sites 7, 8 and 9; circle).
coefficients. Finally, we assessed the interspecific response patterns among the species for the three most relevant plant traits SLA, leaf N and leaf P. Because the same response variables had quite different magnitudes and ranges of variation among the species, we calculated a new standardized measure (i.e. effect size) for each plant functional trait and used a meta-analytical approach to compare their responses across species (Koricheva et al. 2013). For these analyses we only considered response variables in two contrasting conditions (burned and unburned) and used the unbiased standardized mean difference (Hedges’s $d$) between burned and unburned conditions as the effect size. Thus, we pooled data from high and low fire frequencies and used it as the treatment effect (refer to Koricheva et al. 2013 for calculation details). The effect size $d$ can be interpreted as the inverse-variance-weighted difference in plant functional trait values between unburned and burned conditions, measured in units of standard deviation. Negative values of $d$ imply a decrease in the mean value of the trait in burned conditions, whereas a positive $d$ value corresponds to an increase as compared to the unburned conditions.

RESULTS

Soil properties

We found significant differences in water content, NO$_3$ and EC among fire frequency scenarios, with consistently higher values of these three soil parameters in the unburned conditions (Table 2). Soil water content was similar in both low and high fire frequencies. Soil NO$_3$ levels differed only between the unburned and high fire frequency scenarios. Soil EC differed significantly among the three conditions (Table 2). While the rest of the soil parameters showed consistently higher values in the unburned sites, no statistical significant differences were detected among the three fire frequency conditions. Because water content, NO$_3$ and EC were highly correlated among themselves (Pearson’s $r = 0.72–0.88$; $P < 0.001$), we calculated a new integrated soil quality variable by multiplying these three significant soil variables (refer to Aguirre-Acosta et al. 2014). We then used this integrated soil quality variable (log-transformed) to run correlations with leaf traits.

Intraspecific variability in leaf traits

Leaf area and SLA showed consistently the same response pattern in each of the three species. Leaf area and SLA significantly decreased in low and high fire frequency scenarios in comparison to unburned conditions (Table 3; Fig. 3a–f). In contrast, LDMC showed consistent increases in sites of low and high fire frequencies compared to unburned conditions. These differences were significant only for C. broussonetii (Table 3). Site effect (random factor) explained less than 35% of model variance in all response variables in the three species.

Intraspecific variability in leaf nutrients and TLB

A higher leaf N concentration was observed in C. broussonetii and R. edulis in the high fire frequency scenarios, doubling the levels of the unburned scenarios (Table 3; Fig. 3g, h). In contrast, N values in D. uncinatum were similar across all fire conditions. The leaf P concentration in R. edulis differed significantly between the unburned and the high fire frequency scenarios, with higher P levels in the latter (Table 3; Fig. 3i). In both C. broussonetii and D. uncinatum, the P concentrations were similar among fire scenarios. The estimated TLB values showed non-significant differences among fire frequencies in any of the three species. However, the highest TLB average value in each of the three species coincided with either low or high fire frequency scenarios (Table 3). In all these analyses, site identity always explained less than 35% of the variation in response variables in each species.

Intraspecific relationship between soil quality and leaf traits

In general, the integrated soil quality variable was positive and strongly correlated with LA and SLA for each of the three studied species ($r = 0.6–0.87$; $P < 0.05$). Additionally, we observed that sites with highest soil fertility and leaf trait values were consistently found in unburned conditions (Fig. 4a, b). Leaf dry matter content was negatively correlated with soil quality in the three plant species, but it was significant only for C. broussonetii ($r = −0.86$; $P = 0.005$; Fig. 4c). Leaf N was not related to soil quality in C. broussonetii and D. uncinatum but it was negatively related to soil quality in R. edulis ($r = −0.93$; $P = 0.0007$; Fig. 4d). Leaf P was not related with soil quality in any of the three studied species ($P > 0.05$) (Fig. 4e).

