1

An Assessment of Grazing Effects on Soil Carbon Stocks in Grasslands

Gervasio Piñeiro,¹ José M. Paruelo,^{1,2} Martín Oesterheld,^{1,2} and Esteban G. Jobbágy³

Authors are ¹Research Scientists and ²Associated Professors, IFEVA, Laboratorio de Análisis Regional y Teledetección, Facultad de Agronomía, Universidad de Buenos Aires/CONICET, Buenos Aires, Argentina; and ³Research Scientist and Adjunct Professor, Grupo de Estudios Ambientales, IMASL, Universidad Nacional de San Luis–CONICET, San Luis, Argentina.

Abstract

Grazing modifies the structure and function of ecosystems, affecting soil organic carbon (SOC) storage. Although grazing effects on some ecosystem attributes have been thoroughly reviewed, current literature on grazing effects on SOC needs to be synthesized. Our objective was to synthesize the effects of grazing on SOC stocks in grasslands, establishing the major mechanistic pathways involved. Additionally, and because of its importance for C biogeochemistry, we discuss the controls of soil organic nitrogen stocks. We reviewed articles analyzing grazing effects on soil organic matter (SOM) stocks by comparing grazed vs. ungrazed sites, including 67 paired comparisons. SOC increased, decreased, or remained unchanged under contrasting grazing conditions across temperature and precipitation gradients, which suggests that grazing influences the factors that control SOC accumulation in a complex way. However, our review also revealed some general patterns such as 1) root contents (a primary control of SOC formation) were higher in grazed than in their ungrazed counterparts at the driest and wettest sites, but were lower at sites with intermediate precipitation (~400 mm to 850 mm); 2) SOM C:N ratios frequently increased under grazing conditions, which suggests potential N limitations for SOM formation under grazing; and 3) bulk density either increased or did not change in grazed sites. Nearly all sites located in the intermediate precipitation range showed decreases or no changes in SOC. We grouped previously proposed mechanisms of grazing control over SOC into three major pathways that can operate simultaneously: 1) changes in net primary production (NPP pathway), 2) changes in nitrogen stocks (nitrogen pathway), and 3) changes in organic matter decomposition (decomposition pathway). The relative importance of the three pathways may generate variable responses of SOC to grazing. Our conceptual model suggests that rangeland productivity and soil carbon sequestration can be simultaneously increased by management practices aimed at increasing N retention at the landscape level.

Resumen

El pastoreo modifica la estructura y el funcionamiento de los ecosistemas, alterando las reservas de C orgánico del suelo (COS). A pesar de que existen revisiones de los efectos del pastoreo sobre la productividad primaria o los cambios en la composición de especies, no se dispone de una síntesis detallada de los efectos del pastoreo sobre el COS. A su vez, discutiremos los controles de las reservas de nitrógeno orgánico del suelo, debido a su importancia en la regulación del ciclo del C. El objetivo de este trabajo fue sintetizar los efectos del pastoreo sobre el COS y analizar los mecanismos a través de los cuales opera. Realizamos una revisión de los artículos que evalúan, mediante comparaciones de sitios pastoreados y no pastoreados, los efectos del pastoreo sobre el COS, incluyendo 67 comparaciones pareadas. La revisión bibliográfica mostró que: 1) el C en las raíces y otros órganos subterráneos fue mayor en zonas pastoreadas que en las no pastoreadas en sitios secos y húmedos, pero menor en sitos con precipitaciones intermedias; 2) el pastoreo aumentó la relación C:N del suelo, sugiriendo limitaciones de N para la formación de materia orgánica del suelo; y 3) la densidad aparente del suelo fue mayor o igual en los sitios pastoreados, respecto a los no pastoreados. Los efectos del pastoreo sobre el COS operarían a través de tres vías: 1) por medio de cambios en la productividad primaria neta, 2) a través de alteraciones en los stocks de NOS, o 3) aumentando la descomposición del COS. El pastoreo afectaría el COS a través de los tres mecanismos simultáneamente, y la respuesta del COS al pastoreo dependerá de la importancia relativa de cada uno. Finalmente, el modelo conceptual propuesto sugiere que la productividad de la biomasa y la cantidad de carbono almacenado en el suelo podrían aumentarse simultáneamente mediante manejos que apunten a incrementar la retención de N a escala de paisaje.

Key Words: belowground production, C:N, herbivores, nitrogen, reactive nitrogen, roots, soil organic carbon

INTRODUCTION

Grazing modifies the structure and function of ecosystems, affecting their stability and productivity (McNaughton 1985; Sala 1988; Milchunas and Lauenroth 1993; Altesor et al. 1998; Schuman et al. 1999). As a consequence, understanding the effects of grazing at the ecosystem level has been a primary goal of range scientists (Milchunas et al. 1988). However, it has been difficult to synthesize and generalize the effects of grazing

RANGELAND ECOLOGY & MANAGEMENT 63(1) January 2010

Rangeland Ecology and Management rama-63-01-05.3d 24/11/09 14:31:11 1 Cust # 08-255R1

This work was partially funded by grants from the Inter-American Institute for Global Change Research (IAI, CRN II 2031), which is supported by the US National Science Foundation (Grant GEO-0452325), FONCYT- PICT 32399, PICT 32415, and PICT 1764, CONICET (PIP 5961), UBACYT (CS 6179/05 G 078 and G 071), INIA-FPTA 175 (Uruguay), and SENSOR-TTC 003874-2.

Correspondence: Gervasio Piñeiro, Facultad de Agronomía, Universidad de Buenos Aires, IFEVA/CONICET, San Martín 4453, Capital Federal, C1417DSE, Argentina. Email: piñeiro@ ifeva.edu.ar

Manuscript received 5 January 2009; manuscript accepted 28 July 2009.

on grasslands, particularly its effect on the carbon cycle (Milchunas and Lauenroth 1993). Although grazing effects on some ecosystem attributes such as Aboveground Net Primary Production (ANPP) or species composition have been successfully reviewed (Milchunas and Lauenroth 1993; Oesterheld et al. 1999; Cingolani et al. 2005), current literature on grazing effects on soil organic carbon (SOC) needs to be synthesized.

Soil organic matter (SOM) is the main reservoir of SOC and soil organic nitrogen (SON) in rangelands and determines soil fertility, water retention, and soil structure (Tiessen et al. 1994; Lal 2004a). Grassland soils can store more than 100 and 10 tons per hectare of SOC and SON, respectively, in their first meter (Jobbágy and Jackson 2000), and grazing can increase, decrease, or maintain unaltered the size of both pools (Milchunas and Lauenroth 1993; Derner et al. 2006; Piñeiro et al. 2009). Therefore, management practices aimed to increase SOC and SON contents in rangelands are highly relevant as potential strategies for C sequestration (Conant and Paustian 2002; Schuman et al. 2002; Lal 2004a, 2004b).

SOC and SON are tightly associated as structural components of organic matter with a relatively constant mass ratio (C:N) in short- to mid-term time scales (Baisden and Amundson 2003). Although SOM C:N ratios may shift after grazing or other disturbances, SON dynamics frequently constrain C fluxes and SOC accumulation in soils (Wedin 1995; Semmartin and Oesterheld 2001; Baer et al. 2003; Baisden and Amundson 2003; Piñeiro et al. 2006, 2009; Harpole et al. 2007). While most C and N fluxes within ecosystems tend to be coupled, the inputs and outputs of these elements can be decoupled (i.e., photosynthesis and decomposition in the case of C vs. biological fixation and volatilization in the case of N). For this reason input or output processes that involve N exclusively can influence SOC dynamics through cascading effects across within-ecosystem fluxes (Wedin and Tilman 1996; Piñeiro et al. 2006, 2009). However, N is not the only constraint for SOC accumulation. Water availability and C uptake (net primary production) can also limit SOC accumulation, especially in arid sites (Burke et al. 1998).

