

Evidence of shift in C₄ species range in central Argentina during the late Holocene

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Received: 1 March 2011 / Accepted: 13 June 2011 / Published online: 14 July 2011
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Abstract

Aim Millennial-scale biogeographic changes are well understood in many parts of the world, but little is known about long-term vegetation dynamics in subtropical regions. Here we investigate shifts in C₃/C₄ plant abundance occurred in central Argentina during the past few millennia

Methods We determined present day soil organic matter $\delta^{13}\text{C}$ signatures of grasslands, shrublands and woodlands, containing different mixtures of C₃ and C₄ plants. We measured past changes in the relative cover of C₃/C₄ plants by comparing $\delta^{13}\text{C}$ values in

soil profiles with present day $\delta^{13}\text{C}$ signatures. We analyzed ^{14}C activity in soil depths that showed major changes in vegetation.

Results Present day relative cover of C₃/C₄ plants determines whole ecosystem $\delta^{13}\text{C}$ signatures integrated as litter and superficial soil organic matter ($R^2=0.78$; $p<0.01$). Deeper soils show a consistent shift in $\delta^{13}\text{C}$, indicating a continuous replacement of C₄ by C₃ plants since 3,870 (± 210) YBP. During this period, the relative abundance of C₃ plants increased 32% (average across sites) with significant changes being observed in all studied ecosystems.

Conclusions Our results show that C₄ species were more abundant in the past, but C₃ species became dominant during the late Holocene. We identified increases in the relative C₃/C₄ cover in grasslands, shrublands and woodlands, suggesting a physiological basis for changes in vegetation. The replacement of C₄ by C₃ plants coincided with changes in climate towards colder and wetter conditions and could represent a climatically driven shift in the C₄ species optimum range.

Responsible Editor: Hans Lambers.

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Keywords C₃/C₄ · Carbon isotopes · Climate change ·
Vegetation dynamics · SOM · Subtropics

Introduction

Paleovegetation reconstructions have shown that millennial-scale climate variability modulates the distribution of ecosystems, affecting global terrestrial biogeography. It is now clear from the palynological

record that synchronic changes in plant species range occurred in the recent geological past as a response to climate change (Allen et al. 2010; Gajewski 2008; Williams et al. 2004). The vast majority of palynological studies, however, have been conducted in cold (mostly northern) regions and little is known about climatically driven vegetation dynamics in tropical and subtropical ecosystems. In South America, for example, palynological studies have been concentrated in cold zones of high altitudes, as unstable sedimentation, fast decay rates and exceedingly complex flora hinder the characterization of fossil pollen in warmer regions (Birks and Birks 2000; Flenley 1985; Marchant et al. 2002; Thouret et al. 1997). In areas where palynological investigation is possible, extrapolations from single isolated pollen profiles are typically used to infer regional shifts in vegetation (Behling and Pillar 2007; Ledru et al. 1996; Mancini 2009) and, for this reason, we still lack a detailed understanding of how past climate-vegetation interactions occurred.

Recent studies have successfully identified past vegetation changes in tropical and subtropical regions by analyzing soil organic carbon isotope ratios (Dümig et al. 2008; Silva et al. 2008, 2010a). Although more limited than palynological records for the purpose of describing changes in populations of individual species or genera, carbon isotopes can be used to trace paleo events at the ecosystem level. This technique is particularly useful to describe local-scale ecotonal shifts in predominant vegetation, where C_3 - and C_4 -dominated systems coexist. Generally, when light is not a limiting factor, plants with C_4 metabolism dominate warm environments, while C_3 plants predominate in cool climates (Sage 2004). The corollary is that C_4 outcompete C_3 plants in the tropics and subtropics, while the converse is observed in colder regions (Bond 2008). Fluctuations in climate, however, are expected to directly influence the balance between C_3 and C_4 plants, imposing alternate dominance of either metabolic pathway, explaining their co-occurrence through various scales of space and time.

When contrasting metabolic pathways also represent contrasting life forms (e.g. trees *versus* grasses), indirect effects of changes in climate may play a fundamental role on vegetation dynamics. For example, frequent and intense fires expected during warmer and drier periods would favor C_4 grasses at the expense of C_3 woody plants in local scales (Behling et al. 2004;

Hoffmann et al. 2003). On the other hand, high atmospheric CO_2 levels, which also follow periods of dry warm climates, could favor C_3 species because of their lower photosynthetic efficiency when compared with C_4 plants (Epstein et al. 1997; Luo et al. 2006; Sage et al. 2010). Recent studies have reported the expansion of C_3 - (forests) over C_4 -dominated ecosystems (savannas and grasslands) during the late Holocene (Behling et al. 2005, 2004; Dümig et al. 2008; Silva et al. 2008). Unfortunately, these studies were conducted where distinct metabolisms represent differences in life form and for this reason they were not able to disentangle direct and indirect effects of climate fluctuations on vegetation change.

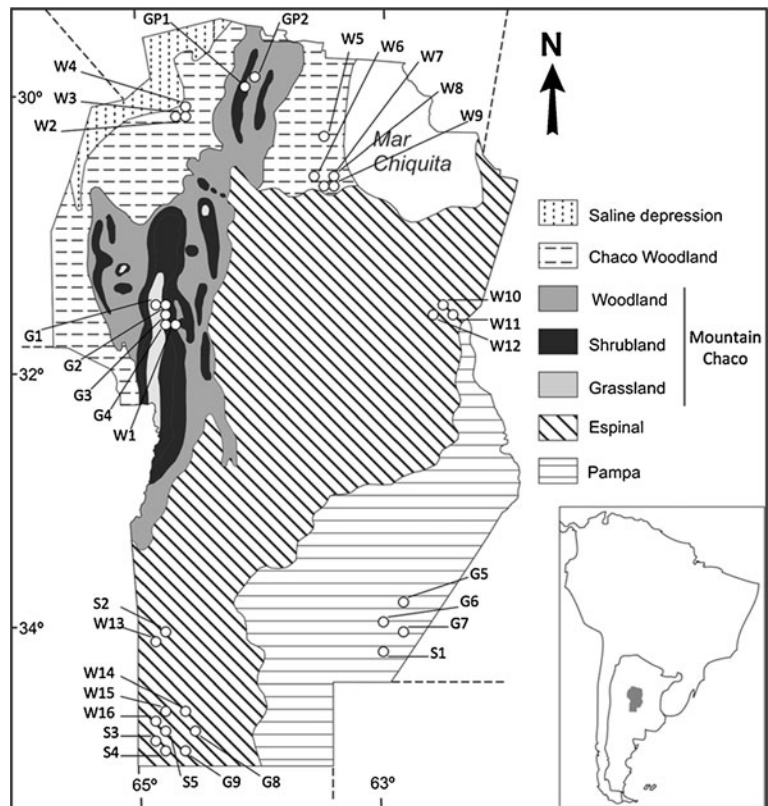
Here we further investigate this issue, asking whether natural climate fluctuations during the Holocene have had a significant effect on subtropical ecosystems, promoting an overall competitive advantage of C_3 over C_4 plants, independently of differences in species life form. To answer this question we sampled a broad region in central Argentina, which corresponds to the present day southernmost limit of C_4 grass distribution within the Americas (Ehleringer et al. 2005). We investigated ecosystems with different structures (grassland, shrubland and woodland) containing various proportions of C_3 and C_4 plants. We quantitatively described shifts on C_3/C_4 vegetation cover occurred in the past using isotopic analysis of organic carbon and discussed the results in relation to previously reported climate reconstructions.