Interspecific response patterns in SLA and leaf nutrients

At the interspecific level, SLA was consistently lower in burned conditions for the three studied species (i.e. confidence intervals did not overlap Hedges’s $d$ cero value). In contrast, leaf N and P significantly increased in burned scenarios, with the exception of D. uncinatum (Fig. 5a) that showed no changes in leaf nutrients (i.e. confidence intervals overlap Hedges’s $d$ cero value). The overall response of SLA to fire across the species was negative and significantly different from cero (Fig. 5b), implying a net decrease in SLA in burned conditions. Leaf N was on average significantly higher

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in burned conditions, and leaf P did not significantly change in burned and unburned conditions across all species, as observed by the overlapping confidence intervals with cero (Fig. 5b).

**DISCUSSION**

The role of fire as modulator of environmental filters is fundamental in structuring and generating plant community and population traits’ variability (Dantas et al. 2013a, 2013b; Ojeda et al. 2010). Natural fire regimes, which involve fire frequency, intensity and time since last fire, can be altered by human activities and seriously affect biodiversity (Keeley et al. 2011). Measuring all of such fire regime aspects simultaneously is seldom achieved in a single study, which is a generalized shortcoming of most fire studies (Whelan 1995). Here we confirmed our initial hypotheses and found that increased fire frequency caused a strong decrease in certain fundamental soil quality parameters, which in turn drastically modified intraspecific leaf functional traits. However, we observed contrasting effects in morphological versus nutritional traits within the same plant species. While LA and SLA decreased, TLB, N, and P were not affected or increased with fire frequency in these three species, as observed by the overlapping confidence intervals with cero (Fig. 5b).

**Table 2.** Soil properties of the three different fire frequency scenarios.

<table>
<thead>
<tr>
<th>Soil property</th>
<th>Unburned</th>
<th>Low fire frequency</th>
<th>High fire frequency</th>
<th>Statistical tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water content (%)</td>
<td>11.05 ± 4.90a</td>
<td>3.36 ± 2.20b</td>
<td>3.18 ± 1.20b</td>
<td>Z = 2.446; P &lt; 0.014</td>
</tr>
<tr>
<td>Organic matter (%)</td>
<td>6.37 ± 0.25</td>
<td>5.23 ± 1.77</td>
<td>5.00 ± 0.70</td>
<td>Z = 0.721; P &gt; 0.471</td>
</tr>
<tr>
<td>Organic carbon (%)</td>
<td>3.70 ± 0.20</td>
<td>3.03 ± 1.08</td>
<td>2.90 ± 0.36</td>
<td>Z = 0.547; P &gt; 0.584</td>
</tr>
<tr>
<td>Total nitrogen (%)</td>
<td>0.27 ± 0.01</td>
<td>0.24 ± 0.07</td>
<td>0.21 ± 0.05</td>
<td>Z = 0.153; P &gt; 0.879</td>
</tr>
<tr>
<td>C:N ratio</td>
<td>13.90 ± 1.41</td>
<td>12.60 ± 1.03</td>
<td>13.80 ± 1.21</td>
<td>F = 0.989; P = 0.425</td>
</tr>
<tr>
<td>Nitrates (ppm)</td>
<td>38.27 ± 19.10a</td>
<td>21.70 ± 6.68ab</td>
<td>8.40 ± 2.46b</td>
<td>F = 4.829; P = 0.050</td>
</tr>
<tr>
<td>Phosphorates (ppm)</td>
<td>6.10 ± 0.50</td>
<td>3.87 ± 2.70</td>
<td>4.17 ± 0.90</td>
<td>F = 1.611; P = 0.275</td>
</tr>
<tr>
<td>Sulphates (ppm)</td>
<td>24.07 ± 15.40a</td>
<td>23.63 ± 5.83</td>
<td>22.23 ± 9.21</td>
<td>F = 0.023; P &gt; 0.977</td>
</tr>
<tr>
<td>pH</td>
<td>6.50 ± 0.30</td>
<td>6.17 ± 0.70</td>
<td>6.30 ± 0.20</td>
<td>F = 0.442; P = 0.662</td>
</tr>
<tr>
<td>Electric conductivity (dS/m)</td>
<td>0.67 ± 0.10a</td>
<td>0.5 ± 0.00b</td>
<td>0.37 ± 0.10c</td>
<td>F = 30.500; P = 0.0007</td>
</tr>
</tbody>
</table>

Values are means ± SD. Different letters in bold indicate statistical significant differences (P < 0.05) in parameter value among fire frequency scenarios.