Our objectives were to review and synthesize the effects of grazing on SOC stocks in grasslands and to identify different pathways by which grazing might alter this soil pool. Additionally, and because of its importance for C biogeochemistry, we discuss the controls of SON stocks in grasslands. We present a series of conceptual models aimed to organize the current knowledge on grazing impacts on SOC and SON pools. We also examine the potential effects of grazers on N emissions into the atmosphere, and finally, we discuss how the conceptual models may help to design management strategies oriented to increase productivity and reduce greenhouse gases emissions.

CONTROLS ON SOIL ORGANIC C AND N ACCUMULATION

In this section we present conceptual models showing the major controls of C and N accumulation in soils, following the conceptual model for decomposition proposed by Chapin et al. (2002, chapter 6; Fig. 1). Our scheme includes context factors and interactive, indirect, and direct controls as well as the disturbance regime. We constructed the diagrams based on reported controls of SOC and SON accumulation from the literature and arranged them following the rationale by Chapin et al. (2002). Based on the model, we explore the main processes through which grazing (but also other disturbances, such as fire or flooding) may affect SOC and SON accumulation at different time scales (Fig. 1).

In the short term, SOC and SON contents are a function of direct controls that represent the main inputs to and outputs from these pools (Fig. 1). Changes in SOC and SON stocks are determined by the balance between C and N inputs and outputs in a given period. SOC stocks are directly controlled by Net Primary Production (NPP; C inputs) and respiration and C lixiviation (C outputs), whereas SON stocks are controlled by N atmospheric depositions and biological N fixation (N inputs) and N volatilization and leaching (N outputs; Fig. 1). NPP, but essentially the proportion of NPP that is allocated to belowground organs, controls SOC storage (Jobbágy and Jackson 2000). NPP and SOC are, in turn, mutually dependent, as discussed below. C outputs (leaching and organic matter respiration) are controlled by soil texture, mainly by clay and silt particles that adsorb organic molecules and favor SOM stability (Hassink 1997). Because the direct controls of SOC and SON accumulation differ, a partial decoupling between SOC and SON can be expected in the short term, and thus C and N stocks may change in opposite directions during this time frame (Asner et al. 1997).

Intermediate and long-term changes in SOC and SON would result from cumulative inputs and outputs, but are constrained by factors that operate on these direct controls (indirect controls and interactive factors; Fig. 1). N availability is a critical indirect control of both SOC and SON accumulation (Fig. 1) because it constrains both inputs and outputs of C and N (i.e., N availability can increase NPP, increasing C inputs to the soil, and may also decrease soil respiration, decreasing C outputs from the soil). The same interactive controls that control SOC also affect SON accumulation, supporting the tight association between the two ecosystem pools. Community composition affects and is affected by SOC and SON contents at intermediate to long-term time scales (Jobbágy and Jackson 2000: Altesor et al. 2006). Soil resources, especially N but also P or other elements (e.g., Ca, Fe), may constrain organic matter accumulation in soils (Hedin et al. 2003). Thus, the depletion or accretion of C or N usually promotes losses or gains of the other element or changes in C:N ratios at intermediate to longterm time scales (Asner et al. 1997; Piñeiro et al. 2006). If interactive controls change in time or are suddenly modified by a disturbance (fire, grazing, etc.), they may modify SOC and SON via cascade effects, affecting indirect and direct controls.

Context factors constrain SOC and SON accumulation, establishing their potential ranges (Chapin et al. 2002; Fig. 1). Profuse evidence shows that climate (Jobbágy and Jackson 2000), biota (Jobbágy and Jackson 2000), time (Vitousek and Reiners 1975), topography, and parent material (specifically soil texture; Burke et al. 1989; Torn et al. 1997) control SOC and SON accumulation. Context factors do not affect input and outputs of C and N directly, but operate via cascade effects through other shorter-term controls, thus ultimately controlling SOC and SON stocks. Different contexts result in different responses to grazing. For example, Piñeiro et al. (2009) found that grazing increases SOC in deep soils but reduces it in

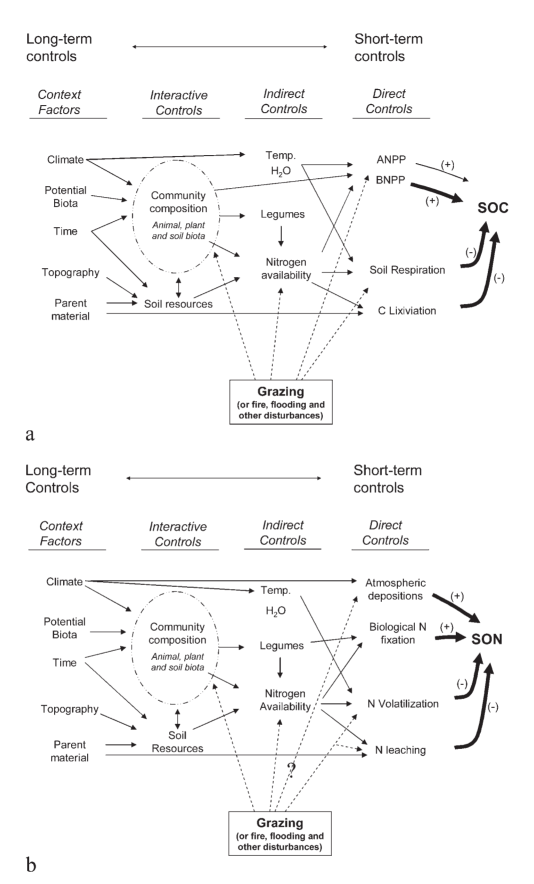


Figure 1. Soil organic carbon (SOC; **a**) and soil organic nitrogen (SON; **b**) controls at different temporal scales. Dashed lines show which controls are affected by grazing. ANPP is aboveground net primary production, and BNPP is belowground net primary production. See text for details.

shallow soils. Recently Derner et al. (2006) showed that in grasslands of North America grazing increased SOC stocks in dry shortgrass steppe ecosystems and decreased them in more humid mid- and tallgrass prairie ecosystems. In our model, similar context factors, interactive controls, and indirect controls affect SOC and SON accumulation, while direct controls differ between SOC and SON (Fig. 1). Thus, SOC and SON responses to grazing may differ in the short term, but will probably converge in the long term.

In addition to context factors the disturbance regime may significantly affect SOC and SON accumulation. SOC and SON response to grazing can be substantially different when interacting with other disturbances such as fire (Briggs et al. 2005; Piñeiro et al. 2006) or flooding (Chaneton et al. 1988). Grazing may modify (increase or decrease) fire frequency and intensity (Oesterheld et al. 1999; Bond and Keeley 2005). Firegrazing interactions can change SOC accumulation, depending on the relative magnitude of nutrient losses promoted by each disturbance (Briggs et al. 2005; Piñeiro et al. 2006). Fire and grazing are not the only factors that may interact and alter SOC contents. In the Flooding Pampas of Argentina, Piñeiro et al. (2009) found that grazing increased SOC contents at lowland sites (frequently flooded) but decreased it in uplands sites (not flooded).

GRAZING EFFECTS ON SOIL ORGANIC C AND N

In this section we present results from a literature review of grazing effects on SOC stocks, performed by compiling papers that report comparisons of grazed vs. ungrazed paired sites. We searched the ISI web of knowledge to identify articles that measured SOC contents under grazed and ungrazed conditions using the keywords "soil and carbon and grazing." We included only papers performed in grasslands and did not consider some located in savannas (e.g., Savadogo et al. 2007). Unfortunately, several papers were excluded from the review because they did not report SOC stocks corrected by soil bulk density or did not provide data to perform the correction (e.g., Zeidler et al. 2002; Beukes and Cowling 2003; Feral et al. 2003). We did not find any papers from Africa or Australia, although we did find three papers reporting the effects of different grazing intensities or rotational vs. continuous grazing in Australia. We finally included 20 articles reporting 67 comparisons that evaluated grazing effects on SOC stocks (Table 1), and we additionally compiled from these papers SOM C:N ratios, roots biomass contents, soil bulk density changes, mean annual precipitation (MAP), and mean annual temperature (MAT) when reported.