Material and methods

Study region

The present study was conducted in central Argentina, within the limits of Córdoba province (Fig. 1). Both C_4 (grasses) and C_3 (herbaceous and/or woody) plants coexist in the five major regional phytogeographical units: Pampa, Espinal, Chaco, Monte and Pastizal de altura (Cabrera 1976). These distinct vegetation types represent a gradient that encompasses grasslands to the eastern lowlands (~100 m ASL), woodlands towards west at intermediate altitudes (300–500 m), xerophytic open shrublands in the semi-arid western highlands (800–1,200 m) and grasslands at higher altitudes (above the tree line, 1,600 m) (Fig. 1).

Fig. 1 The vegetation of Córdoba Province, Argentina, as represented by its most important phytogeographical regions. Circles show the approximate location of the study sites, where W; S and G and GP represent woodland; shrubland, grassland, and grassland with palms respectively



According to the Köppen-Geiger classification, the regional climate is temperate/warm-temperate to subtropical, in average ranging from 10°C to 24°C between winter and summer (summer-time highs ~30°C and winter lows ~4°C), with a marked rainfall gradient from east (annual rainfall >1,000 mm) to west (annual rainfall <400 mm), with 70% of the rainfall occurring from November to March (Cabido et al. 2008).

Vegetation survey

The regional landscape has been severely altered by human activity throughout the past centuries, but relicts of undisturbed vegetation can still be found. Using our own unpublished and other authors' published recent floristic surveys (Cabido et al. 1993, 1997, 2008; Diaz et al. 1994, 1999, 2001; Perez-Harguindeguy et al. 2000; Pucheta et al. 1998; Zak and Cabido 2002), we selected 32 well-preserved sites where the vegetation is representative of the most important regional ecosystems. We classified the distinct vegetation types in four major categories:

woodlands (16 sites); shrublands (5 sites); grasslands (9 sites) and grasslands with palms (2 sites) (W, S, G and GP sites indicated in Fig. 3). The number of sites representing each of these vegetation categories varied according to their floristic complexity. Woodlands comprise both xerophytic and mesophytic ecosystems with high diversity of woody and non-woody plants, dominated by *Aspidosperma* and species of *Prosopis*, while shrublands only represent xerophytic ecosystems dominated by species of *Geoffroea*, *Condalia* or *Acacia* genera. Grasslands encompass both C₃- and C₄-dominated treeless vegetation. In a couple of grassland sites the presence of palms of the species *Trithrinax campestris* was remarkable and for this reason these sites were classified as grassland with palms.

In previous studies we used leaf area index (LAI) measurements to characterize C₃/C₄ relative cover across vegetation gradients (see for example: Silva et al. 2008, 2010a). These studies were conducted, however, in regions where different strata represent distinct metabolic pathways (e.g. herbaceous layer is dominated by C₄ grasses, while shrubs and trees are

C₃). In such conditions, LAI measurements (which do not differentiate between species) are sufficient to describe changes in the relative cover of C₃ and C₄ plants. On the contrary, in the region studied here perennial C₃ and C₄ herbaceous plants can be equally abundant (Zuloaga et al. 2008) and predominant metabolic pathways can only be assessed by detailed floristic surveys. For this reason, we performed a complete census, including all herbaceous, shrub and tree species at each studied site. Our survey was conducted during the growing season (Summer) of 2010, when maximum productivity is typically attained and all potential species are present. The relative contribution of each species for the total vegetation cover was estimated (visual estimation—projection of canopy cover in 400 m² plots—3 plots per site) and from these estimates the relative cover of C₃ and C₄ species was calculated (Cabido et al. 1997, 2008). Plants that have the C₄ photosynthetic pathway were distinguished from those that have the C₃ pathway by examination of the Kranz anatomy in cross-sections of fresh and herbarium specimens and from the literature (Sánchez and Arriaga 1990; Smith and Epstein 1971). A list of the surveyed species, including growth habit and metabolic pathways, is presented in Appendix 1. More details about the regional vascular flora can be found at Zuloaga et al. (2008), but see also their on line updates at: www.darwin.edu.ar.

Isotopic signatures and past vegetation changes

To test whether and how much C₃ and C₄ relative vegetation cover has changed we relied on two well-known natural processes: (i) during photosynthesis C₃ plants discriminate more against CO₂ molecules that contain the stable isotope ¹³C than do C₄ plants. For this reason different proportions of C₃/C₄ contribution to the total biomass yield differences in the mean carbon isotopic ratios ($\delta^{13}\text{C}$) in the soil organic matter (SOM) (Ehleringer et al. 2000; Marshall et al. 2007; Silva et al. 2010a, b, 2008; Smith and Epstein 1971; Von Fischer et al. 2008); (ii) SOM in soil profiles represents a chronological sequence of vegetation signature, with past vegetation recorded at deeper levels (Dümig et al. 2008; Nordt et al. 2007; Sanaiotti et al. 2002; Silva et al. 2010a, 2008; Victoria et al. 1995). Based on these we characterized past shifts in vegetation structure by, firstly,

determining the current vegetation cover at each study site (described in the previous section), secondly, by determining the carbon isotopic signatures ($\delta^{13}\text{C}$) in the litter and superficial SOM and, finally, by examining the $\delta^{13}\text{C}$ of SOM at different depths in soil profiles.

We used 3 to 5 soil profiles at each site to sample: fresh litter (any decaying organic matter found at the uppermost soil layer), superficial (0–1 cm depth) and deep soil (10, 20, 30, 50 and 100 cm depth). Prior to analysis litter was dried for two days at 70°C and coarsely ground, while superficial and deep soils were dried at room temperature after which fine roots were removed by sieving through a 0.8 mm mesh and not analyzed. The carbon isotope ratios of litter and soil samples were determined at the Laboratory of Stable Isotope Ecology (LSIETE) at the University of Miami. Samples (10 mg) were loaded in tin cups, which were placed in an automated elemental analyzer connected to a continuous flow isotope ratio mass spectrometer. ¹³C abundances were expressed as:

$$\delta^{13}\text{C} (\text{‰}) = (R_{\text{sample}}/R_{\text{PDB}} - 1) \times 1000$$

where R_{sample} and R_{PDB} represent the ¹³C/¹²C ratios and PeeDee standard respectively. The precision of analysis was $\pm 0.1\text{‰}$ (\pm SD).

We used regression analysis to describe the effect of present day C₃/C₄ relative vegetation cover on litter and superficial SOM $\delta^{13}\text{C}$ values (signatures). Based on the equation that best described this relationship (Fig. 2) we estimated past C₃/C₄ relative cover, by

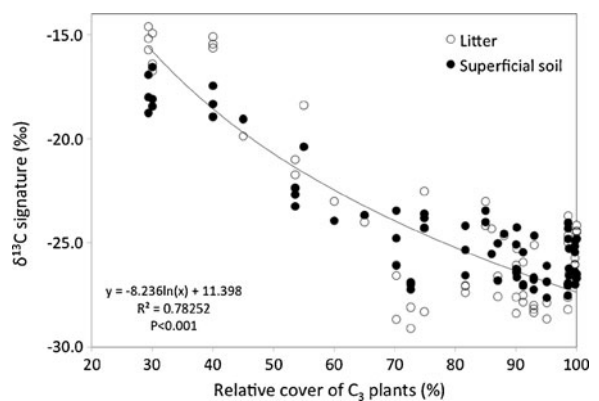


Fig. 2 Relationship between the relative vegetation cover of C₃ plants and the isotopic composition ($\delta^{13}\text{C}$) of litter and superficial soil organic matter (0–1 cm depth), across all study sites

examining SOM $\delta^{13}\text{C}$ signatures at deeper SOM in soil profiles. We estimated past vegetation cover for each study site and soil depth (1, 10, 20, 30, 50 and 100 cm). By calculating percent differences between the C_3/C_4 relative cover estimated in superficial (0–1 cm) and in deep soils, where major shifts in vegetation were observed (50 cm), we determined the total vegetation change. We also measured the slope (angular coefficient) of least square regressions, performed between estimates of C_3/C_4 cover and their respective soil depth at each study site. All soil depths were used for this calculation, here named degree of vegetation change. We used two-way analysis of variance (ANOVA) to test the effect of ecosystem, altitude and their interaction on both metrics of vegetation change (total percent and degree of vegetation change) across sites. Analysis of variance and regression analysis were performed according to standard methods, using the software JMP Version 9 (JMP 2009). For brevity, only statistically significant results are presented.