**Table 3.** Functional traits of three native perennial plant species in three different fire frequency scenarios.

<table>
<thead>
<tr>
<th>Species</th>
<th>Trait</th>
<th>Unburned</th>
<th>Low fire frequency</th>
<th>High fire frequency</th>
<th>Statistical tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cologania broussetti</strong></td>
<td>LA (cm²)</td>
<td>29.08 ± 7.09a</td>
<td>10.97 ± 3.80b</td>
<td>11.99 ± 3.06b</td>
<td>F = 63.294; P = 0.0001</td>
</tr>
<tr>
<td></td>
<td>SLA (cm²/g)</td>
<td>294.85 ± 70.07a</td>
<td>182.05 ± 22.31b</td>
<td>182.48 ± 33.81b</td>
<td>F = 13.088; P = 0.007</td>
</tr>
<tr>
<td></td>
<td>LDMC (mg/g)</td>
<td>263.12 ± 24.02a</td>
<td>305.44 ± 33.92ab</td>
<td>316.76 ± 39.06b</td>
<td>F = 4.954; P = 0.027</td>
</tr>
<tr>
<td></td>
<td>TLB (g)</td>
<td>1.72 ± 0.81</td>
<td>1.46 ± 0.93</td>
<td>2.25 ± 2.02</td>
<td>P = 0.554; P = 0.602</td>
</tr>
<tr>
<td></td>
<td>N (mg/g)</td>
<td>4.03 ± 3.80a</td>
<td>8.01 ± 2.60ab</td>
<td>8.88 ± 2.24b</td>
<td>Z = 4.457; P = 0.035</td>
</tr>
<tr>
<td></td>
<td>P (mg/g)</td>
<td>1.86 ± 1.34</td>
<td>2.58 ± 0.90</td>
<td>2.25 ± 0.72</td>
<td>F = 0.808; P = 0.489</td>
</tr>
<tr>
<td><strong>Desmodium uncinatum</strong></td>
<td>LA (cm²)</td>
<td>31.99 ± 7.68a</td>
<td>20.26 ± 6.68b</td>
<td>21.17 ± 6.90b</td>
<td>F = 9.528; P = 0.014</td>
</tr>
<tr>
<td></td>
<td>SLA (cm²/g)</td>
<td>320.21 ± 78.47a</td>
<td>210.40 ± 32.00b</td>
<td>214.03 ± 31.16b</td>
<td>F = 25.234; P = 0.001</td>
</tr>
<tr>
<td></td>
<td>LDMC (mg/g)</td>
<td>276.41 ± 27.47</td>
<td>304.71 ± 22.07</td>
<td>307.41 ± 23.30</td>
<td>F = 3.150; P &gt; 0.116</td>
</tr>
<tr>
<td></td>
<td>TLB (g)</td>
<td>2.26 ± 1.65</td>
<td>3.13 ± 2.07</td>
<td>3.64 ± 2.09</td>
<td>F = 0.863 P = 0.468</td>
</tr>
<tr>
<td></td>
<td>N (mg/g)</td>
<td>8.36 ± 2.23</td>
<td>8.68 ± 2.64</td>
<td>8.18 ± 2.97</td>
<td>F = 0.232; P = 0.800</td>
</tr>
<tr>
<td></td>
<td>P (mg/g)</td>
<td>2.21 ± 1.03</td>
<td>1.80 ± 0.67</td>
<td>1.62 ± 0.51</td>
<td>F = 0.986; P = 0.426</td>
</tr>
<tr>
<td><strong>Rhynchosia edulis</strong></td>
<td>LA (cm²)</td>
<td>33.11 ± 9.05a</td>
<td>14.98 ± 4.06b</td>
<td>13.44 ± 3.65b</td>
<td>F = 56.186; P = 0.0001</td>
</tr>
<tr>
<td></td>
<td>SLA (cm²/g)</td>
<td>259.59 ± 70.19a</td>
<td>170.71 ± 19.27b</td>
<td>174.07 ± 34.77b</td>
<td>F = 9.776; P = 0.013</td>
</tr>
<tr>
<td></td>
<td>LDMC (mg/g)</td>
<td>354.81 ± 29.15</td>
<td>371.31 ± 26.94</td>
<td>370.60 ± 38.75</td>
<td>F = 1.167; P = 0.373</td>
</tr>
<tr>
<td></td>
<td>TLB (g)</td>
<td>2.84 ± 1.95</td>
<td>3.90 ± 3.16</td>
<td>3.11 ± 2.41</td>
<td>F = 0.311; P = 0.744</td>
</tr>
<tr>
<td></td>
<td>N (mg/g)</td>
<td>2.40 ± 1.45a</td>
<td>3.26 ± 1.49a</td>
<td>4.77 ± 1.20b</td>
<td>F = 11.326; P = 0.009</td>
</tr>
<tr>
<td></td>
<td>P (mg/g)</td>
<td>0.78 ± 0.32a</td>
<td>0.95 ± 0.19b</td>
<td>1.18 ± 0.34b</td>
<td>F = 3.307; P = 0.043</td>
</tr>
</tbody>
</table>