SOC increased, decreased, or remained unchanged under contrasting grazing conditions across temperature and precipitation gradients, suggesting that grazing influences the factors that control SOC in a complex way (Table 1). However, our review also revealed some general patterns, such as 1) root contents (a primary control of SOC formation) were higher in grazed than in their ungrazed counterparts at the driest and wettest sites, but were lower at sites with intermediate precipitation (\sim 400 mm to 850 mm); 2) SOM C:N ratios consistently increased under grazing conditions; and 3) bulk density either increased or did not change in grazed sites.

Our review showed how grazing affected differentially belowground biomass across one context factor: MAP. Following Burke et al.'s (1998) conceptual model for plantsoil interactions, we suggest that grazing results in the stimulation of root production by changing both allocation (Gao et al. 2008) or species abundances (Derner et al. 2006) in dry sites for the capture of water and in humid sites for the capture of soil nutrients. The mechanism operating at intermediate precipitation sites remains unclear but could be probably related to light limitation of NPP. Higher root biomass have been suggested to increase SOC contents because of greater C inputs to the soil (Derner et al. 2006). However, SOC changes in grazed sites were not associated with variations in belowground biomass as expected initially (Table 1). Our review suggests that grazing effects on SOC will depend not only on belowground allocation but also on other site characteristics (probably soil properties) and on grazinginduced modifications of other C and N stocks and fluxes.

Grazing increased or maintained unchanged SOM C:N ratio at nearly all sites, despite increases or decreases in SOC contents (Table 1; grazing increased organic matter C:N ratios at 25 sites, maintained unchanged at 22 sites, and decreased it at two sites, whereas no data were available for 17 sites). This consistent trend in C:N ratios suggests a potential N limitation under grazing. Increases in SOM C:N ratios often occur in the more labile fractions, because recalcitrant fractions have low and constant C:N ratios near 10. Piñeiro et al. (2009) reported increases in the C:N ratios of the particulate (less decomposed) organic matter fraction after grazing removal. Conversely, organic matter fractions associated with mineral particles (a more recalcitrant pool) maintained similar and low C:N ratios at grazed and ungrazed sites, despite the higher C:N ratios of litter and root inputs in grazed sites. These results suggest that the higher C:N ratios of plant debris are only increasing the C:N ratio of the labile SOM fractions. Then, to accumulate C in the recalcitrant SOM fractions, more C will be respired to achieve the low C:N ratios of these fractions, or in other words, N would be limiting SOM accumulation in stable pools. Because grasslands accumulate the majority of SOC in recalcitrant fractions, grazing may reduce SOM formation via N limitation. However, N limitation is not the only factor constraining SOC accumulation under grazing, because either increases or decreases of SOC occurred when grazing significantly increased SOM C:N ratios (Table 1).

As expected, soil bulk density increased or remained invariant at grazed compared to ungrazed sites. Soil compaction under grazing is attributed to trampling by herbivores (Taboada and Lavado 1993). Soil compaction reduces water infiltration and increases runoff under grazing conditions, decreasing water available for plant growth (Abdel-Magid et al. 1987; Savadogo et al. 2007). Additionally, less pore space can limit gas exchange and reduce root growth. Both mechanisms suggest that soil compaction reduces plant production and thus SOC storage. However, soil compaction is not the only way by which herbivores alter SOC stocks, because both increases and decreases of SOC have been reported with increases in soil bulk density (Table 1). Thus, as for N limitation, other mechanisms

 Table 1. Grazing effects on different soil properties and belowground biomass in grasslands.

Author	Mean annual precipitation	Mean annual temperature	$\Delta \ {\rm SOC^1}$	Δ C:N SOM ¹	Δ Bulk density ¹	$\Delta \ \rm roots^1$	<i>n</i> (total 67)	Sampling depth	Experiment type, and age (yr)	Observations
Frank et al. (1995)	n/d	n/d	Decreases	Increases	n/d	=	1	0–100	Exclosures, 75	Moderate grazing
Frank et al. (1995)	n/d	n/d	=	=	n/d	Increases	1	0–100	Exclosures, 75	Heavy grazing
Neff et al. (2005)	207	11.6	Decreases	Increases, $=$	=	n/d	1	0–30	Never grazed	Wind erosion
Shrestha and Stahla (2008)	213	n/d	=	=	=	n/d	1	0–15	Exclosures, 43	High elevation
Shrestha and Stahla (2008)	200	n/d	=	=	=	n/d	1	0–15	Exclosures, 47	High elevation
Shrestha and Stahla (2008)	203	n/d	=	=	=	n/d	1	0–15	Exclosures, 41	High elevation
Shrestha and Stahla (2008)	203	n/d	=	Decreases	=	n/d	1	0–15	Exclosures, 41	High elevation
Smoliak et al. (1972)	310	n/d	Increases	Increases	=	Increases	1	0–10	Exclosures, 42	Three grazing intensities
Derner et al. (2006)	321	8.6	Increases	n/d	=	=	1	0–30	Exclosures, 65	Shortgrass
Reeder et al. (2004)	325	n/d	Increases	Increases	Increases	Increases	1	0–90	Exclosures, 56	Heavy grazing
Manley et al. (1995)	338	n/d	Increases	n/d	Increases	n/d	2	0–90	Grazing, 11	Significant only at 0–7.6
Wu et al. (2008)	345	1.1	Decreases	Decreases, =	n/d	n/d	5	0–40	Exclosures, 2 to 27	Chronosequence
Cui et al. (2005)	350	0.2	=	n/d	n/d	Increases	2	0–80	Exclosures, 20	Natural vegetation
Cui et al. (2005)	350	0.2	Decreases	n/d	n/d	n/d	1	0–80	Exclosures, 10	Degraded
Schuman et al. (1999)	384	n/d	Increases	Increases, =	Increases	Decreases	2	0–60	Grazing, 12 (40 Exclosures)	Heavy grazing
Henderson et al. (2004)	350–480	6.25	=	n/d	Increases	n/d	8	0–100	Exclosures, 20 to 71	Correction a to equal mass
Xie and Wittig (2004)	411	5.9	Decreases	n/d	Increases	Decreases	1	0–40	Exclosures, 18	Four levels of grazing
Mikhailova et al. (2000)	587	5.4	=	Increases	Increases	=	1	0–10	Exclosures, 72	Correction a to equal mass
Derner et al. (2006)	588	11.9	=	n/d	=	Decreases	1	0–30	Exclosures, 96	Midgrass
Fuhlendorf et al. (2002)	766	18.3	Decreases	Increases	n/d	Decreases	1	0–10	Exclosures, 50	Restoration
Derner et al. (2006)	835	12.8	Decreases	n/d	=	Decreases	1	0–30	Exclosures, 35	Tallgrass
Potter et al. (2001)	842	17	Increases	=	Increases	n/d	1	0–60	Exclosures, 11	Clay soils
Potter et al. (2001)	842	17	Decreases	Increases	=	n/d	1	0–60	Exclosures, 11	Sandy soils
Piñeiro et al. (2009)	861	15	Increases	Increases	=	Increases	2	0–100	Exclosures, max 30	Lowlands
Chaneton and Lavado (1996)	~ 900	14.5	Increases	Increases	n/d	n/d	1	0–10	Exclosures, 12	Lowlands
Chaneton and Lavado (1996)	~ 900	14.5	=	Increases	n/d	n/d	1	0–10	Exclosures, 16	Uplands
Lavado et al. (1995)	~ 900	14.5	=	=	n/d	n/d	1	0–15	Exclosures, 13	Lowlands
Gill (2007)	932	1.3	=	=	=	n/d	11	0–15	Exclosures, \sim 90	Subalpine
Piñeiro et al. (2009)	1 099	15	Increases	Increases	=	Increases	1	0–100	Exclosures, 5	Lowlands
Piñeiro et al. (2009)	1 099	15	Decreases		Increases	Increases	3	0–100	Exclosures, max 11	Deep soils
Piñeiro et al. (2009)	1 099	15	Increases		Increases	Increases	2	0–100	Exclosures, max 7	Shallow soils
Piñeiro et al. (2009)	1 161	15	Decreases		Increases	Increases	1	0–100	Exclosures, 13	Deep soils
Welker et al. (2004)	1 200	n/d	Increases		=	n/d	1	0–10	Exclosures, 73	No significant difference from 0–60
Piñeiro et al. (2009)	1 300	15	Decreases	Increases	Increases	Increases	3	0–100	Exclosures, 30	Deep soils
Piñeiro et al. (2009)	1 406	15	Decreases		Increases	Increases	1	0–100	Exclosures, 8	Deep soils
(/		15						0–100	Exclosures, 18	Shallow soils

¹"Increases" means that SOC, roots, SOM C:N, or bulk density are higher under grazed conditions. SOC is soil organic carbon; SOM, soil organic matter; n/d, no data.

