Soil Organic Carbon Increases in Semi-Arid Regions while it Decreases in Humid Regions Due to Woody-Plant Encroachment of Grasslands in South Africa

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Grasslands and savannas are experiencing intensive land-cover change due to woody plant encroachment. This change in land cover is thought to alter soil carbon (C) and nitrogen (N) storage in these ecosystems. Some studies have reported a negative correlation between soil C and N and mean annual precipitation while others have indicated that there is no relationship with mean annual precipitation. We quantified the changes in C and N pools and $\delta^{13}$C and $\delta^{15}$N values to a depth of 1 m in pairs of encroached and adjacent open grassland sites along a precipitation gradient from 300 mm to 1500 mm per annum in South Africa. Our study showed a negative correlation between changes in soil organic C stocks in the 0–100 cm soil layer and mean annual precipitation (MAP). The most humid site (1500 mm MAP) had less C in shrub-encroached sites while the drier sites (300–350 mm MAP) had more C than their paired open grasslands. This study generally showed soil organic C gains in low precipitation areas, with a threshold value between 750 mm and 900 mm. Our threshold value was higher than that found in North America, suggesting that one cannot extrapolate across continents.

Soils are a major reservoir of terrestrial carbon (C), storing more C (2 344 Gt C, up to 3 m depth) than terrestrial biomass (560 Gt C) and atmospheric pools (750 Gt C)1–3. A small change in soil organic C (SOC) may have a great impact on atmospheric C and subsequently on the earth’s climate4. Land-cover change is one of the main factors that can alter SOC content5,6. One of the major sources of land-cover change, especially in tropical and subtropical regions, is encroachment by native woody plant species. In South Africa, this problem is particularly acute; encroachment affects about 10–20 million ha, seriously reducing the productivity for a country where >70% of its agricultural area is largely grazing lands5.

Jackson et al.8 examined the effects of vegetation change on SOC at six paired grassland and invaded woody sites along a precipitation gradient in the south-western USA, testing relationships between biomass C gains and SOC losses for common native woody invaders, such as Prosopis (mesquite), Larrea (creosote) and Juniperus (juniper) species. In that study, they found that there was a greater increase in SOC stored in the encroached sites than in the open grasslands in drier locations (i.e., below ~400 mm mean annual precipitation (MAP)). However, Jackson et al.8 also observed lower values of SOC in encroached sites than in open grasslands in the more humid locations. Other data show that generally organic C and total N pools in soils beneath shrub canopies increase linearly with time since shrub establishment9. Findings from several studies10,11 indicated that woody plant encroachment is a potentially, but highly uncertain, carbon sink. A subsequent meta-analysis of SOC data on woody plant encroachment found that the change in soil organic C in response to woody plant encroachment was highly variable and unrelated to MAP12.

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Soil bulk density and clay content mediate the magnitude and direction of change in SOC with woody encroachment. In encroached grasslands SOC decreased as soil bulk density increased\(^{12,18}\), while SOC accumulated with increasing clay content\(^{12,14}\). Barger et al.\(^{13}\) reported that SOC accumulated linearly with increasing clay content \(r^2 = 0.76\), presumably reflecting greater occlusion and protection of organic matter afforded by clay micelles\(^{13,14}\). They also found that soil C accumulation decreased as soil bulk density increased, with C gains \(r^2 = 0.52\) being confined to soil densities less than 1.6 g cm\(^{-3}\).

Woody plant encroachment in tropical grasslands usually alters the ecosystem from C\(_4\) plants (African grasses use this photosynthetic pathway to C\(_3\)-dominated woody vegetation\(^{13,14}\). The change in the dominant photosynthetic pathway offers a unique opportunity to use the natural abundance of \(^{13}\)C to evaluate changes in C cycles due to woody plant encroachment of grasslands. Variations in \(^{13}\)C/\(^{12}\)C ratios of plants utilising the C\(_4\) and C\(_3\) photosynthetic pathways provides a natural tracer when a C\(_4\) community type (\(\delta^{13}\)C \(\approx -14\)%o) is replaced by a C\(_3\) community (\(\delta^{13}\)C \(\approx -27\)%o) or vice versa\(^{15,16}\), allowing for the differentiation of the C derived from C\(_4\) vegetation from C derived from C\(_3\)\(^{8,17,18}\).

SOC in encroached grasslands can also be enhanced by an increase in available soil nitrogen. Change in nitrogen availability in ecosystems is likely to drive net primary productivity and thus C sequestration\(^{31}\). The frequent increase of N\(_2\)-fixing woody species in grasslands\(^{20,21}\) has strong potential for altering the N-cycle, primary production, and other key ecosystem processes\(^{21,22}\). Plant species are likely to influence ecosystem N cycling via alterations in N-use efficiency and by changing N inputs and losses\(^{22}\). Changes to the soil total N pool via the shift in the balance of N inputs versus losses should be apparent in the natural abundance of \(^{15}\)N in the plant-soil system\(^{31}\).