Carbon dating

We analyzed ^{14}C activity of carbon in the SOM of the soil profiles that showed major shifts in C_3/C_4 vegetation cover at each ecosystem (grassland, shrubland and woodland: G4, G8, GP1, S1, W7 and W13; all at 50 cm depth; Fig. 1). Because roots from the modern day vegetation can exude recent ^{14}C into the deeper soil matrix, dates acquired by analyzing SOM ^{14}C may be underestimated. For this reason, SOM ^{14}C measurements should be considered as a proxy for determining the minimum possible date of past vegetation changes (Dümig et al. 2008; Martinelli et al. 1996; Silva et al. 2010a, 2008; Trumbore 2000; Victoria et al. 1995). Analyses of ^{14}C activity were performed using accelerator mass spectrometry at the Beta Analytic Radiocarbon Dating Laboratory (Miami, USA). The results were then calibrated to represent actual calendar years of the mean age of SOM. The calibration was performed using the newest (2004) calibration database (Reimer et al. 2004). Calibrated dates expressed as years before present (YBP) were consistent among sites/ecosystems and here we present the average date and standard deviation of measurements that represent major vegetation shifts across sites.

Results

Current vegetation cover and $\delta^{13}\text{C}$ signatures

Differences in the present day relative contribution of C_3 and C_4 plants to total vegetation cover explain variations in whole ecosystems' $\delta^{13}\text{C}$ signatures, integrated as litter and superficial soil organic matter ($R^2=0.78$; $p<0.01$) (Fig. 2). Similar relationships have been found in previous studies conducted in different ecosystems (Lloyd et al. 2008; Silva et al. 2010a, 2008) and represent an expected gradient determined by variations in the amount of C_3 - and C_4 -originated biomass and their mixture. Due to intrinsic differences in discrimination, C_3 plants $\delta^{13}\text{C}$ signature (ranging from -20 to -35‰) are unmistakably distinguishable from C_4 plants $\delta^{13}\text{C}$ signature (ranging from -9 to -15‰) (Epstein et al. 1997). The $\delta^{13}\text{C}$ values found here ranged from -15‰ in ecosystems dominated by C_4 grasses to -29‰ in ecosystems dominated by C_3 plants (either herbaceous or woody plants). This supports our estimates of vegetation cover as being representative of the total in-situ biomass production, attributable to either C_3 or C_4 metabolic pathways. Both litter and superficial SOM $\delta^{13}\text{C}$ significantly reflected the current vegetation cover and, thus, could be used as reference to trace past changes in vegetation recorded in the SOM at deeper soil layers.

Past vegetation change

Using the equation presented on Fig. 2 and isotopic measurements performed at different depths of soil profiles, we calculated changes in the relative C_3 and C_4 vegetation cover at each study site. Our results show a consistent signal of continuous vegetation change, with C_4 being replaced by C_3 plants. Measurements of ^{14}C activity show that this process began approximately 3,870 (± 210) years before present and, since then, the relative abundance of C_3 plants increased 32% in average (Fig. 3). All woodlands and shrublands, currently dominated by C_3 plants ($>$ than 70% of the total vegetation cover), had a greater C_4 cover in the past. In some cases C_4 grasses were the predominant vegetation, while C_3 plants represented less than 20% of the total vegetation cover (see sites W1, W9, W13; Fig. 3). The least noticeable shift observed in woodlands corresponds to

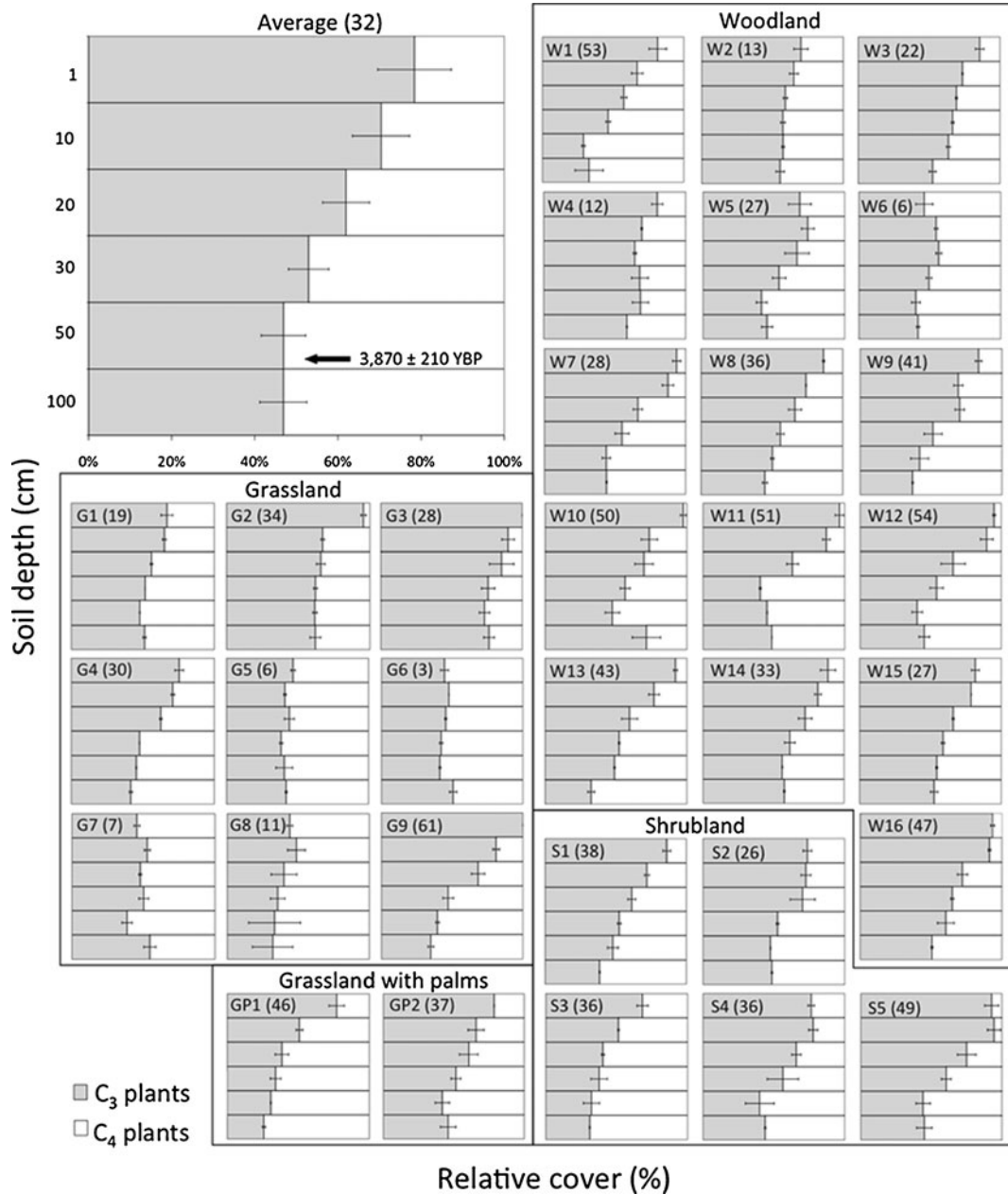


Fig. 3 Changes in the relative cover of C₃ and C₄ plants, estimated from δ¹³C values measured in the soil organic matter (SOM) at different depths of soil profiles. The conversion of δ¹³C values into percent cover followed the relationship found between present day vegetation cover and δ¹³C signatures measured in the litter and superficial SOM (Fig. 2). Grey and

white bars represent average values of C₃ and C₄ cover at each study site (3 to 5 profiles per site) and error bars correspond to one standard deviation. Numbers in brackets show total percent change in vegetation cover since 3,870 (±210) years before present. Site locations are shown on Fig. 1

a 6% change towards a greater C₃ cover in the present (W6; Fig. 3). In other woodland sites C₃ cover increased at least 12%, often reaching values greater than 40%. Similarly, the replacement of C₄ by C₃

plants in shrublands ranged from 26 to 49% in the past few thousand years (Fig. 3). In grasslands, vegetation cover remained nearly unaltered in a couple of sites (see G6, G7; Fig. 3), but in most

cases we observed shifts in C_3/C_4 cover of the same magnitude (in some cases higher) than those observed in shrub and woodland sites (see for example G9=61%; Fig. 3).