Values are means ± SD. Different letters in bold indicate statistical significant differences (P < 0.05) in parameter values among fire frequency scenarios determined by linear mixed-effect models with fire frequency as fixed effect and site as random factor.

LA, leaf area; LDMC, leaf dry matter content; N, leaf total nitrogen concentration; P, leaf total phosphorous concentration; SLA, specific leaf area; TLB, total leaf biomass.
species. At the interspecific level, as a consequence of sharing ecological and life-history traits, the overall response of the three species was similar.

High fire recurrence enhances soil hydrophobicity affecting water infiltration into soil and increasing runoff events (Certini 2005; González-Pelayo et al. 2015), which leads to soil erosion and overall lower soil moisture (Shakesby & Doerr 2006). Moreover, coarse-textured soils, as found at our study sites, typically have low available water content; thus, small variations may significantly affect plant growth (Albanesi & Anriquez 2003; Thompson & Troeh 1988). Decreased soil water content in frequently burned sites can directly lower SLA as a way of maximizing the limited and scarce water available with a conservative resource use strategy (Pérez-Ramos et al. 2012). Furthermore, decreased EC as a result of high fire frequency results from reduced inorganic ions from the soil’s cation exchange capacity (extractable Mg and Ca), which are essential plant nutrients (Brye 2006). In contrast, total soil N, P, organic C and C:N did not significantly change across burned and unburned sites. These soil characteristics have been observed to be rough proxies of nutrient supply to the vegetation, because most nutrient stocks can be occluded in recalcitrant forms (Aerts & Chapin 2000). Mineralized forms of nitrogen, such as nitrates, together

Fig. 3. Leaf area (a, b, c), specific leaf area (SLA; d, e, f), leaf nitrogen (N; g, h) and phosphorous (P; i) of three native plant species in nine sites with three different fire regimes: unburned (white bars), low fire frequency (light grey bars) and high fire frequency (dark grey bars). Values are means ± standard error.