0

should be co-occurring with compaction to generate the observed patterns in SOC changes.

Based on the influence of the three factors outlined before, and considering that all operate simultaneously, we would expect decreases in SOC contents under grazing in the sites located in the middle of the typical grasslands' mean annual precipitation range ($\sim 400-850$ mm). In this range all the factors would decrease SOC stocks: roots biomass decreases, soil compaction increases, and N limitation increases. Articles reporting data for 15 sites included in this precipitation range showed either a decrease or no changes in SOC contents under grazing compared to the herbivore exclosures, except one site located near the 850 precipitation edge (Table 1).

The studies we reviewed that evaluate SOC changes using the grazing-exclosure approach present at least four shortcomings. The first two are related to the experimental setting, and the last two relate to the type of observation made. The grazing vs. exclosure approach relies on the assumption that differences in space (ungrazed vs. grazed plots) are equal to changes through time (pregrazing vs. postgrazing). Soil heterogeneity can add noise to these comparisons, curtailing their statistical power, and independent variables that are changing during the exclosure treatment period can interact with grazing effects in a complex way (e.g., the increasing atmospheric concentration of CO₂ could have differential effects on grazed vs. ungrazed plots). An additional assumption made in the exclosure approach has to do with the symmetry and reversibility of grazing-exclosure treatments. Strictly, exclosures test the effects of removal of grazing, not the effects of addition of grazing (Burke et al. 1997). Are exclosures representing pregrazing conditions? Or can they just tell us how grazing interruption works, with no clues about the reverse treatment? Observational limitations often arise when SOC data are not accompanied by bulk density measurements, allowing us to estimate C stocks in an equivalent soil mass (Davidson and Ackerman 1993). This often leads to an overestimation of SOC in grazed treatments because of soil compaction (Henderson et al. 2004). Finally a substantial number of studies evaluated only grazing effects on surface soil (first 5 cm or 15 cm), although some more recent studies have explored the whole soil profile (Henderson et al. 2004; Reeder et al. 2004). Surface changes in SOC can differ from whole profile analysis (Baker et al. 2007). All the limitations mentioned above may also contribute to the contradictory effects of grazing on SOM and should be further investigated.

PATHWAYS OF GRAZING EFFECTS: SOC AND SON

To synthesize grazing effects on SOC contents, we grouped the mechanisms proposed by each author to explain their observed SOC changes at grazed vs. ungrazed sites according to different pathways. Most of the papers reviewed explained their results based on one or more mechanisms that could be clumped in three main pathways: grazing altered SOC contents by changing the magnitude or the proportion of NPP that reaches the soil (*NPP pathway*), by changing SON stocks (*nitrogen pathway*), and/or by altering SOM decomposition (*decomposition pathway*; Fig. 2). We propose that grazing may alter

SOC contents simultaneously via these three pathways and thus generate variable responses.

Several authors discussed how grazing affected SOC contents by changing NPP or by affecting the amount of NPP that actually reaches the soil (see the NPP pathway in Fig. 2). Normally an important portion of ANPP is consumed and respired by herbivores, increasing C outputs from the ecosystems and decreasing C available for SOC formation (Fig. 2). Grazing may additionally decrease NPP by removing large portions of aboveground biomass and decreasing leaf area and light interception. In contrast, herbivores also remove or break down standing dead biomass that shades green leaves, and they consume relatively old plant tissues that have low radiation use efficiency (McNaughton 1983; Knapp and Seastedt 1986). The net direct effect of defoliation on NPP will result from the balance of these two opposite mechanisms.

Grazing can also change C allocation patterns affecting the amount of C entering the soil. Belowground biomass directly enters the soil and contributes more to SOM formation than aboveground tissues (Johnson and Matchett 2001). Thus, grazing may affect the amount of C available for SOC formation by changing the proportion of NPP that is allocated to below or aboveground organs. Grazing effects on ANPP are variable, but mostly negative, and do not seem to vary across a precipitation gradient (Oesterheld et al. 1999). In contrast, BNPP seems to increase with grazing (McNaughton et al. 1998; 14 Pucheta et al. 2004; Derner et al. 2006), but again, some studies reported lower BNPP (or root biomass) under grazing (Johnson and Matchett 2001; Kauffman et al. 2004). Our review showed that grazing effects on root biomass varied across a precipitation gradient (Table 1), and that grazing frequently increases the proportion of C allocated to belowground organs, potentially increasing SOM formation.

On the other hand, grazers may change NPP indirectly by altering species composition or soil resources or by decreasing water availability (Fig. 2). Grazing affects community composition in several ways depending on the history of grazing and moisture gradients (Milchunas and Lauenroth 1993; Cingolani et al. 2005). Several mechanisms by which herbivores alter plant community composition have been proposed: selective grazing, changes in resource availability, physical breakdown of fragile species, etc. (Semmartin and Oesterheld 2001; Bakker et al. 2003). Leoni et al. (2008) found that grazing-tolerant species have lower relative growth rates than species present in the ungrazed sites, determining the lower ANPP of grazed sites. The long or short history of grazing also constrains NPP responses to grazing, because sites with a long history of grazing usually have a similar species composition in grazed and ungrazed plots, whereas sites with a short history of grazing usually have contrasting species composition in grazed and ungrazed plots (Milchunas and Lauenroth 1993; Adler et al. 2004; Cingolani et al. 2005). Based on differing grazingresistance plan traits, Adler et al. (2004) suggested that productivity of the upland plant communities in the Patagonian steppe (with a long history of grazing) will be less affected by grazing than the productivity of the sagebrush steppe in the United States (with a short history of grazing). In addition, soil compaction may decrease soil moisture and reduce NPP (Savadogo et al. 2007; Fig. 2). Finally, grazing may also

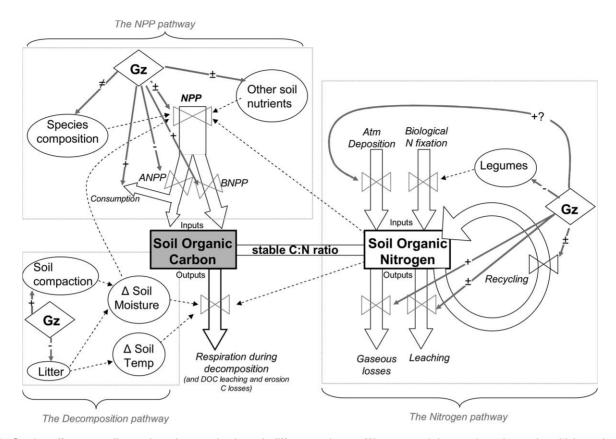


Figure 2. Grazing effects on soil organic carbon stocks through different pathways. We separated three main pathways by which grazing modifies soil organic carbon stocks in rangelands: via affecting net primary production, by changing nitrogen stocks and cycling, or by altering soil organic carbon decomposition. Gz indicates grazing; NPP, net primary production; ANPP, aboveground NPP; BNPP, belowground NPP; and DOC, dissolved organic carbon.

constrain NPP by altering soil nutrients, especially N, as discussed below (Fig. 2).