In this study, we quantified and compared C and N pools (measured as concentrations and mass stocks) between adjacent woody-encroached and open-grassland plots along a precipitation gradient (300 mm to 1500 mm per annum) in South Africa. The major encroaching species in our sites (except at the 300 mm MAP site in Middelburg) were potentially N\(_2\)-fixing shrubs (\(V\). \textit{karroo}, \(S\). \textit{mellifera}, \(V\). \textit{tortilis} and \(V\). \textit{erioloba}). We focused on responses in C sequestration, and in particular its relationship with MAP, as a key continental scale driver of ecosystem function\(^{24}\). We recorded SOC, N, \(^{13}\)C and \(^{15}\)N at depth intervals of 0–10, 10–30, 30–60 and 60–100 cm and determined the \(^{13}\)C and \(^{15}\)N isotopic ratio in plant leaves and plant litter. We also determined the contribution of C\(_4\) plants to SOC in the encroached grasslands. Our study sites contained mostly warm-season C\(_4\) grasses (Table 1). The sites in decreasing order of precipitation were KwaMbonambi (1500 mm yr\(^{-1}\)), Stanger (900 mm yr\(^{-1}\)), Bergville (750 mm yr\(^{-1}\)), Bloemfontein (500 mm yr\(^{-1}\)), Pniel (350 mm yr\(^{-1}\)) and Middelburg (300 mm yr\(^{-1}\)). At each site, we examined six encroached and six open grasslands. We predicted that in encroached grasslands there would be a negative correlation between changes in SOC and MAP\(^{33}\). We also predicted a negative correlation between C change due to woody plant encroachment and soil bulk density and a positive correlation between C change due to woody encroachment and silt and clay content\(^{10}\). We predicted similar relationships for soil N because of the tight relationship between soil C and N\(^{33}\). We also predicted that \(^{13}\)C will become less enriched in shrub-encroached grasslands compared to open grasslands because of transition from C\(_4\) grasses to C\(_3\) woody-shrubs and trees\(^{15,16}\).

At the most humid site (1500 mm MAP), in the 0–10 cm layer, SOC concentration and SOC stocks were significantly greater \(F_{1,10} = 83.70, P < 0.0001\) and \(F_{1,10} = 103.9, P < 0.0001\) in the open grasslands than in shrub-encroached grasslands (Fig. 1). This humid site lost 50.5% of SOC stocks in the 0–10 cm soil layer (Table 2). The loss in SOC stocks to 100 cm depth due to woody plant encroachment at this humid site was almost 50% at 1500 mm MAP compared to 0–10 cm. At the 900, 750 and 500 mm MAP sites there were generally no differences in SOC concentrations and stocks between encroached and open grasslands in the uppermost soil layer (Fig. 1, Table 2). However, the 750 and 500 mm MAP sites gained carbon stocks slightly at 0–1 m depth, ranging between 9.2% and 25%. In the semi-arid sites of 350 and 300 mm MAP, SOC concentration in the 0–10 cm soil layer was higher \(F_{1,10} = 5.06, P = 0.048\) and \(F_{1,10} = 16.73, P = 0.002, r^2 = 0.62\) and 0.83, respectively) in encroached grassland than in open grasslands (Fig. 1). The 350 and 300 mm MAP sites had higher SOC stocks \(F_{1,10} = 11.26, P < 0.007\) and \(F_{1,10} = 11.26, P < 0.022\) in the encroached grasslands compared to open grasslands in the upper soil layer.

There was a significant negative correlation between change in SOC stocks in open versus encroached grasslands in the 0–100 cm soil layer and mean annual precipitation (MAP) \(r^2 = 0.83, P = 0.012\) (Fig. 2a)). In contrast, there was no significant relationship between the change in SOC stocks (between open and encroached grasslands) and soil bulk density \(r^2 = 0.05, P = 0.68\) (Fig. 2b)) and between change SOC stocks and silt and clay content \(r^2 = 0.02, P = 0.77\) (Fig. 2c)).

Nitrogen was significantly higher in shrub-encroached grasslands than in the open grasslands in the 0–10 cm soil layer in the 900, 500, 350 and 300 mm MAP sites (Fig. 3). The correlation between the change in total N stocks of encroached versus open grasslands in the 0–100 cm soil layer and MAP tended to be negative and marginally significant \(r^2 = 0.62, P = 0.064, r^2 = 0.28\) (data not shown). There was no significant relationship between total N change between encroached and open grasslands and soil bulk density \(r^2 = 0.001, P = 0.89\) or between total N change and silt and clay content \(r^2 = 0.026, P = 0.76\).

In the encroached grasslands, \(^{15}\)N was generally more negative in the plant foliage and plant litter compared to the top soil. The open grassland showed the opposite trend, with the plant material being more enriched in \(^{15}\)N than the top soil (Fig. 4). \(^{15}\)N was generally more enriched in the top soil compared to the plant material in both the encroached and open grasslands (Fig. 4).