It is important to note that these results only represent records of past vegetation at each given site. However, the fact that noticeable changes in vegetation occurred in most sites and always in the same direction (C_3 replacing C_4 plants) suggests regional scale shifts in vegetation. Two-way ANOVA models indicate that changes in vegetation, measured as percent differences between past and recent C_3/C_4 cover, occurred independently of ecosystem (grassland, shrubland and woodland), altitude (sites ranged from 100 to 2,012 m asl) or their interaction. Likewise, the degree of vegetation change across sites, measured as the angular coefficient of least square regressions between C_3/C_4 cover and soil depths, were not affected by ecosystem, altitude or their interaction (Appendix 2).

Discussion

$\delta^{13}C$ signatures and past vegetation change

We found a highly significant relationship between the relative cover of C_3 (herbaceous and woody) and C_4 (grasses) with $\delta^{13}C$ values measured in the organic matter across sites. It has been well documented that differences in the total contribution of C_3 - and C_4 -originated biomass to whole ecosystems production yield distinguishable carbon isotopic signatures (Dümig et al. 2008; Ehleringer et al. 2000; Lloyd et al. 2008; Sanaiotti et al. 2002; Silva et al. 2010a, b, 2008; Victoria et al. 1995; Von Fischer et al. 2008). Similarly, here we show that differences in vegetation cover, with respect to metabolic pathways, explain variations in soil organic matter $\delta^{13}C$ signatures. Litter carbon isotope ratios varied with C_3/C_4 cover, remaining nearly unaltered after decomposition. The spectrum of $\delta^{13}C$ ratios reported here, varying up to 15‰ depending on the vegetation cover, is consistent with those described in different C_3 - and C_4 -dominated ecosystems (Lloyd et al. 2008; Silva et al. 2008; Von Fischer et al. 2008) and represent reliable signatures that can be used to trace shifts in vegetation through time.

Although processes other than changes in vegetation may alter isotopic ratios before and after litter

deposition, major paleoecological events can be identified with good confidence in soil profiles, because organic matter $\delta^{13}C$ ratios tend to be stable long after decomposition, conserving vegetation signatures throughout millennia (Silva et al. 2008; Von Fischer et al. 2008). Changes in the isotopic composition of atmospheric CO_2 could have affected values of $\delta^{13}C$ prior to deposition, but with the exception of past century, $\delta^{13}C$ variations in atmospheric CO_2 have been small (<1.0‰ within the last 10,000 years; Flückiger et al. 2002). Variations in $\delta^{13}C$ due to changes in photosynthetic discrimination are also possible, but would not have produced similar changes across sites/ecosystems and even if so, such variations would have been more than one order of magnitude smaller than the range observed here (Martin-Benito et al. 2010; Nock et al. 2010; Silva et al. 2010c, 2009).

Differences in root depth between trees and grasses could have contributed to soil organic matter isotopic enrichment with depth. However, the presumably deeper rooting habit of trees could have depleted (C_3 signal), but not enriched (C_4 signal), SOM $\delta^{13}C$. Moreover, differences in root depth between C_3 and C_4 plants typically yield small changes in $\delta^{13}C$ and isotopic offsets that exceed 3‰ are usually interpreted as a shift in vegetation (Krull et al. 2002; Von Fischer et al. 2008). Additionally, because changes in vegetation were observed in different ecosystems (grassland, shrubland and woodland), differences in root depth cannot explain changes in $\delta^{13}C$ observed here. Post depositional differential degradation of chemical compounds and fractionation associated with microbial activity could also have affected soil organic matter isotopic ratios (Marshall et al. 2007). However, this influence is typically observed only within the top 20 cm of the soil profile, correlated with total soil organic carbon content (Ehleringer et al. 2000; Krull et al. 2002). We observed changes in $\delta^{13}C$ values at much deeper layers of the soil profile that were not correlated with soil carbon content. Total soil carbon in the study region range from 2 to 26 $g\ kg^{-1}$ according to vegetation type and altitudinal/climatic variations (Alvarez and Lavado 1998), yet we observed consistent variations in $\delta^{13}C$ across sites, which did not change significantly in different ecosystems or altitudes. For these reasons, we conclude that changes in $\delta^{13}C$ values observed with soil depth in the present study were caused by past changes in vegetation.

A possible northward shift in the C_4 species range

We detected a convergent signal of soil organic matter isotopic signatures, showing a continuous increase in the relative abundance of C_3 plants since 3,870 (± 210) years ago. Previous palynological and isotopic studies from southern South America have identified the expansion of C_3 -dominated (forests) over C_4 -dominated ecosystems (savannas and grasslands) during the same period (Behling et al. 2005, 2004; Dümig et al. 2008; Mancini 2009; Silva et al. 2008; Wille and Schabitz 2009). While these studies have described local shifts in forest-savanna or forest-grassland ecotones, the results presented here show that vegetation changed at a regional scale and independently of ecosystem structure. In the past, C_4 grasses greatly contributed to the total biomass of sites now dominated by distinct groups of C_3 plants. Because changes in the predominant metabolic pathway occurred in woodlands, shrublands and grasslands, they were probably not related to intrinsic differences in plant life forms but rather had a physiological basis.

Changes in ecosystem structure, from dense to open vegetation, can be site-specific and determined by local environmental gradient or disturbance history (Crisci et al. 1991; Da Silva and Bates 2002). For example, it has been shown that spatial and/or temporal changes in vegetation structure may be attributed to variations in edaphic properties (Haridasan 2008), water availability (Bush et al. 2004; Furley et al. 1992; Silva et al. 2009), fire regime (Accatino et al. 2010; Hoffmann et al. 2003; Moreira 2000) and grazing intensity (Diaz et al. 2001). We suspect that all these factors have contributed to configure the current structure of the vegetation at each studied site. However, a simultaneous shift in the predominant metabolic pathway across sites, suggests that a major force has promoted large scale changes in plant communities during the late Holocene.