Herbivory can reduce SOC stocks by increasing decomposition through micrometeorological changes (soil temperature and moisture; see the decomposition pathway in Fig. 2). A grazing-induced reduction of litter pools (Savadogo et al. 2007) would indirectly increase soil temperature and evaporation, which in turn may increase SOM decomposition (Bremer et al. 1998; Burke et al. 1998). Soil temperature in summer can be 10°C higher at grazed sites compared to ungrazed sites (G. Piñeiro, unpublished data). Increases in soil bulk density (see Table 1) reduce infiltration rates and increases soil moisture variations under grazing, potentially increasing SOM decomposition (Abdel-Magid et al. 1987; Savadogo et al. 2007; Fig. 2).

Changes in SOC contents at grazed sites should be related to changes in SON that, in turn, affect both NPP and SOC decomposition rates (see the N pathway in Fig. 2). Grazers can alter SON stocks by both increasing or decreasing N inputs and N outputs. Regarding outputs, grazers promote higher N losses from urine and dung patches (Whitehead and Raistrick 1993) but can also stimulate N retention by decreasing N losses through greater root allocation. Grazing promotes an increase in root biomass at both extremes of the precipitation gradient (Table 1), and such an increase in belowground allocation may result in greater N uptake, lower N losses, and higher N retention (Heckathorn and Delucia 1996; Derner et al. 2006; Piñeiro et al. 2009). Regarding inputs, grazing can decrease N inputs by decreasing legume biomass or cover. Higher N volatilization would increase N redeposition from the atmosphere, partially compensating for N losses, as discussed further (Figs. 1 and 2).

Additionally grazing may either increase or decrease N recycling (Fig. 2). Herbivores accelerate N returns in urine and dung patches. As most grasslands experience some level of N limitation, faster N returns have been suggested as a possible mechanism to explain grazing-induced increases in productivity (Frank and Evans 1997; McNaughton et al. 1997; Holdo et al.

 Table 2. N emissions from herbivores excretions in the Río de la Plata Grasslands.

Productivity $kg \cdot ha^{-1} \cdot yr^{-1}$ (Paruelo et al., this issue)	5 800	
Percentage of aboveground net primary production consumed	58	
(Oesterheld et al. 1992)		
Consumption of forage kg \cdot ha ⁻¹ \cdot yr ⁻¹	3 364	
C consumed kg \cdot ha ⁻¹ \cdot yr ⁻¹	1 682	
C:N ratio of consumed forage (estimated based on a	20	
green:standing dead ratio of 4)		
N consumed kg \cdot ha ⁻¹ \cdot yr ⁻¹	84	
Percentage of N consumed that returns to the soil (Whitehead	70-80	
and Raistrick 1993)		
N emissions kg \cdot ha ⁻¹ \cdot yr ⁻¹	17–25	

Nitrogen Cycle in Grazed Ecosystems

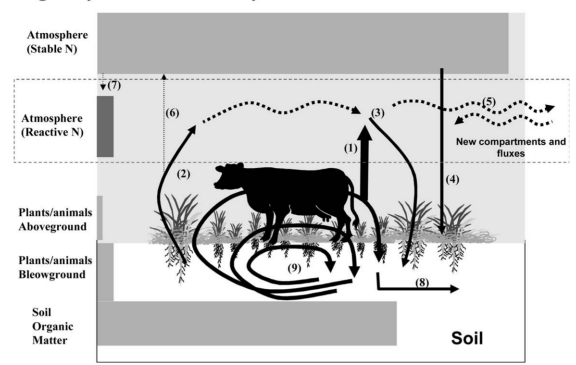


Figure 3. Diagram of the nitrogen cycle in grazed ecosystem, showing traditional fluxes (solid arrows) and pools (light gray boxes) and new fluxes (dashed arrows) and pools (dark gray box) proposed to better understand N cycling at regional scales. The numbers in the figure are 1) N volatilization from urine and dung patches, 2) N reactive emissions from the soil, 3) redeposition, 4) biological N fixation, 5) N exports to adjacent ecosystems, 6) N₂ emissions, 7) conversion of N₂ to N reactive by lightning, 8) leaching, and 9) N recycling through soil biota, vegetation, and herbivores.

2007; Semmartin et al. 2007). However, grazing may accelerate N recycling in nutrient-rich systems but decelerate N recycling in nutrient-poor systems (Frank and McNaughton 1993; McNaughton et al. 1997; Augustine and McNaughton 2006). Lower SON contents or accelerated recycling under grazing could explain observed increases in SOM C:N ratios. If grazing determines a net loss of N, then we would expect an increase in SOM C:N ratios and a decrease in total SON contents. C:N ratios should decrease if C limits SOM formation. Our review shows that grazing mostly increased or did not alter SOM C:N ratios (Table 1), despite increases or decreases in SOC contents. Our result supports the hypothesis that grazing increases N limitation in grasslands.

GRAZING EFFECTS ON ATMOSPHERIC REACTIVE N

Our conceptual models suggest that in grazed systems the atmospheric reactive N should be included as an important N pool. Reactive N pools refer to those N forms that are not stable in atmosphere and biosphere and consequently have significant effects on N dynamics (Schlesinger 1991); they include NH₃, NH₄, NO_x, N₂O (and other inorganic oxidized forms), and other organic compounds (Galloway et al. 2004). As suggested before, grazing may accelerate N recycling but increase N losses. N losses by grazing are mainly gaseous NH₃, a reactive N form. Very simple calculations for the Río de la

Plata Grasslands in South America suggest that 17 kg to 25 kg of N per ha are lost each year from these grasslands (Table 2). Some of this N may be lost by leaching, but probably half or more is lost as NH₃ volatilization. NH₃ has a short residence time in the atmosphere (~ 5 d), and thus it is redeposited near emission sources. In agreement, similar but lower values (~7 kg of N-NH₄ · ha⁻¹ · yr⁻¹) were measured as wet deposition in rain events in Río de la Plata Grasslands, suggesting that some N is being redeposited within the region, but some has been exported out of the region (Piñeiro et al. 2007). Similar results were obtained in North America, indicating a positive trend in regional N deposition associated with increased N emissions generated by larger pig stocks (Aneja et al. 2003).

We suggest that this new compartment (atmospheric reactive N) and four new fluxes should be included explicitly when analyzing N cycling in rangelands. The new fluxes are N emissions in reactive forms, N emissions in nonreactive forms (N₂), local redeposition, and reactive N export to adjacent ecosystems (Fig. 3). Higher reactive emissions (Flux 1 in Fig. 3) can be locally redeposited (Flux 3). Thus, higher N emissions can be partially attenuated by redeposition, but can also be lost by atmospheric transport to other regions (Flux 5), increasing N loadings on them. As a result, grazing "opens" the N cycle, reducing internal recycling (Flux 9), and increases ecosystem dependence on atmospheric loadings. Finally, because N volatilization in grazed ecosystems occurs mainly as NH_3 (Ruess and McNaughton 1988; Whitehead and Raistrick 1993; Frank and Zhang 1997), while N volatilization in ecosystems

without herbivores occurs mainly as N₂O and NO_x (Zheng et al. 2002), herbivores may shift N volatilization from N₂O and NO_x forms to NH₃, potentially reducing the unwanted N₂O and NO_x emissions (Piñeiro et al. 2006). While NO_x has a short residence time in the atmosphere (~ 1 d), N₂O has a relatively long life (~ 150 yr), being more stable in the atmosphere and therefore being a strong greenhouse gas that also depletes O₃ in the stratosphere (Schlesinger 1991). Additionally, NO_x emission generates O₃ in the troposphere, which is also unwanted. Therefore, grazing may alter N fluxes and emission to the atmosphere, altering regional biogeochemistry with implications for climate change and ozone formation (Bouwman et al. 2002; Neff et al. 2002). Soil N storage in grasslands can be a potential solution for decreasing atmospheric reactive N, enhancing both NPP and SOC accumulation.