At 0–10 cm depth, the \(^{13}\)C values were significantly different between encroached and open grasslands in the 1500, 900, 500 and 350 mm MAP sites while in the 300 and 750 mm MAP sites there was no significant difference in \(^{13}\)C between the two encroached and open grasslands (Fig. 5). There was generally \(^{15}\)C enrichment with increasing depth across the precipitation gradient. There was no significant correlation between the change in \(^{13}\)C between encroached and open grasslands and mean annual precipitation (MAP) \(r^2 = 0.03, P = 0.35\). There was no significant enrichment of \(^{15}\)N with increasing depth in both the encroached and open grasslands (Fig. 6).
However, we found a significant and negative correlation between rainfall and MAP. Additionally, our findings also show that shrub encroachment sequesters more C at MAP compared to the more arid sites because the encroached grasslands had greater biomass than open grasslands. Greater primary productivity supports the idea that shrubs develop extensive root systems that extract nutrients from the horizontal and vertical planes, leading to local nutrient recycling and greater biomass production associated with organic matter accumulation. The accumulation of SOC in the drier shrub-encroached grasslands can also be due to greater rates of primary productivity in the more arid sites because the encroached grasslands had greater biomass than open grasslands. Greater primary productivity increases soil organic matter input in the soil beneath deep-rooted shrubs through increased surface litter and accumulation of complex organic compounds, which are resistant to decomposition.

Our study lacked a direct test of SOC change independent of other covarying factors, such as fire, grazing intensity and microbial activities. Nonetheless, our findings agree with several studies that found a negative correlation between change in SOC and MAP due to shrub encroachment. Our results, however, were contrary to a review by Barger et al. who showed that C response to shrub encroachment was highly variable and unrelated to MAP. Additionally, our findings also show that shrub encroachment sequesters more C at MAP compared to the more arid site (230 mm MAP) having more C than our 500 mm MAP site.

Assuming that there is no isotopic fractionation during decomposition, the isotopic-mixing model indicated that the percentage contribution of C₃ plants (i.e. woody species) to SOC in the encroached grassland increased from 24.7% in the 0–10 cm soil layer to 31.6% in the 60–100 cm soil layer at the 1500 mm MAP site. From the 900 to the 300 mm MAP sites, the contribution of C₃ plants to SOC generally decreased with increasing depth in the encroached grasslands, as shown in Table 3. The C₃ contribution to SOC ranged from 64.6% to 24.7% in the uppermost soil layer and between 8.59 to 35.7% in the 60–100 cm soil layer. The lowest C₃ contribution was at the 1500 mm MAP site which ranged from 3.6 to 17.6%. We acknowledge that the isotopic-mixing model used here does not take into account the fractionation of δ¹⁵N during decomposition. Although there was a positive relationship between δ¹⁵N change and silt and clay content, the relationship was marginally significant ($r^2 = 0.46$). Although there was a positive relationship between δ¹⁵N change and MAP, the variance was also not significantly correlated with MAP ($r^2 = 0.06$, $P = 0.06$). However, we found a significant and negative correlation between δ¹⁵N change and soil bulk density ($r^2 = −0.86$, $P = 0.008$).

Table 1. Study site descriptions.

| Site          | GPS coordinates | Annual Rainfall (mm) | Temperature (°C) | Biome | Soil Characteristics | Major plant species | General land management |
|--------------|-----------------|----------------------|------------------|-------|----------------------|---------------------|------------------------|
| Kwambonambi  | 28°49'61"S 32°16'97"E | 1500 | 3.5–35 | Maputaland wooded grasslands | Quaternary redistributed sands supporting yellowish redistributed sands of the Borea formation | Sporobolus fimbriatus, Digitaria sanguinalis (grass species); Diospyros lyocoles (shrub); Terminalia sericea (tree) | Protected area, privately managed, generally excluded from fires, herbivory activities from wild and livestock |
| Stanger      | 29°18'59"S 31°22'13"E | 900  | 5.8–32.6 | KwaZulu-Natal Coastal Belt | Ordovician Natal group sandstone | Themeda triandra, Aristida junciformis (grasses); Vachellia karroo (tree) | Protected area, privately managed, generally excluded from fires, herbivory from wild animals and livestock |
| Bergville    | 28°7'96"S 29°38'98"E | 750  | 5.8–32.6 | KwaZulu-Natal moist grasslands | Ordovician Natal group sandstone | Themeda triandra, Hysteranthia hirta (grasses); Vachellia karroo, Vachellia sieberiana (trees) | Protected area, generally excluded from fire, herbivory from wild animals and livestock |
| Bloemfontein | 28°59'17"S 26°16'54"E | 450  | 0–32 | Bloemfontein dry grasslands | Sedimentary mudstones and layers of sandstone | Aristida congesta, A. diffusa, Cynodon dactylon (grass); Vachellia karroo (tree) | Protected area, privately managed, generally excluded from fire, herbivory from wild animals and livestock |
| Pietermaritzburg | 28°34'50"S 24°30'30"E | 350  | −4.1–37.5 | Sandy-loam soils of the Hutton soil form | Eragrostis curvula, Schmidia pappophoroides (grasses); Vachellia erioloba, Vachellia tortilis, Vachellia karroo, Senegalia mellifera, Tarchonanthus camphoratus (trees) | Protected area, privately managed, generally excluded from fire, herbivory from wild animals and livestock |
| Middelburg   | 31°25'98"S 24°58'82"E | 300  | −7.2–36.1 | Sandy to loamy soils of the Hutton soil form | Aristida and Eragrostis (grasses); Searsia nesos, S. burchellii, Diospyros lyocoles and Eriocapnos ericoides (shrubs) | Protected area, generally excluded from fire, herbivory from wild animals |