with the controlling factors of nutrient cycling, such as soil moisture and texture, may be better proxies of plant nutrient supply (Aerts & Chapin 2000; Ordoñez et al. 2009). This may be the situation in our low and high fire frequency scenarios, where available water and nitrates for plants to absorb were seriously compromised, lowering LA and SLA trait values and increasing levels of LDMC consistently in the three studied species. This is the first study to our knowledge that assessed soil quality and intraspecific variability in plant functional traits in different fire frequency conditions. An assessment of fire recurrence on plant functional traits has been documented at the intraspecific level in the Cerrado ecosystem but they did not evaluate soil quality parameters (Loiola et al. 2010; Gianciaruso et al. 2012; Dantas et al. 2013a). These studies concur with our findings of decreased LA and SLA as fire frequency increased. However, these authors also found decreased plant nutrients with increased fire frequency, which contrasts with our results. At the interspecific level, the three studied species responded similarly, which is expected when species are phylogenetically closely related and share similar life-history traits, as observed elsewhere (Anacker et al. 2011; Dantas et al. 2013b).

Many Fabaceae species have different strategies to cope in nutrient-stressed environments, maintaining optimal levels of main nutritional traits with key implications for ecosystem functioning (Carreira & Niell 1992; Goergen & Chambers 2009). For example, long-lived resprouter plants, such as these three native Fabaceae species, have a conservative use of nutrients and allocate more resources to underground organs such as rhizomes, which facilitates regrowth (Saura-Mas & Lloret 2009). Moreover, in resprouter plants, fire eliminates aerial biomass while roots survive. Thus, high fire recurrence increases root:above-ground biomass ratios, allowing resprouters to reach higher foliar nutrient concentrations, regardless of nutrient uptake efficiency or soil fertility (Chapin 1980; Carreira & Niell 1992). An additional mechanism for maintaining and even increasing N leaf nutrient content in particular with increasing fire frequency in these three species can be related to their biological N2-fixation capability. In fact, increased light availability after fire and decreased nitrates in frequent burned sites may favour biological N2 fixation, enhancing the rate of N2 fixation and thereby of plant N content (Carreira & Niell 1992; Casals et al. 2005; Goergen & Chambers 2009). Around 20% of Fabaceae
species are known to establish a symbiotic relationship with N₂-fixing *Rhizobium* bacteria (Sprent & Sprent 1990), and Fabaceae usually show higher frequency, density and diversity of species in post-fire scenarios as a result of their biological N₂-fixation capability (Hendricks & Boring 1999; Guinto et al. 2000). For example, species of the genus *Rhynchosia* (Fabaceae) increased levels of biological N₂ fixation after fire, showing a positive association between absorbed N from the atmosphere and total biomass per plant (Hiers & Mitchell 2007). Such responses can be expected because soil mineral deficiencies rarely affect biological N₂-fixation rates (Raison 1979; Aranibar et al. 2003); in contrast, high soil N availability produces a negative effect on nodulation (Casals et al. 2005), presumably to reduce the high energetic cost of biological N₂ fixation (Aerts & Chapin 2000).

When plant species with biological N₂ fixation have high N content in their leaves, they may contribute to enriching the underground environment through root exudates and decomposition of their tissues (Casals et al. 2005; Reverchon et al. 2012). In sites with low fertility occasioned by recurrent fires, even low rates of biological N₂ fixation may represent a valuable source for restoring soil N levels (Raison 1979; Caon et al. 2014). Thus, biological N₂ fixation represents a key functional trait for restoring soil nutrient cycling in degraded burned areas as has been shown in a few Fabaceae such as *Cologania*, *Rhynchosia* and *Lupinus* (Hiers et al. 2003; Goergen & Chambers 2009; Grau et al. 2009). By maintaining or increasing nutrients in their tissue in burned scenarios, these native species can shape successional dynamics and contribute to nutrient cycling in post-fire microenvironments, improving abiotic conditions and allowing other species to emerge. However, transferring nutrients back to the soil requires time periods long enough to cycle N and P through litterfall (Reverchon et al. 2012). Thus, active management of burned areas is needed to allow sufficient time for the completion of nutrient cycling to restore soil fertility, which may vary depending on the system.