MANAGEMENT IMPLICATIONS

Identifying the pathway by which grazing may alter SOM stocks in different regions is a key step for designing locally adapted management practices to increase SOC storage. For example, the change in species composition that results as a consequence of grazing may be a key factor for determining the net effect of grazing on SOC changes in a region, since both the NPP pathway and the nitrogen pathway are affected by changes is species composition. It has been shown that intensive grazing can increase forbs with low NPP and thus decrease C inputs to the soil. On the other hand, in other regions shrub encroachment or increased abundance of highly productive species can enhance NPP and thus increase C inputs to the soil under grazing. Ranchers and decision makers should use regional evidence of grazing effects to evaluate which of the three pathways suggested by our model is the most important for determining SOC storage, and then apply management practices to decrease grazing impacts on that/those pathway/s. For example, grazing systems able to manage community composition through seasonal and annual resting periods may counteract the negative impact of grazing on community composition that leads to decreases in SOC, additionally increasing forage quality and accessibility.

Our model can be used to help organize and analyze the complex effects of grazing on rangeland functioning and suggests that one of the key aspects to maximize SOC storage is N dynamics. Net primary production, and particularly BNPP, is an important pathway to increase SOC contents, but the positive effects of grazing on C inputs and belowground allocation would be sustainable only if N losses are reduced or N inputs augmented. Several alternatives aimed to manage N losses and inputs can be implemented; however, caution should be put in avoiding unwanted N emissions such as N2O and NO_x, which can derive from excessive N loadings. A better understanding of source, residence time, and transferences of reactive N within a region would provide a critical piece of information allowing for the design of local management practices aimed at conserving and redistributing N at the landscape level. Higher N conservation will allow grater C sequestration in soils and increased soil fertility, which opens the possibility of a win-win situation in terms of ranchers' economy and global greenhouse gases reduction initiatives.

ACKNOWLEDGMENTS

We thank Emilio Laca and Mary Brooke McEachern who supported and encouraged this work. We also thank four anonymous reviewers for their insightful comments and suggestions on our manuscript.

LITERATURE CITED

- ABDEL-MAGID, A. H., G. E. SCHUMAN, AND R. H. HART. 1987. Soil bulk density and water infiltration as affected by grazing systems. *Journal of Range Management* 40:307–309.
- ADLER, P., D. MILCHUNAS, W. K. LAUENROTH, O. E. SALA, AND I. C. BURKE. 2004. Functional traits of graminoids in semi-arid steppes: a test of grazing histories. *Journal of Applied Ecology* XX:653–663.
- ALTESOR, A. I., E. DI LANDRO, H. MAY, AND E. EZCURRA. 1998. Long-term species change in a Uruguayan grassland. *Journal of Vegetation Science* 9:173–180.
- ALTESOR, A. I., G. PIÑEIRO, F. LEZAMA, R. B. JACKSON, M. SARASOLA, AND J. M. PARUELO. 2006. Ecosystem changes associated with grazing in sub-humid grasslands of South America. *Journal of Vegetation Science* 17:323–332.
- ANEJA, V. P., D. R. NELSON, P. A. ROELLE, J. T. WALKER, AND W. BATTYE. 2003. Agricultural ammonia emissions and ammonium concentrations associated with aerosols and precipitation in the southeast United States. *Journal of Geophysical Research-Atmospheres* 108.
- ASNER, G. P., T. R. SEASTEDT, AND A. R. TOWNSEND. 1997. The decoupling of terrestrial carbon and nitrogen cycles. *BioScience* 47:226–234.
- AUGUSTINE, D., AND S. MCNAUGHTON. 2006. Interactive effects of ungulate herbivores, soil fertility, and variable rainfall on ecosystem processes in a semi-arid savanna. *Ecosystems* 9:1242–1256.
- BAER, S. G., J. N. BLAIR, S. L. COLLINS, AND A. K. KNAPP. 2003. Soil resources regulate productivity and diversity in newly established tallgrass prairie. *Ecology* 884:724–735.
- BAISDEN, W. T., AND R. AMUNDSON. 2003. An analytical approach to ecosystem biogeochemistry modelling. *Ecological Applications* 13:649–663.
- BAKER, J. M., T. E. OCHSNER, R. T. VENTEREA, AND T. J. GRIFFIS. 2007. Tillage and soil carbon sequestration—what do we really know? *Agriculture, Ecosystems & Environment* 118:1–5.
- BAKKER, C., J. M. BLAIR, AND A. K. KNAPP. 2003. Does resource availability, resource heterogeneity or species turnover mediate changes in plant species richness in grazed grasslands? *Oecologia* 137:385–391.
- BEUKES, P. C., AND R. M. COWLING. 2003. Non-selective grazing impacts on soilproperties of the Nama Karoo. *Journal of Range Management* 56:547–552.
- BOND, W. J., AND J. E. KEELEY. 2005. Fire as a global "herbivore": the ecology and evolution of flammable ecosystems. *Trends in Ecology & Evolution* 20:387–394.
- BOUWMAN, A. F., L. J. M. BOUMANS, AND N. H. BATJES. 2002. Estimation of global NH3 volatilization loss from synthetic fertilizers and animal manure applied to arable lands and grasslands. *Global Biogeochemical Cycles* 16:1024.
- BREMER, D. J., J. M. HAM, C. E. OWENSBY, AND A. K. KNAPP. 1998. Responses of soil respiration to clipping and grazing in a tallgrass prairie. *Journal of Environmental Quality* 27:1539–1548.
- BRIGGS, J. M., A. K. KNAPP, J. M. BLAIR, J. L. HEISLER, G. A. HOCH, M. S. LETT, AND J. K. McCARRON. 2005. An ecosystem in transition: causes and consequences of the conversion of mesic grassland to shrubland. *BioScience* 55:243–254.
- BURKE, I. C., W. K. LAUENROTH, AND D. MILCHUNAS. 1997. Biogeochemistry of managed grasslands in central North America. *In:* E. A. Paul, K. Paustian, E. T. Elliott, and C. V. Cole [EDS.]. Soil organic matter in temperate agroecosystems. Boca Raton, FL. USA: CRC Press. p. 85–102.
- BURKE, I. C., W. K. LAUENROTH, M. A. VINTON, P. B. HOOK, R. H. KELLY, H. E. EPSTEIN, M. R. AGUIAR, M. D. ROBLES, M. O. AGUILERA, K. L. MURPHY, AND R. A. GILL. 1998. Plant-soil interactions in temperate grasslands. *Biogeochemistry* 42:121– 143.
- BURKE, I. C., C. M. YONKER, W. J. PARTON, C. V. COLE, K. FLACH, AND D. S. SCHIMEL. 1989. Texture, climate, and cultivation effects on soil organic content in U.S. grassland soils. *Soil Science Society of America Journal* 53:800–805.

63(1) January 2010

6

7

- CHANETON, E. J., J. M. FACELLI, AND R. J. C. LEON. 1988. Floristic changes induced by flooding on grazed and ungrazed lowland grasslands in Argentina. *Journal of Range Management* 41:495–500.
- CHANETON, E. J., AND R. S. LAVADO. 1996. Soil nutrients and salinity after long-term grazing exclusion in a flooding pampa grasslands. *Journal of Range Management* 49:182–187.
- CHAPIN, F. S., P. A. MATSON, AND H. A. MOONEY. 2002. Principles of terrestrial ecosystem ecology. New York, NY, USA: Springer. 436 p.
- CINGOLANI, A. M., I. NOY-MEIR, AND S. DIAZ. 2005. Grazing effects on rangeland diversity: a synthesis of contemporary models. *Ecological Applications* 15:757–773.
- CONANT, R. T., AND K. PAUSTIAN. 2002. Potential soil carbon sequestration in overgrazed grassland ecosystems. *Global Biogeochemical Cycles* 16:1143.
- CUI, X. Y., Y. F. WANG, H. S. NIU, J. WU, S. P. WANG, E. SCHNUG, J. ROGASIK, J. FLECKENSTEIN, AND Y. H. TANG. 2005. Effect of long-term grazing on soil organic carbon content in semiarid steppes in Inner Mongolia. *Ecological Research* 20:519–527.
- DAVIDSON, E. A., AND I. L. ACKERMAN. 1993. Changes in soil carbon inventories following cultivation of previously untilled soils. *Biogeochemistry* 20:161–193.