Across all sites. Encroached grasslands had a significantly higher δ¹⁵N than open grasslands in the 500 MAP site ($F_{1, 10} = 8.72$, $P = 0.01$) in the 0–10 cm soil layer (Fig. 6). δ¹⁵N change between encroached and open grasslands was also not significantly correlated with MAP ($r^2 = 0.20$, $P = 0.46$). Although there was a positive relationship between δ¹⁵N change and silt and clay content, the relationship was marginally significant ($r^2 = 0.66$, $P = 0.06$). However, we found a significant and negative correlation between δ¹⁵N change and soil bulk density ($r^2 = −0.86$, $P = 0.008$).
Based on the carbon isotopes, this study also indicated that the current organic matter inputs are not in isotopic equilibrium with soil organic matter in shrub-encroached grasslands. The δ¹³C values of the plant foliage was generally lower than those of soil organic carbon across all sites. In the shrub-encroached grasslands, δ¹³C values of organic inputs were characteristic of C₃ plants (−32 to −28‰) while δ¹³C values of the associated SOC was −25 to −14‰. These contrasting values suggest that our shrub-encroached grasslands dominated by C₃ plants were once occupied by C₄ grasses. δ¹³C soil values in our open grasslands (−18 to −12‰) where generally comparable to other C₄ grasslands and savannas across the world (−27 ± 2.2‰)³⁴. Thus, the δ¹³C values across all our six study sites showed that SOC in open grasslands came from a C₄ dominated ecosystem. The discrepancy between δ¹³C values of vegetation and soils in open grasslands of our drier study sites may reflect lateral intrusion of C₃ plant roots from the shrub-encroached grasslands. Shrubs in semi-arid region have been shown to have extensive roots that can extend to several meters³⁵.

In our drier sites (500, 350 and 300 mm MAP sites), a strong memory of C₄ plants that once dominated these sites was shown by SOC δ¹³C values at depths deeper than 30 cm, while the humid sites (1500 and 900 mm MAP) showed a significant δ¹³C change across the whole range of depth studied. This may indicate a quick SOC turnover in humid sites compared to semi-arid sites. It may also suggest that shrub-encroachment is more recent in

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**Figure 1.** Soil organic carbon concentrations (g C kg⁻¹ soil) vs. depth (cm) along a precipitation gradient (300–1500 mm MAP) in encroached grasslands (black circles) and open grasslands (open circles). Error bars represent standard errors. Some encroached and open grassland data points at the 350 and 300 mm sites completely overlap. Means within the same depth different letters are significantly different (Tukey post hoc test, p < 0.05).
Table 2. Soil organic C and N stocks (Mg/ha) to 1 m depth and % change in C and N after shrub encroachment.

| MAP (mm) | 0–10 cm | 10–30 cm | 30–60 cm | 60–100 cm | 0–100 cm |
|----------|---------|----------|----------|-----------|----------|
|          | Shrub   | Grass    | %change   | Shrub     | Grass    | %change   | Shrub     | Grass     | %change   | Shrub     | Grass     | %change   |
| 1500     | ±0.2    | 9.9 ± 0.4 | 0.5       | ±0.2      | 9.9 ± 0.4 | 0.5       | ±0.2      | 9.9 ± 0.4 | 0.5       | ±0.2      | 9.9 ± 0.4 | 0.5       |
| 900      | ±1.7    | 30.5 ± 1.5 | 9.5       | ±1.7      | 30.5 ± 1.5 | 9.5       | ±1.7      | 30.5 ± 1.5 | 9.5       | ±1.7      | 30.5 ± 1.5 | 9.5       |
| 750      | ±0.7    | 40.0 ± 1.7 | 18.0      | ±0.7      | 40.0 ± 1.7 | 18.0      | ±0.7      | 40.0 ± 1.7 | 18.0      | ±0.7      | 40.0 ± 1.7 | 18.0      |
| 500      | ±1.0    | 12.8 ± 0.5 | 24.8      | ±1.0      | 12.8 ± 0.5 | 24.8      | ±1.0      | 12.8 ± 0.5 | 24.8      | ±1.0      | 12.8 ± 0.5 | 24.8      |
| 350      | ±0.5    | 5.0 ± 0.5 | 108.9     | ±0.5      | 5.0 ± 0.5 | 108.9     | ±0.5      | 5.0 ± 0.5 | 108.9     | ±0.5      | 5.0 ± 0.5 | 108.9     |
| 300      | ±1.2    | 3.9 ± 0.4 | 300       | ±1.2      | 3.9 ± 0.4 | 300       | ±1.2      | 3.9 ± 0.4 | 300       | ±1.2      | 3.9 ± 0.4 | 300       |

Figure 2. Relationship between changes in SOC stocks (%) at 0–100 cm depth with woody encroachment in pairs of encroached and open grasslands and (a) mean annual precipitation (MAP) in mm; (b) silt and clay content as percentage; (c) bulk density in g cm\(^{-3}\). A positive value is achieved because there is more SOC in encroached soils than in open grassland soils and vice versa.

semi-arid sites relative to the humid sites. Recent changes in C\(_4\) to C\(_3\) productivity might be evident in the \(^{13}\)C of SOC near the surface were SOC turnover is most rapid and current organic input are concentrated.