We believe that changes in climate were responsible for the shift in vegetation reported here. Worldwide, the extant C_4 genera occupy a wider range of drier and warmer habitats than their C_3 counterparts because the C_4 pathway represents a pre-adaptation to hot and arid conditions (Osborne and Freckleton 2009). Reconstructions of paleoclimate have shown a significant drop in global temperatures during the past few thousand years (Fig. 4). This climate cooling followed the retreat of glaciers in the northern

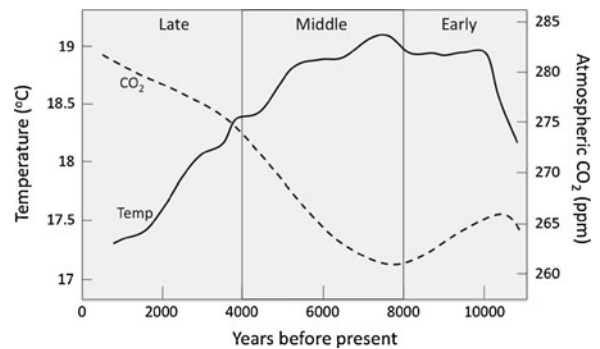


Fig. 4 Temperature and CO_2 records for the Holocene time period. Temperature record according to alkenone measurements (core GeoB 7139–2) of north-central Chile (De Pol-Holz et al. 2006) and CO_2 record described from ice core analyses (EPICA ice cores; Flückiger et al. 2002)

hemisphere during the Holocene climatic optimum, which also led to substantial water input in tropical and subtropical systems (Markgraf et al. 1992; Vimeux et al. 2009; Wanner et al. 2008). Fossil pollen, charcoal and isotopic evidence from different South American biomes indicate that C_3 plants became more abundant during the late Holocene, coinciding with changes in climate towards colder and wetter conditions (Behling et al. 2005, 2004, Mancini 2009; Silva et al. 2008).

Presently, the composition of C_3 and C_4 species in the study region follow temperature parameters more strongly than rainfall (Cavagnaro 1988; Cabido et al. 1997). Along altitudinal gradients C_4 appear in greater number than C_3 species in locations where summers are warmer than $18^\circ C$, while in colder areas

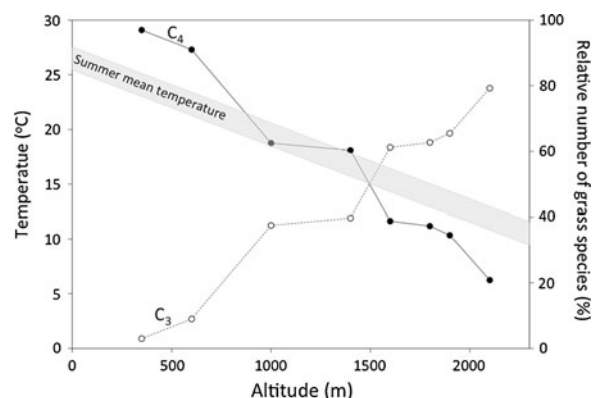


Fig. 5 Changes in the present day relative species composition of C_3 and C_4 grasses across an altitudinal gradient in central Argentina, as defined by summer (January) average temperatures (adapted from Cabido et al. 1997)

the number of C_3 species represent more than 60% of the total grass diversity (Fig. 5). Although the number of species is not necessarily linked to the total cover of either C_3 or C_4 plants, it seems fair to assume that colder temperatures (average annual temperatures dropped from $>18.5^\circ\text{C}$ to $<17.5^\circ\text{C}$; Fig. 4) increased the competitive advantage and, consequently, the relative abundance of C_3 plants at the latitude studied here. Increases in atmospheric CO_2 concentrations could also have favored C_3 at the expenses of C_4 plants (Epstein et al. 1997; Luo et al. 2006; Sage et al. 2010). Atmospheric CO_2 levels have increased monotonically (~ 20 ppm in total) since the mid Holocene (Fig. 4). The reasons are debatable, but decreases in total terrestrial biomass in response to colder temperatures (Indermühle et al. 1999) and a drop in oceanic carbonate ion concentration (Broecker et al. 2001) are possible causes behind this natural CO_2 enrichment. However, even though C_3 plants are more responsive to elevation in CO_2 than C_4 plants, enrichments of this magnitude should not have altered the competitive balance between these two metabolic pathways (Wand et al. 1999). Therefore, changes in temperature are the most likely explanation for a regional scale vegetation displacement.

Despite increasingly higher atmospheric CO_2 levels, warmer global temperatures are expected to promote the expansion of C_4 species range in the near future (Sage 2004; Sage et al. 2010; Wand et al. 1999). This

could represent a return to C_4 -dominated systems in the latitude studied here, as observed during warmer periods in the past. It is important to note however that the velocity of climate change during the Holocene allowed time for species to migrate in locked step with their optimal range and the consequences of the much faster rates of recent global warming on plant communities remain unknown. There is still great uncertainty in predicting the future of C_3 - and C_4 -dominated ecosystems globally (Bond 2008). So far attempts to predict climate change impacts on terrestrial ecosystems have been limited to correlations between the current climate and species distribution (Pearson and Dawson 2003). The search for long-term convergent patterns as those described here, could lead to an improved picture of climate-vegetation interactions, increasing our ability to predict the future impacts of climate change worldwide.

Acknowledgments This research was supported by the Natural Sciences and Engineering Council of Canada, Canada Research Chairs, Canadian Foundation for Innovation grants to M. A., SECyT, Universidad Nacional de Córdoba and Inter-American Institute for Global Change Research (IAI CRN II No. 2005) grants to M.A. and M.C. We are thankful to Diego Gurvich from the Instituto Multidisciplinario de Biología Vegetal (FCEfyN, UNC-CONICET), Argentina, Leandro Duarte and Mark Leithhead, from the Universidade Federal do Rio Grande do Sul, Brazil, for valuable comments and to the organizers of the IAI (2009) collaborative meeting in Córdoba, Argentina, where this collaboration began.

Appendix 1

Table 1 Identity, life form and metabolic pathway of all plant species present in the studied sites. Plants with C_4 photosynthetic pathway were distinguished from those that have the C_3

pathway by examination of the Kranz anatomy in cross-sections of fresh and herbarium specimens and from the literature

Family/species	Life form	Metabolic pathway
Acanthaceae		
<i>Dicliptera squarrosa</i> Nees	herb	C_3
<i>Justicia squarrosa</i> Griseb.	herb	C_3
<i>Stenandrium dulce</i> (Cav.) Nees	herb	C_3
Amaranthaceae		
<i>Amaranthus hybridus</i> L. ssp. <i>hybridus</i>	herb	C_3
<i>Guilleminea densa</i> (Willd. ex Roem. & Schult.) Moq.	herb	C_3
<i>Iresine diffusa</i> Humb. & Bonpl. ex Willd. var. <i>diffusa</i>	herb	C_3
<i>Pfaffia gnaphaliodes</i> (L. f.) Mart.	herb	C_3
Amaryllidaceae		

Table 1 (continued)