DERNER, J., T. BOUTTON, AND D. BRISKE. 2006. Grazing and ecosystem carbon storage in the North American Great Plains. *Plant and Soil* 280:77–90.

- FERAL, C. J. W., H. E. EPSTEIN, L. OTTER, J. N. ARANIBAR, H. H. SHUGART, S. A. MACKO, AND J. RAMONTSHO. 2003. Carbon and nitrogen in the soil-plant system along rainfall and land-use gradients in southern Africa. *Journal of Arid Environments* 54:327–343.
- FRANK, A. B., D. L. TANAKA, L. HOFMANN, AND R. F. FOLLETT. 1995. Soil carbon and nitrogen of northern great plains grasslands as influenced by long-term grazing. *Journal of Range Management* 48:470–474.
- FRANK, D. A., AND R. D. EVANS. 1997. Effects of native grazers on grassland N cycling in Yellowstone National Park. *Ecology* 78:2238–2248.
- FRANK, D. A., AND S. J. MCNAUGHTON. 1993. Evidence for the promotion of aboveground grassland production by native large herbivores in Yellowstone National Park. *Oecologia* 96:157–161.
- FRANK, D. A., AND Y. M. ZHANG. 1997. Ammonia volatilization from a seasonally and spatially variable grazed grassland: Yellowstone National Park. *Biogeochemistry* 36:189–203.
- FUHLENDORF, S. D., H. ZHANG, T. R. TUNNELL, D. M. ENGLE, AND A. F. CROSS. 2002. Effects of grazing on restoration of southern mixed prairie soils. *Restoration Ecology* 10:401–407.
- GALLOWAY, J. N., F. J. DENTENER, D. G. CAPONE, E. W. BOYER, R. W. HOWARTH, S. P. SEITZINGER, G. P. ASNER, C. C. CLEVELAND, P. A. GREEN, E. A. HOLLAND, D. M. KARL, A. F. MICHAELS, J. H. PORTER, A. R. TOWNSEND, AND C. J. VOOSMARTY. 2004. Nitrogen cycles: past, present, and future. *Biogeochemistry* 70:153–226.
- GAO, Y., M. GIESE, S. LIN, B. SATTELMACHER, Y. ZHAO, AND H. BRUECK. 2008. Belowground net primary productivity and biomass allocation of a grassland in Inner Mongolia is affected by grazing intensity. *Plant and Soil* 307:41–50.
- GILL, R. A. 2007. Influence of 90 years of protection from grazing on plant and soil processes in the subalpine of the Wasatch Plateau, USA. *Rangeland Ecology & Management* 60:88–98.
- HARPOLE, W. S., D. L. POTTS, AND K. N. SUDING. 2007. Ecosystem responses to water and nitrogen amendment in a California grassland. 2341–2348.
- HASSINK, J. 1997. The capacity of soils to preserve organic C and N by their association with clay and silt particles. *Plant and Soil* 191:77-87.

HECKATHORN, S. A., AND E. H. DELUCIA. 1996. Retranslocation of shoot nitrogen to rhizomes and roots in prairie grasses may limit loss of N to grazing and fire during drought. *Functional Ecology* 10:396–400.

- HEDIN, L. O., P. M. VITOUSEK, AND P. A. MATSON. 2003. Nutrient losses over four million years of tropical forest development. *Ecology* 84:2231–2255.
- HENDERSON, D., B. ELLERT, AND M. NAETH. 2004. Grazing and soil carbon along a gradient of Alberta rangeland. *Journal of Range Management* 57:402–410.
- HOLDO, R. M., R. D. HOLT, M. B. COUGHENOUR, AND M. E. RITCHIE. 2007. Plant productivity and soil nitrogen as a function of grazing, migration and fire in an African savanna. *Journal of Ecology* 95:115–128.

- JOBBÁGY, E. G., AND R. B. JACKSON. 2000. The vertical distribution of soil organic carbon and its relation to climate an vegetation. *Ecological Applications* 10:423–436.
- JOHNSON, L. C., AND J. R. MATCHETT. 2001. Fire and grazing regulate belowground processes in tallgrass prairie. *Ecology* 82:3377–3389.
- KAUFFMAN, J. B., A. S. THORPE, AND E. N. J. BROOKSHIRE. 2004. Livestock exclusion and belowground ecosystem responses in riparian meadows of eastern Oregon. *Ecological Applications* 14:1671–1679.
- KNAPP, A. K., AND T. R. SEASTEDT. 1986. Detritus accumulation limits productivity of tallgrass prairie. *BioScience* 36:662–668.
- LAL, R. 2004a. Soil carbon sequestration impacts on global climate change and food security. *Science* 304:1623–1627.
- LAL, R. 2004b. Soil carbon sequestration to mitigate climate change. *Geoderma* 123:1–22.
- LAVADO, R. S., J. O. SIERRA, AND P. N. HASHIMOTO. 1995. Impact of grazing on soil nutrients in a Pampean grassland. *Journal of Range Management* 49:452–457.
- LEONI, E., A. I. ALTESOR, AND J. M. PARUELO. 2008. Relative growth rates of grazing tolerant and sensible grass species. *Journal of Vegetation Science*. (in press).
- MANLEY, J. T., G. E. SCHUMAN, J. D. REEDER, AND R. H. HART. 1995. Rangeland soil carbon and nitrogen responses to grazing. *Journal of Soil and Water Conservation* 50:294–299.
- McNaughton, S., F. BANYIKWA, AND M. McNaughton. 1997. Promotion of the cycling of diet-enhancing nutrients by African grazers. *Science* 278:1798–1800.
- McNAUGHTON, S. J. 1983. Compensatory plant growth as a response to herbivory. *Oikos* 40:329–336.
- MCNAUGHTON, S. J. 1985. Ecology of a grazing ecosystem: the Serengeti. *Ecological Monographs* 55:259–294.
- McNaughton, S. J., F. F. BANYIKWA, AND M. M. McNaughton. 1998. Root biomass and productivity in a grazing ecosystem: the Serengeti. *Ecology* 79:587–592.
- MIKHAILOVA, E. A., R. B. BRYANT, D. J. R. CHERNEY, C. J. POST, AND I. I. VASSENEV. 2000. Botanical composition, soil and forage quality under different management regimes in Russian grasslands. *Agriculture, Ecosystems & Environment* 80:213–226.
- MILCHUNAS, D. G., AND W. K. LAUENROTH. 1993. Quantitative effects of grazing on vegetation and soils over a global range of environments. *Ecological Monographs* 63:327–366.
- MILCHUNAS, D. G., O. E. SALA, AND W. K. LAUENROTH. 1988. A generalized model of the effects of grazing by large herbivores on grassland community structure. *American Naturalist* 132:87–106.
- NEFF, J. C., R. L. REYNOLDS, M. J. BELNAP, AND P. LAMOTHE. 2005. Multi-decadal impacts of grazing on soil physical and biogeochemical properties in southeast Utah. *Ecological Applications* 15:87–95.
- NEFF, J. C., A. R. TOWNSEND, G. GLEIXNER, S. J. LEHMAN, J. TURNBULL, AND W. D. BOWMAN. 2002. Variable effects of nitrogen additions on the stability and turnover of soil carbon. *Nature* 419:915–917.
- OESTERHELD, M., J. LORETI, M. SEMMARTIN, AND J. PARUELO. 1999. Grazing, fire, and climate effects on primary productivity of grasslands and savannas. *In:* L. Walker [ED.]. Ecosystems of disturbed ground. Amsterdam, the Netherlands: Elsevier. p. 287–306.
- PIÑEIRO, G., E. G. JOBBÁGY, R. B. JACKSON, C. S. SANTONI, S. I. PORTELA, AND C. DI BELLA. 2007. RP-RainNet: the Rio de la Plata Atmospheric Deposition Network: setup and preliminary results. Meeting of the American Geophysical Union–Joint Assembly; Acapulco, Mexico. Eos Trans. AGU, 88(23), Jt. Assem. Suppl. Oral.
- PIÑEIRO, G., J. M. PARUELO, E. G. JOBBÁGY, R. D. JACKSON, AND M. OESTERHELD. 2009. Grazing effects on belowground C and N stocks along a network of cattle exclosures in temperate and subtropical grasslands of South America. *Global Biogeochemical Cycles*. (in press).
- PIÑEIRO, G., J. M. PARUELO, AND M. OESTERHELD. 2006. Potential long-term impacts of livestock introduction on carbon and nitrogen cycling in grasslands of southern South America. *Global Change Biology* 12:1267–1284.
- POTTER, K., J. DANIEL, W. ALTOM, AND H. TORBERT. 2001. Stocking rate effect on soil carbon and nitrogen in degraded soils. *Journal of Soil and Water Conservation* 56:233–236.
- PUCHETA, E., I. BONAMICI, M. CABIDO, AND S. DIAZ. 2004. Below-ground biomass and productivity of a grazed site and a neighboring ungrazed exclosure in a grassland in central Argentina. *Austral Ecology* 29:201–208.