A particularly strong memory of C\(_4\) plants that occupied the shrub-encroached grassland was very evident in \(^{13}\)C values for our 750 mm MAP sites, which ranged from −14 to −12‰ across the soil depth. This finding
indicates that organic carbon at this shrub-encroached site was exclusively from C4 plants and that little SOC was derived from the present C3 vegetation. This very strong C4 memory can be attributed to the high clay content of the soil at the site. The finding is consistent with other studies which showed slow turnover of clay associated organic carbon. Organic carbon associated with clay soils contains aliphatic hydrocarbons which are inert from carbon turnover.

In the 1500 mm MAP site the δ13C signature in encroached grasslands indicated that new SOC from woody plants has been incorporated into the soil. However, the finding that this humid site had low SOC may indicate that the rate at which SOC is being lost is higher than that being added by the C3 plants. The woody plants have added approximately 24.7% of SOC compared to 50.5% lost due to encroachment. This humid site had also the lowest SOC and SN compared to our other humid sites. The isotopic mixing model also shows that C3 SOC contribution for this site increased with depth from 24.3% in the 0–10 cm depth to 31.6% in the 60–100 cm depth.

These findings may indicate high nutrient leaching accelerated by the sandy soil type and high MAP. Although the encroached grasslands are dominated by C3 plants, the bulk of the SOC likely comes from the C4 plants according to the mixed-isotopic model we used. The higher SOC δ13C deeper in the soil may be interpreted as supporting this conclusion, although this depends on there being limited discrimination against 13C during respiration/decomposition.

Figure 3. Soil total-N concentrations (g N kg−1 soil) with depth (cm) along a precipitation gradient (300–1500 mm MAP) in encroached grassland (black circles) and open grasslands (open circles). Means at the same depth with different letters are significantly different (Tukey post hoc test, p < 0.05).
of the sites (300 mm, 350 mm and 750 mm MAP) were open grasslands 20 to 50 years ago. Although 1500 mm MAP site possibly indicated a decrease in the contribution of the C4 plants in SOC with increasing depth, the bulk of the SOC is still potentially derived from the C4 plants. This decrease could also mean that the 1500 mm MAP site has been encroached for a much longer period than in the other sites.

Soil N was higher in the encroached grasslands compared to open grasslands at the 900, 500, 350 and 300 mm MAP sites over the 0–10 cm soil depth probably because most of the encroaching woody species at these sites are leguminous N-fixing plants such as Vachellia erioloba, Vachellia karroo and Senegalia mellifera. N2-fixing plants ameliorate and increase N in the soil. However, at the 1500 mm MAP site, soil nitrogen was significantly higher in the open compared to the encroached grasslands, which is consistent with the SOC trends at this site. This finding is consistent with results from other African savanna ecosystems where sites encroached by woody species have been compared with open grasslands.

There was a general decrease of plant material δ15N values with MAP. This finding is consistent with many other studies that showed a negative correlation between δ15N and MAP. A plausible explanation for this decrease in δ15N with increasing MAP is that photosynthetic activity become more water-than nutrient-limited. Arid environments have high nutrient availability therefore have a more open N cycle resulting in a less negative δ15N compared to humid ecosystems.

There were no significant differences in δ15N values between shrub-encroached and open grasslands across all depths as would be expected if the source of the increase in soil N pools was N fixation. This might suggest other processes affecting 15N concentration, such as N pumping from deeper soil layers and isotope fractionation during litter decomposition and denitrification, to be contributing to soil nitrogen levels in the shrub-encroached grasslands.

This study generally showed shrub encroachment may result in SOC gains in areas with a threshold value between 750 mm and 900 mm MAP. Most of the SOC in the shrub-encroached grasslands came from the C4 plants that previously occupied the area. The significant difference between our study and the North American study of Jackson et al. which showed a threshold of 400 mm indicates that extrapolation across continents is problematic, as has been indicated for savannas elsewhere.