Family/species	Life form	Metabolic pathway
<i>Zephyranthes longistyla</i> Pax	herb	C3
Anacardiaceae		
<i>Lithraea molleoides</i> (Vell.) Engl.	tree	C3
<i>Schinus bumeloides</i> I.M. Johnst.	shrub	C3
<i>Schinus fasciculatus</i> (Griseb.) I. M. Johnst. var. <i>fasciculatus</i>	shrub	C3
Apiaceae		
<i>Ammi visnaga</i> (L.) Lam.	herb	C3
<i>Bowlesia incana</i> Ruiz & Pav.	herb	C3
<i>Conium maculatum</i> L.	herb	C3
<i>Eryngium agavifolium</i> Griseb.	herb	C3
<i>Eryngium horridum</i> Malme	herb	C3
<i>Eryngium nudicaule</i> Lam.	herb	C3
Apocynaceae		
<i>Amblyopetalum coccineum</i> (Griseb.) Malme	herb	C3
<i>Aspidosperma quebracho-blanco</i> Schtdl.	tree	C3
<i>Ditassa buchellii</i> Hook. & Arn.	epiphyte	C3
<i>Metastelma tubatum</i> Griseb.	epiphyte	C3
<i>Morreria odorata</i> (Hook. & Arn.) Lindl.	epiphyte	C3
Arecaceae		
<i>Trithinax campestris</i> (Burmeist.) Drude & Griseb.	palm	C3
Asteraceae		
<i>Ambrosia tenuifolia</i> Spreng.	herb	C3
<i>Baccharis coridifolia</i> DC.	herb	C3
<i>Baccharis pingraea</i> DC.	herb	C3
<i>Baccharis salicifolia</i> (Ruiz & Pav.) Pers.	herb	C3
<i>Baccharis stenophylla</i> Ariza	herb	C3
<i>Baccharis ulicina</i> Hook. & Arn.	herb	C3
<i>Bidens andicola</i> var. <i>decomposita</i> Kuntze	herb	C3
<i>Bidens pilosa</i> L. var. <i>pilosa</i>	herb	C3
<i>Bidens subalternans</i> DC.	herb	C3
<i>Carduus acanthoides</i> L.	herb	C3
<i>Carduus thoermeri</i> Weinm.	herb	C3
<i>Chaptalia nutans</i> (L.) Pol.	herb	C3
<i>Cirsium vulgare</i> (Savi) Ten.	herb	C3
<i>Conyza bonaeriensis</i> (L.) Cronquist	herb	C3
<i>Conyza primulifolia</i> (Lam.) Cuatrec. & Lourteig	herb	C3
<i>Eupatorium inulifolium</i> Kunth	herb	C3
<i>Eupatorium patens</i> D. Don ex Hook. & Arn.	herb	C3
<i>Eupatorium viscidum</i> Hook. & Arn.	herb	C3
<i>Galinsoga parviflora</i> Cav.	herb	C3
<i>Gamochaeta falcata</i> (Lam.) Cabrera	herb	C3
<i>Gamochaeta</i> sp.	herb	C3
<i>Gnaphalium gaudichardianum</i> DC.	herb	C3
<i>Heterosperma ovatifolium</i> Cav.	herb	C3
<i>Heterotheca subaxillaris</i> (Lam.) Britton & Rusby	herb	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
<i>Hypochaeris caespitosa</i> Cabrera	herb	C3
<i>Lactuca serriola</i> L.	herb	C3
<i>Schkuria pinnata</i> (Lam.) Kuntze ex Thell.	herb	C3
<i>Senecio ceratophylloides</i> Griseb.	herb	C3
<i>Senecio pampeanus</i> Cabrera	herb	C3
<i>Sonchus oleraceus</i> L.	herb	C3
<i>Synedrellopsis grisebachii</i> Hieron. & Kuntze	herb	C3
<i>Tagetes minuta</i> L.	herb	C3
<i>Taraxacum officinale</i> G. Weber ex F. H. Wigg.	herb	C3
<i>Verbesina encelioides</i> (Cav.) Benth. & Hook. f. ex A. Gray	herb	C3
<i>Vernonia incana</i> Less.	herb	C3
<i>Vernonia nudiflora</i> Less.	herb	C3
<i>Xanthium spinosum</i> L. var. <i>spinosum</i>	herb	C3
Basellaceae		
<i>Anredera cordifolia</i> (Ten.) Steenis	epiphyte	C3
Bignoniaceae		
<i>Amphilophium cynanchoides</i> (DC.) L. G. Lohmann	epiphyte	C3
<i>Dolichandra cynanchoides</i> Cham.	epiphyte	C3
Brassicaceae		
<i>Descurainia argentina</i> O.E. Schulz	herb	C3
<i>Exhalimolobos weddellii</i> (E. Fourn.) Al-Shehbaz & C. D. Bailey	herb	C3
<i>Lepidium bonariense</i> L.	shrub	C3
Bromeliaceae		
<i>Tillandsia aizoides</i> Mez	epiphyte	C3
<i>Tillandsia bryoides</i> Griseb. ex Baker	epiphyte	C3
<i>Tillandsia capillaris</i> Ruiz & Pav.	epiphyte	C3
<i>Tillandsia duratii</i> Vis. var. <i>duratii</i>	epiphyte	C3
<i>Tillandsia xiphioides</i> Ker Gawl. var. <i>xiphioides</i>	epiphyte	C3
Calyceraceae		
<i>Boopis anthemoides</i> Juss.	herb	C3
Campanulaceae		
<i>Wahlenbergia linarioides</i> (Lam.) A. DC.	herb	C3
Capparaceae		
<i>Capparis atamisquea</i> Kuntze	shrub	C3
Caryophyllaceae		
<i>Cardionema ramosissima</i> (Weinm.) A. Nelson & J. F. Macbr.	epiphyte	C3
<i>Polycarpon tetraphyllum</i> (L.) L.	herb	C3
<i>Silene antirrhina</i> L. var. <i>antirrhina</i>	herb	C3
Celastraceae		
<i>Moya spinosa</i> Griseb.	shrub	C3
Celtidaceae		
<i>Celtis ehrenbergiana</i> (Klotzch) Liebm.	tree	C3
Chenopodiaceae		
<i>Allenrolfea patagonica</i> (Moq.) Kuntze	herb	C3
<i>Atriplex lampa</i> (Moq.) D. Dietr.	herb	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
Chenopodiaceae sp.	herb	C3
Chenopodium album L.	herb	C3
Salsola kali L.	herb	C3
Sarcocornia ambigua (Michx.) M.A. Alonso & M.B. Crespo	herb	C3
Commelinaceae		
Commelina erecta L.	herb	C3
Convolvulaceae		
Cressa truxillensis Kunth	herb	C3
Dichondra microcalyx (Hallier f.) Fabris	herb	C3
Evolvulus sericeus Sw.	herb	C3
Ipomoea hieronymi (Kuntze) O'Donell	epiphyte	C3
Ipomoea nil (L.) Roth	epiphyte	C3
Ipomoea sp.	epiphyte	C3
Cucurbitaceae		
Sicyos malvifolius Griseb.	epiphyte	C3
Cyperaceae		
Bulbostylis juncooides (Vahl) Kük. ex Herter var. juncooides	sedge	C3
Carex sororia Kunth ssp. sororia	sedge	C3
Cyperaceae sp.	sedge	C3
Cyperus aggregatus (Willd.) Endl. var. aggregatus	sedge	C3
Cyperus entrerianus Boeck.	sedge	C3
Cyperus hermaphroditus (Jacq.) Standl.	sedge	C3
Cyperus incomtus Kunth	sedge	C3
Ephedraceae		
Ephedra americana Humb. & Bonpl. ex Willd.	epiphyte	C3
Ephedra triandra Tul. emend. J. H. Hunz.	epiphyte	C3
Euphorbiaceae		
Acalypha communis Müll. Arg.	herb	C3
Croton lachnostachyus Baill.	herb	C3
Euphorbia acerensis Boiss.	herb	C3
Euphorbia dentata Michx.	herb	C3
Euphorbia lorentzii Müll. Arg.	herb	C3
Euphorbia serpens Kunth	herb	C3
Tragia geraniifolia Klotzch ex Baill.	herb	C3
Tragia hieronymii Pax & K. Hoffm.	herb	C3
Fabaceae		
Acacia aroma Gillies ex Hook & Arn.	shrub	C3
Acacia caven (Molina) Molina	shrub	C3
Acacia gilliesii Steud.	shrub	C3
Acacia praecox Griseb.	shrub	C3
Adesmia bicolor (Poir.) DC.	epiphyte	C3
Cercidium praecox (Ruiz & Pav. ex Hook.) Harms glaucum Ssp. (Cav.) Burkart & Carter	tree	C3
Cologamia broussonetii (Balb.) DC.	epiphyte	C3
Coursetia hassleri Chodat	herb	C3
Desmanthus sp.	herb	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
<i>Desmodium uncinatum</i> (Jacq.) DC.	herb	C3
Fabaceae sp.	herb	C3
<i>Geoffraea decorticans</i> (Gillies ex Hook. & Arn.) Burkart	tree	C3