Rangeland Ecology & Management

11

12

0

9

- REEDER, J. D., G. E. SCHUMAN, J. A. MORGAN, AND D. R. LECAIN. 2004. Response of organic and inorganic carbon and nitrogen to long-term grazing of the shortgrass steppe. *Environmental Management* 33:485–495.
- RUESS, R. W., AND S. J. MCNAUGHTON. 1988. Ammonia volatilization and the effects of large grazing mammals on nutrient loss from East African grasslands. *Oecologia* 77:382–386.
- SALA, O. E. 1988. The effect of herbivory on vegetation structure. *In:* M. J. A. Werger, P. J. M. van der Aart, H. J. During, and J. T. A. Verboeven [EDS.]. Plant form and vegetation structure. The Hague: SPB Academic Publishing. p. 317–330.
- SAVADOGO, P., L. SAWADOGO, AND D. TIVEAU. 2007. Effects of grazing intensity and prescribed fire on soil physical and hydrological properties and pasture yield in the savanna woodlands of Burkina Faso. *Agriculture, Ecosystems & Environment* 118:80–92.
- SCHLESINGER, W. H. 1991. Biogeochemistry. An analysis of global change. San Diego, CA, USA: Academic Press. 588 p.
- SCHUMAN, G. E., H. H. JANZEN, AND J. E. HERRICK. 2002. Soil carbon dynamics and potential carbon sequestration by rangelands. *Environmental Pollution* 116:391–396.
- SCHUMAN, G. E., J. D. REEDER, J. T. MANLEY, R. H. HART, AND W. A. MANLEY. 1999. Impact of grazing management on the carbon and nitrogen balance of a mixed-grass rangeland. *Ecological Applications* 9:65–71.
- SEMMARTIN, M., AND M. OESTERHELD. 2001. Effects of grazing pattern and nitrogen availability on primary production. *Oecologia* 126:225–230.
- SEMMARTIN, M., M. OYARZABAL, J. LORETI, AND M. OESTERHELD. 2007. Controls of primary productivity and nutrient cycling in a temperate grassland with yearround production. *Austral Ecology* 32:416–428.
- SHRESTHA, G., AND P. D. STAHLA. 2008. Carbon accumulation and storage in semiarid sagebrush steppe: effects of long-term grazing exclusion Agriculture Ecosystems & Environment 125.
- SMOLIAK, S., J. F. DORMAAR, AND A. JOHNSTON. 1972. Long-term grazing effects on Stipa-Bouteloua prairie soils. *Journal of Range Management* 25:246–250.

- TABOADA, M., AND R. LAVADO. 1993. Influence of cattle trampling on soil porosity under alternate dry and ponded situations. *Soil Use and Management* 9:139–143.
- TIESSEN, H., E. CUEVAS, AND P. CHACON. 1994. The role of soil organic matter in sustaining soil fertility. *Nature* 371:783–785.
- TORN, M. S., S. E. TRUMBORE, O. A. CHADWICK, P. M. VITOUSEK, AND D. M. HENDRICKS. 1997. Mineral control of soil organic carbon storage and turnover. *Nature* 389:170–173.
- VITOUSEK, P., AND W. REINERS. 1975. Ecosystem succession and nutrient retention. *BioScience* 25:376–381.
- WEDIN, D. A. 1995. Species, nitrogen, and grassland dynamics: the constraints of stuff. *In*: C. Jones and J. H. Lawton [EDS.]. Linking species and ecosystems. New York, NY, USA: Chapman & Hall. p. 253–262.
- WEDIN, D. A., AND D. TILMAN. 1996. Influence of nitrogen loading and species composition on the carbon balance of grasslands. *Science* 274:1720–1723.
- WELKER, J. M., J. T. FAHNESTOCK, K. L. POVIRK, C. J. BILBROUGH, AND R. E. PIPER. 2004. Alpine grassland CO2 exchange and nitrogen cycling: grazing history effects, Medicine Bow Range, Wyoming, U.S.A. Arctic, Antarctic, and Alpine Research 36:11–20.
- WHITEHEAD, D. C., AND N. RAISTRICK. 1993. The volatilization of ammonia from cattle urine applied to soils as influenced by soil properties. *Plant Soil* 148:43–51.
- Wu, L., N. HE, Y. WANG, AND X. HAN. 2008. Storage and dynamics of carbon and nitrogen in soil after grazing exclusion in Leymus chinensis grasslands of northern China. *Journal of Environmental Quality* 37:663–668.
- XIE, Y., AND R. WITTIG. 2004. The impact of grazing intensity on soil characteristics of Stipa grandis and Stipa bungeana steppe in northern China (autonomous region of Ningxia). Acta Oecologica 25:197–204.
- ZEIDLER, J., S. HANRAHAN, AND M. SCHOLES. 2002. Land-use intensity affects range condition in arid to semi-arid Namibia. *Journal of Arid Environments* 52:389–403.
- ZHENG, X., C. FU, X. XU, X. YAN, Y. HUANG, S. HAN, F. HU, AND G. CHEN. 2002. The Asian nitrogen cycle case study. *Ambio* 31:79–87.

Authors Queries

Journal: Rangeland Ecology and Management Paper: rama-63-01-05 Title: An Assessment of Grazing Effects on Soil Carbon Stocks in Grasslands

Dear Author

During the preparation of your manuscript for publication, the questions listed below have arisen. Please attend to these matters and return this form with your proof. Many thanks for your assistance

Query Reference	Query	Remarks
1	Author: This article has been lightly edited for grammar, style, and us- age. Please compare it with your original document and make any corrections on these pages. Please limit your corrections to substantive changes that affect meaning. If no change is required in response to a question, please write "OK as set" in the margin. CE	
2	AU: In the author affiliations, is 'Associated Professor' instead of 'Associate Professor' correct? CE	
3	AU: Please include URL of this in your references. CE	
4	AU: Will readers know what 'correc- tion a' means? CE	
5	AU: Please give volume no. CE	
6	AU: Please give page range. CE	
7	AU: Correct that only 1 page? CE	
8	AU: Correct that only 1 page? CE	
9	AU: Please give journal name and volume. CE	
10	AU: Please update publication infor- mation. CE	
11	AU: Please clarify abbreviations at end of this reference and give page range of article. CE	
12	AU: Please update publication infor- mation. CE	

13	AU: Please update publication infor- mation. CE	
14	AU: Please spell out BNPP at first use.	
15	AU: Please provide year for G. Piñeiro unpublished data.	