**Methods**

Soil samples were collected between October 2013 and March 2014 at six different sites. The site are described in Table 1 according to Mucina and Rutherford (2010). Each site was divided into encroached and unencroached sections, using adjacent sections to minimize differences in topography. In three of our sites (300, 350 and 750 mm MAP) there is documented evidence, while for the other three sites we have strong oral evidence.
suggesting that our sites were once open grasslands 20–50 years ago. Encroached sites were defined as those areas with >40% shrub cover. Six 1 m-deep and 1 m-wide pits were dug at random at each site in each section. Within each pit, two soil cores of 100 cm³ each were sampled (drilled horizontally into the side of the pit) at centre of each of the four different depth intervals (0–10, 10–30, 30–60 and 60–100 cm). One core was used to quantify soil bulk density (BD) and soil texture, and the other core was used to determine soil organic carbon (SOC), total nitrogen (TN), δ¹³C and δ¹⁵N values. Bulk density and soil texture were quantified using the core method and pipette method, respectively. The soil cores designated for SOC, TN, δ¹³C and δ¹⁵N were air-dried and passed through a 2 mm sieve. Soil samples for SOC and δ¹³C were treated to remove soil carbonates. To remove soil carbonates, soil samples were treated with 1N H₂SO₄/5% FeSO₄ and double-checked for complete inorganic C removal. Soil samples were air-dried at 40 °C and then sieved (2-mm mesh). Generally, our soil samples were free of rocks. Total SOC and N were measured using a Europar elemental analyser (Germany) at BemLabs (Somerset West, Western Cape). The SOC and N stocks were calculated using the following equation:

$$S_x = x_1 x_2 x_3$$

where $S_x$ is the SOC or nitrogen stocks (Mg C ha⁻¹); $x_1$ is the C or N concentration in the soil material (%); $x_2$ is the soil bulk density (Mg m⁻³); $x_3$ is the thickness of the soil layer (cm).

Figure 5. δ¹³C (‰) values in encroached grassland (black circles) and open grasslands (open circles) with depth (cm) in South Africa. Error bars are standard errors of the mean. The reference standard was Pee Dee Belemnite. Means at the same depth with different letters are significantly different (Tukey post hoc test, p < 0.05).
Figure 6. δ¹⁵N (‰) values in encroached grassland (black circles) and open grasslands (open circles) with depth (cm) in South Africa. Error bars are standard errors of the mean. The reference standard was atmospheric nitrogen. Means at the same depth with the same letter are not significantly different (Tukey post hoc test, p < 0.05).

Table 3. Carbon derived from encroaching C₃ woody plants in encroached sites.
We collected plant foliage from the five most abundant species at each site in each landcover area (encroached and open grassland). We sampled litter layer composed of plant debris in different stages of decomposition using a 15 cm × 15 cm wooden frame around each station. Both the plant material and the litter layer were air dried for 3–4 days and then oven dried at 60 °C. The plant foliage and the litter-layer material were then ground before the determination of their δ13C and δ15N signatures.

For δ13C and δ15N values, c. 40 mg of soil, 2 mg of plant foliage and 5 mg of litter-layer material was weighed into tin capsules (Elemental Microanalysis Ltd., Devon, UK) and combusted in a Thermo Flash EA 1112 series elemental analyser; the gases were fed into a Delta Plus XP isotope ratio mass spectrometer (Thermo Electron Corporation, Milan, Italy). The C isotopic ratio of a sample was expressed relative to the Pee Dee Belemnite standard35 and the N isotopic ratios was expressed relative to atmospheric nitrogen36.

Assuming complete and unbiased mixing in the soil, we used the δ13C values of soils (Fig. 5) and our mean δ13C values for the plant material of C3 (shrub) and C4 (grass) species (Fig. 4) present at our sites to estimate the relative proportion (%) of soil organic matter derived from C4 and C3 photosynthetic pathway sources with an isotopic-mixing model46:

\[
FC_3 = \frac{\delta^{13}C_{soil} - \delta^{13}C_{C4}}{(\delta^{13}C_{C3} - \delta^{13}C_{C4})}
\]

where FC3 is the carbon fraction derived from C3 sources, \(\delta^{13}C_{soil}\) is the measured δ13C of the soil sample, \(\delta^{13}C_{C4}\) is the mean δ13C (%) of C4 sources, and \(\delta^{13}C_{C3}\) is the mean δ13C (%) of C3 sources37. The isotopic mixing model used here does not consider δ13C fractionation with depth due to decomposition. Several studies have shown no indication of isotopic fractionation during decomposition45,26, while several other studies have shown various δ13C fractionations due to decomposition27.

The data on SOC and N concentration, and SOC stocks, δ13N and δ15N values attributable to the effects of shrub encroachment and soil depth were analysed using a multivariate ANOVA analysis (MANOVA) to reduce problems with Type I error because we examined several dependent variables simultaneously. The datasets used in this study are available from the corresponding author on request.