<i>Gleditsia triacanthos</i> L.	tree	C3
<i>Medicago lupulina</i> L.	herb	C3
<i>Mimosa detinens</i> Benth.	shrub	C3
<i>Mimozyanthus carinatus</i> (Griseb.) Burkart	shrub	C3
<i>Prosopis alba</i> Griseb.	tree	C3
<i>Prosopis caldenia</i> Burkart	tree	C3
<i>Prosopis flexuosa</i> DC.	tree	C3
<i>Prosopis nigra</i> (Griseb.) Hieron. var. <i>nigra</i>	tree	C3
<i>Prosopis pugionata</i> Burkart	tree	C3
<i>Prosopis torquata</i> (Cav. ex Lag.) DC.	tree	C3
<i>Rhynchosia senna</i> Griseb. ex Hook.	epiphyte	C3
<i>Senna aphylla</i> (Cav.) H. S. Irwin & Barneby	herb	C3
Geraniaceae		
<i>Geranium dissectum</i> L.	herb	C3
Juncaceae		
<i>Juncus acutus</i> L.	sedge	C3
<i>Juncus imbricatus</i> Laharpe	sedge	C3
<i>Juncus</i> sp	sedge	C3
Lamiaceae		
<i>Clinopodium gilliesii</i> (Benth.) Kuntze	shrub	C3
<i>Hyptis floribunda</i> (Briq.) Briq. ex Micheli	herb	C3
<i>Hyptis mutabilis</i> (Rich.) Briq.	herb	C3
<i>Leonurus japonicus</i> Houtt.	herb	C3
Lythraceae		
<i>Cuphea glutinosa</i> Cham. & Schldl.	herb	C3
<i>Heimia salicifolia</i> (Kunth) Link	herb	C3
Malpighiaceae		
<i>Cordobia argentea</i> (Griseb.) Nied.	epiphyte	C3
<i>Janusia guaranítica</i> (A. St.-Hil.) A. Juss.	epiphyte	C3
<i>Tricomaria usillo</i> Hook. & Arn.	shrub	C3
Malvaceae		
<i>Krapovicasea flavescens</i>	herb	C3
<i>Malvastrum coromandelianum</i> (L.) Garcke	herb	C3
<i>Modiolastrum malvifolium</i> (Griseb.) K. Schum.	herb	C3
<i>Pseudoabutilon pedunculatum</i> (R. E. Fr.) Krapov.	herb	C3
<i>Sida dictyocarpa</i> Griseb. ex K. Schum.	herb	C3
<i>Sida rhombifolia</i> L.	herb	C3
<i>Sida spinosa</i> L.	herb	C3
Moraceae		
<i>Morus alba</i> L.	tree	C3
Olacaceae		
<i>Ximenia americana</i> L.	shrub	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
Onagraceae		
<i>Oenothera affinis</i> Cambess.	herb	C3
<i>Oenothera</i> sp.	herb	C3
Orchidaceae		
<i>Cyclopogon elatus</i> (Sw.) Schltr.	herb	C3
<i>Habenaria</i> sp.	herb	C3
Oxalidaceae		
<i>Oxalis conorrhiza</i> Jacq.	herb	C3
Passifloraceae		
<i>Passiflora mooreana</i> Hook. f.	epiphyte	C3
Phytolaccaceae		
<i>Petiveria alliacea</i> L.	herb	C3
<i>Rivinia humilis</i> L.	herb	C3
Plantaginaceae		
<i>Plantago tomentosa</i> Lam.	herb	C3
Poaceae		
<i>Agrostis montevidensis</i> Spreng. ex Nees f. <i>montevidensis</i>	grass	C3
<i>Aristida laevis</i> (Nees) Kunth	grass	C4
<i>Aristida mendocina</i> Phil.	grass	C4
<i>Bothriochloa barbinodis</i> (Lag.) Herter	grass	C4
<i>Bothriochloa laguroides</i> (DC.) Herter ssp. <i>laguroides</i>	grass	C4
<i>Bothriochloa springfieldii</i> (Gould) Parodi	grass	C4
<i>Bromus catharticus</i> Vahl var. <i>catharticus</i>	grass	C3
<i>Cenchrus myosuroides</i> Kunth var. <i>myosuroides</i>	grass	C4
<i>Cenchrus spinifex</i> Cav.	grass	C4
<i>Chascolytrum subaristatum</i> (Lam.) Desv.	grass	C3
<i>Chloris halophila</i> Parodi var. <i>halophila</i>	grass	C4
<i>Chloris</i> sp.	grass	C4
<i>Cortaderia selloana</i> (Schult. & Schult. f.) Asch. & Graebn.	grass	C3
<i>Cynodon dactylon</i> (L.) Pers. var. <i>dactylon</i>	grass	C4
<i>Deyeuxia hieronymi</i> (Hack.) Türpe	grass	C3
<i>Digitaria californica</i> (Benth.) Henrard var. <i>californica</i>	grass	C4
<i>Digitaria sacchariflora</i> (Nees) Henrard	grass	C4
<i>Digitaria sanguinalis</i> (L.) Scop.	grass	C4
<i>Distichlis scoparia</i> (Kunth) Arechav. var. <i>scoparia</i>	grass	C4
<i>Distichlis spicata</i> (L.) Greene var. <i>spicata</i>	grass	C4
<i>Eleusine tristachya</i> (Lam.) Lam.	grass	C4
<i>Eragrostis curvula</i> (Schrud.) Nees	grass	C4
<i>Eragrostis lugens</i> Nees	grass	C4
<i>Eragrostis mexicana</i> (Hornem.) Link ssp. <i>mexicana</i>	grass	C4
<i>Eustachys retusa</i> (Lag.) Kunth	grass	C4
<i>Festuca hieronymi</i> Hack. var. <i>hieronymi</i>	grass	C3
<i>Gouinia paraguayensis</i> (Kuntze) Parodi	grass	C4
<i>Hordeum stenostachys</i> Godr.	grass	C3
<i>Jarava plumosa</i> (Spreng.) S. W. L. Jacobs & J. Everett	grass	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
<i>Jarava pseudoichu</i> (Caro) F. Rojas	grass	C3
<i>Leptochloa chloridiformis</i> (Hack.) Parodi	grass	C4
<i>Lolium perenne</i> L.	grass	C3
<i>Muhlenbergia peruviana</i> (P. Beauv.) Steud.	grass	C4
<i>Nassella hyalina</i> (Nees) Barkworth	grass	C3
<i>Nassella tenuissima</i> (Trin.) Barkworth	grass	C3
<i>Nassella trichotoma</i> (Nees) Hack. ex Arechav.	grass	C3
<i>Neobouteloua lophostachya</i> (Griseb.) Gould	grass	C4
<i>Pappophorum pappiferum</i> (Lam.) Kuntze	grass	C4
<i>Paspalum dilatatum</i> (Poir.) ssp. <i>dilatatum</i>	grass	C4
<i>Paspalum malacophyllum</i> Trin.	grass	C4
<i>Paspalum notatum</i> Flügge	grass	C4
<i>Paspalum quadrifarium</i> Lam.	grass	C4
<i>Piptochaetium</i> sp.	grass	C3
<i>Poa</i> sp	grass	C3
<i>Setaria lachnea</i> (Nees) Kunth	grass	C4
<i>Setaria oblongata</i> (Griseb.) Parodi	grass	C4
<i>Setaria pampeana</i> Parodi ex Nicora	grass	C4
<i>Setaria parviflora</i> (Poir.) Kerguelen	grass	C4
<i>Sorghum halepense</i> (L.) Pers.	grass	C3
<i>Spartina spartinae</i> (Trin.) Merr. ex Hitchc.	grass	C3
<i>Sporobolus indicus</i> (L.) R. Br.	grass	C4
<i>Sporobolus pyramidatus</i> (Lam.) Hitchc.	grass	C4
<i>Tragus berteronianus</i> Schult.	grass	C4
<i>Trichloris crinita</i> (Lag.) Parodi	grass	C4
Polygonaceae		
<i>Rumex acetosa</i> L.	herb	C3
Portulacaceae		
<i>Portulaca oleraceae</i> L.	herb	C3
<i>Talinum fruticosum</i> (L.) Juss.	herb	C3
<i>Talinum paniculatum</i> (Jacq.) Gaertn.	herb	C3
<i>Talinum polygaloides</i> Gillies ex Arn.	herb	C3
Ranunculaceae		
<i>Clematis montevidense</i> Spreng.	epiphyte	C3
Rhamnaceae		
<i>Condalia buxifolia</i> Reissek	shrub	C3
<i>Condalia microphylla</i> Cav.	shrub	C3
<i>Condalia montana</i> A. Cast.	shrub	C3
<i>Ziziphus mistol</i> Griseb.	tree	C3
Rosaceae		
<i>Lachemilla pignata</i> (Ruiz & Pav.) Rothm.	herb	C3
Rubiaceae		
<i>Borreria eryngioides</i> var. <i>ostenii</i> (Standl.) E.L. Cabral & Bacigalupo	herb	C3
<i>Galium richardianum</i> (Gillies ex Hook. & Arn.) Endl. ex Walp.	herb	C3
<i>Mitracarpus megapotamicus</i> (Spreng.) Kuntze	herb	C3