Fig. 5. Interspecific (a) and overall (b) responses of increased fire frequency on specific leaf area (SLA), leaf nitrogen (N) and leaf phosphorous (P). Values are mean and 95% confidence intervals. *Cologania broussonetii* (white), *Desmodium uncinatum* (horizontal strips) and *Rhynchosia edulis* (dotted). Dotted line denotes Hedges’s *d* = 0. Mean values with confidence intervals overlapping zero imply that they are not significantly different from zero.
Over the past decades, there has been a net increase of wildfires and burned areas in certain subtropical and temperate regions, such as forests from eastern Australia (Bradstock 2010) and the Chaco region in central Argentina (Kunst & Bravo 2003; Bravo et al. 2010). Because fire operates as an ecological filter, such changes in fire regime can significantly alter plant community structure and composition, selecting species with particular combinations of morphological and functional traits (Bradstock 2010; Keeley et al. 2011). Thus, plant species with fire-persistence traits are selected and overrepresented in frequently burned communities, thereby delimiting the species composition and plant functional diversity found in a given community (Pausas & Verdú 2008). Currently, as a result of increased frequent fires, many forest ecosystems around the globe are being driven towards savannas, increasing the dominance of herbaceous species and reducing shrub and tree layers (e.g. Bradstock 2010). A similar process is likely to be occurring in our studied burned sites, where the plant community is mostly dominated by herbaceous species. In fact, in burned sites, there is less than 50% of the shrub and tree species found in unburned sites (Carbone 2015). Shrub and tree species remaining in these burned scenarios also share common life-history traits such as resprouting (e.g. L. molleoides, Celtis ehrenbergiana, Condalia spp., Baccharis spp.), and nitrogen-fixing capacities (e.g. Acacia spp, Geoffroea decorticans). Therefore, quality-depleted and drier soils that result from increased fire frequencies in the Chaco Serrano region may not only affect trait variation at the intraspecific level, but can also drive the community to a predominance of plant species with conservative resource-use strategies (Pérez-Ramos et al. 2012; Reverchon et al. 2012). Such processes will lead to a homogenization of the community where slow-growing, resprouter plant species with reduced and sclerophyllous leaves, and a more efficient use of water will be favoured and predominate against fast-growing species, with broad leaves and acquisitive resource-use strategies. In the end, these plant community level changes triggered by recurrent fire will affect important ecosystem processes such as litter decomposition and productivity (Diaz et al. 1999; Anacker et al. 2011).

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The authors declare that they have no conflict of interest.

REFERENCES


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nutrient stoichiometry or composition in tropical savanna. 

handbook for standardised measurement of plant functional 

Pérez-Ramos I. M., Roumet C., Cruz P., Blanchard A., Autran 
economics spectrum' driven by nutrient and water limitations 
in a Mediterranean rangeland of southern France. J. Ecol. 
100, 1315–27.

Pinheiro J., Bates D., Debroy S., Sarkar D. & The R Core team 
(2009) nlme: linear and nonlinear mixed effects models. R 
package version 3, 1–96.

Prado-Júnior J., Schiavini I., Vale V., Lopes S., Arantes C. & 
Oliveira A. P. (2015) Functional leaf traits of understory spe-
cies: strategies to different disturbance severities. Braz. J. 
Biol. 75 (2), 339–46.

R Core Team (2015) R: A Language and Environment for Statistical 
Computing. R Foundation for Statistical Computing, Vienna, 
Austria. Available from URL http://www.R-project.org/

Raison R. J. (1979) Modification of the soil environment by vegeta-
tion fires, with particular reference to nitrogen transformations: 
a review. Plant Soil 51, 73–108.

Reverchon F., Xu Z., Blumfield T. J., Chen C. & Abdullah K. M. 
(2012) Impact of global climate change and fire on the occur-
rence and function of understory legumes in forest 

correlates with temperature: new evidence of trait variation at 
the population, species and community levels. Alp. Bot. 
125 (2), 79–86.

strategy and leaf nutrient content in Mediterranean woody 


sponse of microbial biomass, dissolved organic matter, 
respiration, and N mineralization in mineral soil to fire in forest 

Press, Cambridge, UK.