References
1. Galy, V. & Eglington, T. Protracted storage of biogenic carbon in the Ganges-Brahmaputra basin. *Nature Geoscience* 4, 843–847, https://doi.org/10.1038/ngeo1293 (2011).
2. Amundson, R. The carbon budget in soils. *Annual Review of Earth and Planetary Sciences* 29, 535–562, https://doi.org/10.1146/annurev.earth.29.1.535 (2001).
3. Dorji, T., Odeh, I. O. A., Field, D. J. & Baillie, I. C. Digital soil mapping of soil organic carbon stocks under different land use and land cover types in montane ecosystems, eastern Himalayas. *Forest Ecology and Management* 318, 91–102, https://doi.org/10.1016/j.foreco.2014.01.003 (2014).
4. Galy, V., Peucker-Ehrenbrink, B. & Eglington, T. Global carbon export from the terrestrial biosphere controlled by erosion. *Nature* 521, 204–207, https://doi.org/10.1038/nature14400 (2015).
5. Edmondsion, J. L., Davies, Z. G., McCormack, S. A., Gaston, K. J. & Leake, J. R. Land-cover effects on soil organic carbon stocks in a European city. *Science of the Total Environment* 472, 444–453, https://doi.org/10.1016/j.scitotenv.2013.11.025 (2014).
6. Zhang, M. et al. Impact of land use type conversion on carbon storage in terrestrial ecosystems of China: a spatial-temporal perspective. *Scientific Reports* 5, 101233 (2015).
7. Hoffman, M. T. & Ashwell, A. Nature divided: land degradation in South Africa. (University of Cape Town Press, 2001).
8. Jackson, R. B., Banner, J. L., Jobbagy, E. G., Pockman, W. T. & Wall, D. H. Ecosystem carbon loss with woody plant invasion of grasslands. *Nature* 418, 623–626, https://doi.org/10.1038/nature00910 (2002).
9. Archer, S., Boutton, T. W. & McMurtry, C. R. In *Global environmental change in the ocean and on land* (eds M. Shiyoumi et al.) 359–373 (Terra pub, 2004).
10. Pacala, S. W. et al. Consistent land- and atmosphere-based U.S. carbon sink estimates. *Science* 292, 2316–2320, https://doi.org/10.1126/science.1057320 (2001).
11. Houghton, R. A., Hackler, J. L. & Lawrence, K. T. Changes in terrestrial carbon storage in the United States: 2. the role of fire and fire management. *Global Ecology and Biogeography* 9, 145–170, https://doi.org/10.1046/j.1365-2699.2000.00164.x (2000).
12. Barger, N. N. et al. Woody plant proliferation in North American drylands: a synthesis of impacts on ecosystem carbon balance. *Journal of Geophysical Research: Biogeochemistry* 116, G00K07, https://doi.org/10.1029/2010JG001506 (2011).
13. Gill, R. A. & Burke, I. C. Ecosystem consequences of plant life form changes at three sites in the semiarid United States. *Oecologia* 121, 551–563, https://doi.org/10.1007/s0044200905962 (1999).
14. Liao, J. D., Boutton, T. W. & Jastrow, J. D. Storage and dynamics of carbon and nitrogen in soil physical fractions following woody plant invasion of grassland. *Soil Biology and Biochemistry* 38, 3184–3196, https://doi.org/10.1016/j.soilbio.2006.04.003 (2006).
15. Balesdent, J., Wagner, G. H. & Mariotti, A. Soil organic matter turnover in long-term field experiments as revealed by carbon-13 natural abundance. *Soil Science Society of America Journal* 52, 118–124, https://doi.org/10.2136/sssaaj1988.03615995005200010021x (1988).
16. Liao, J. D., Boutton, T. W. & Jastrow, J. D. Organic matter dynamics in soil physical fractions following woody plant invasion of grassland: evidence from natural 13C and 15N. *Soil Biology and Biochemistry* 38, 3197–3210, https://doi.org/10.1016/j.soilbio.2006.04.004 (2006).
17. Bernoux, M., Cerri, C. C., Neill, C. & de Moraes, J. F. Tree species effects on soils: implications for global change (eds D. Binkley & O. Menyailo) 155–164 (Springer, 2005).
18. Archer, S., Scifres, C., Bassham, C. R. & Maggio, R. Autogenic succession in a subtropical savanna: conversion of grassland to thorn woodland. *Ecological Monographs* 58, 111–127, https://doi.org/10.2307/1942463 (1988).
19. Boutton, T. W. & Liao, J. D. Changes in soil nitrogen storage and δ15N with woody plant encroachment in a subtropical savanna parkland landscape. *Journal of Geophysical Research: Biogeosciences* 115, G03019, https://doi.org/10.1002/2009JG001184 (2010).
20. Stock, W. D., Wienand, K. T. & Baker, A. C. Impacts of invading *Acacia* species on patterns of nutrient cycling in two Cape ecosystems: evidence from soil incubation studies and 15N natural abundance values. *Ecology* 101, 375–382, https://doi.org/10.1017/BEF00328825 (1995).
21. Knops, J. M. H., Bradley, K. L. & Wedin, D. A. Mechanisms of plant species impacts on ecosystem nitrogen cycling. *Ecology Letters* 5, 454–466, https://doi.org/10.1046/j.1461-0248.2002.00332.x (2002).
24. Knapp, A. K. et al. Shrub encroachment in North American grasslands: shifts in growth form dominance rapidly alters control of ecosystem carbon inputs. Global Change Biology 14, 615–623, https://doi.org/10.1111/j.1365-2486.2007.01512.x (2008).

25. Bosством, R., Comstedt, D. & Eklad, A. Isotope fractionation and 13C enrichment in soil profiles during the decomposition of soil organic matter. Oecologia 153, 89–98, https://doi.org/10.1007/s00442-007-0750-8 (2007).