Table 1 (continued)

Family/species	Life form	Metabolic pathway
Rutaceae		
<i>Zanthoxylum coco</i> Gillies ex Hook. f. et Arn.	tree	C3
Santalaceae		
<i>Jodina rhombifolia</i> (Hook. & Arn.) Reissek	shrub	C3
Sapindaceae		
<i>Cardiospermum halicacabum</i> L.	epiphyte	C3
<i>Urvillea chacoënsis</i> Hunz.	epiphyte	C3
Selaginellaceae		
<i>Selaginella peruviana</i> (Milde) Hieron.	fern	C3
<i>Selaginella sellowii</i> Hieron.	fern	C3
Simaroubaceae		
<i>Castela coccinea</i> Griseb.	tree	C3
Solanaceae		
<i>Capsicum chacoënsis</i> Hunz.	herb	C3
<i>Cestrum parqui</i> L'Hér.	shrub	C3
<i>Datura ferox</i> L.	herb	C3
<i>Grabowskia</i> aff. <i>duplicata</i> Arn.	shrub	C3
<i>Lycium chilense</i> Miers ex Bertero	shrub	C3
<i>Lycium ciliatum</i> Schltdl.	shrub	C3
<i>Lycium elongatum</i> Miers.	shrub	C3
<i>Lycium gilliesianum</i> Miers	shrub	C3
<i>Nierembergia aristata</i> D. Don	herb	C3
<i>Petunia axillaris</i> (Lam.) Britton, Stern & Poggenb.	herb	C3
<i>Salpichroa origanifolia</i> (Lam.) Baill.	herb	C3
<i>Solanum angustifidum</i> Bitter	herb	C3
<i>Solanum argentinum</i> Bitter & Lillo	herb	C3
<i>Solanum chenopodioides</i> Lam.	herb	C3
<i>Solanum elaeagnifolium</i> Cav.	herb	C3
<i>Solanum sisymbriifolium</i> Lam.	herb	C3
<i>Solanum stuckertii</i> Bitter	herb	C3
<i>Vassobia breviflora</i> (Sendtn.) Hunz.	shrub	C3
Urticaceae		
<i>Parietaria debilis</i> G. Forst.	herb	C3
<i>Urtica</i> sp.	herb	C3
Verbenaceae		
<i>Aloysia gratissima</i> (Gillies & Hook. ex Hook.) Tronc.	shrub	C3
<i>Glandularia peruviana</i> (L.) Small	herb	C3
<i>Lantana grisebachii</i> Seckt var. <i>grisebachii</i>	herb	C3
<i>Lippia turbinata</i> Griseb. f. <i>turbinata</i>	shrub	C3
<i>Phyla canescens</i> (Kunth) Greene	herb	C3
<i>Verbena bonariensis</i> L.	herb	C3
Zygophyllaceae		
<i>Larrea divaricata</i> Cav.	shrub	C3
<i>Porlieria mycophylla</i> (Baill.) Descole, O'Donell & Lourteig	shrub	C3

Appendix 2

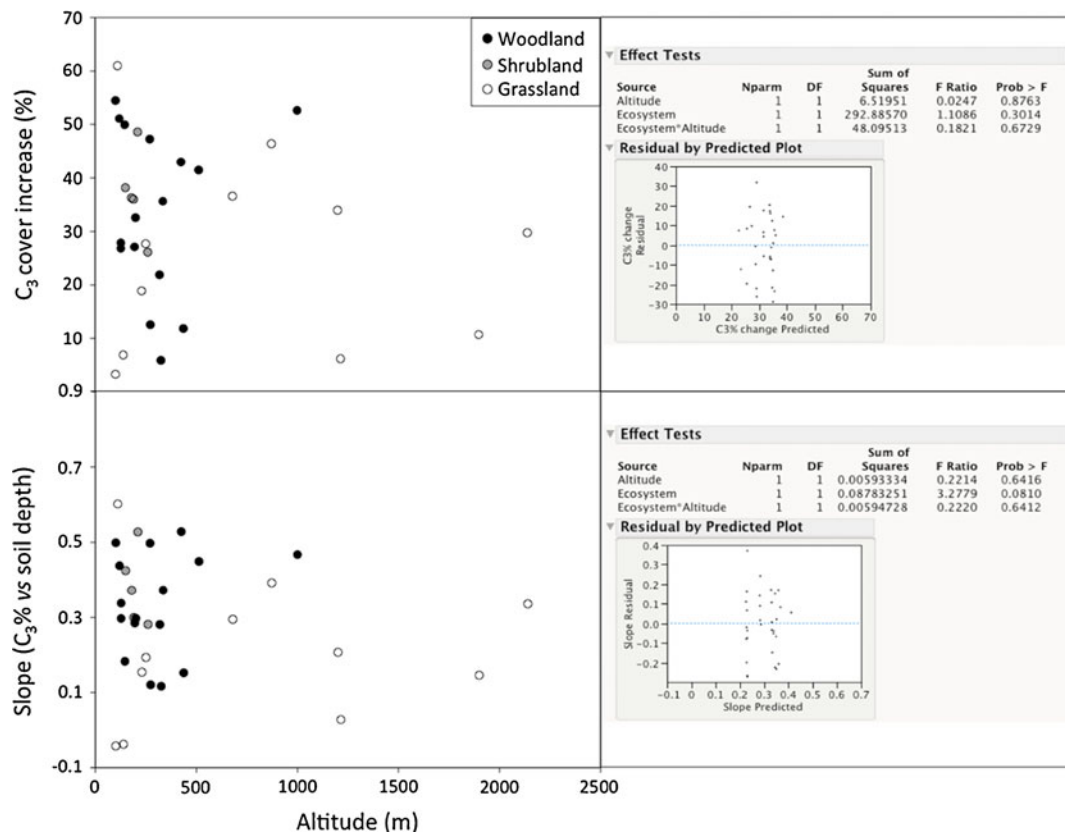


Fig. 6 Changes in vegetation cover through time across ecosystems and altitudes, as measured by percent differences between the C_3/C_4 relative cover estimated in superficial (0–1 cm) and in deep soils (50 cm) and by slope (angular

coefficient) of least square regressions, performed between estimates of C_3/C_4 cover and their respective soil depth at each study site. Note that no significant effects on vegetation change are observed for ecosystem, altitude or their interaction

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