26. Cleveland, C. C., Neff, J. C., Townsend, A. R. & Hood, E. Composition, dynamics, and fate of leached dissolved organic matter in terrestrial ecosystems: results from a decomposition experiment. Ecosystems 7, 175–285, https://doi.org/10.1007/s10021-003-0236-7 (2004).

27. Fernandez, I., Mahieu, N. & Cadisch, G. Carbon isotopic fractionation during decomposition of plant materials of different quality. Global Biogeochemical Cycles 17, 1075–1086, https://doi.org/10.1029/2001GB001833 (2003).

28. Geesing, D., Felker, P. & Bingham, R. L. Influence of mesquite (Prosopis glandulosa) on soil nitrogen and carbon development: implications for global carbon sequestration. Journal of Arid Environments 46, 157–180, https://doi.org/10.1006/jare.2000.0661 (2000).

29. Wheeler, C. W., Archer, S. R., Asner, G. P. & McMurtry, C. R. Climatic/edaphic controls on soil carbon/nitrogen response to shrub encroachment into grasslands: importance of spatial heterogeneity and vegetation change in a subtropical savanna ecosystem. Geoderma 82, 5–41, https://doi.org/10.1016/S0016-7061(97)00095-5 (1998).

30. Zhou, Y., Boutton, T. W. & Wu, X. B. Soil carbon response to woody plant encroachment: importance of spatial heterogeneity and deep soil storage. Journal of Ecology 105, 1738–1749, https://doi.org/10.1111/1365-2745.12770 (2017).

31. Archer, S. T., Boutton, W. & Hibbard, K. Trees in grasslands: biogeochemical consequences of woody plant expansion in Global biogeochemical cycles in the climate system (ed. E. D. Schulze et al.) 115–138 (Academic, 2001).

32. Allington, G. R. H. & Valone, T. J. Islands of fertility: a byproduct of grazing? Ecosystems 17, 127–141, https://doi.org/10.1007/s10021-013-1041-y (2014).

33. Chitt, T. et al. Impact of woody encroachment on soil organic carbon storage in the Lope National Park, Gabon. Biotropica 49, 9–12, https://doi.org/10.1111/btt.12369 (2017).

34. Boutton, T. W., Archer, S. R., Midwood, A. J., Zitter, S. F. & Bol, R. δ13C values of soil organic carbon and their use in documenting vegetation change in a subtropical savanna ecosystem. Geoderma 82, 5–41, https://doi.org/10.1016/S0016-7061(97)00095-5 (1998).

35. Belskya, A. et al. Influence of trees on savanna productivity: tests of shade, nutrients, and tree-grass competition. Ecology 75, 922–932, https://doi.org/10.2307/1939416 (1994).

36. Bol, R. et al. The 14C age and residence time of organic matter and its lipid constituents in a stagnohumic gley soil. European Journal of Soil Science 47, 213–222, https://doi.org/10.1111/j.1365-2389.1996.tb01396.x (1996).

37. Blaser, W. J., Shanungu, G. K., Edwards, P. J. & Otled Venter, H. Woody encroachment reduces nutrient limitation and promotes soil carbon sequestration. Ecology and Evolutions 4, 1423–1438, https://doi.org/10.1002/ecet.1024 (2014).

38. Hagos, M. G. & Smitt, G. N. Soil enrichment by Acazia melifera subsp. detinos on nutrient poor sandy soil in a semi-arid southern African savanna. Journal of Arid Environments 61, 47–59, https://doi.org/10.1016/j.jaroen.2004.08.003 (2005).

39. Ludwig, F., de Kroon, H., Berendse, F. & Prins, H. H. T. The influence of savanna trees on nutrient, water and light availability and the understory vegetation. Plant Ecology 170, 93–105, https://doi.org/10.1023/B:VEGE.0000019023.29636.92 (2004).

40. Vitousek, P. M., Menge, D. N. L., Reed, S. C. & Cleveland, C. C. Biological nitrogen fixation: rates, patterns and ecological controls in terrestrial ecosystems. Philosophical Transactions of the Royal Society B: Biological Sciences 368, https://doi.org/10.1098/rstb.2013.0119 (2013).

41. Swap, R. J. et al. Natural abundance of 13C and 15N in C3 and C4 vegetation of southern Africa: patterns and implications. Global Change Biology 10, 350–358, https://doi.org/10.1111/j.1365-2486.2003.00702.x (2004).

42. Handley, L. L. et al. The 13N natural abundance (δ13N) of ecosystem samples reflects measures of water availability. Functional Plant Biology 26, 185–199, https://doi.org/10.1071/PP98146, Admore - DW (1999).

43. Hobbie, E. & Ouimette, A. Controls of nitrogen isotope patterns in soil profiles. Biogeochemistry 95, 355–371, https://doi.org/10.1007/s10533-009-9328-6 (2009).

44. Lehmann, C. et al. Savanna vegetation-fire-climate relationships differ among continents. Science 343, 548–552, https://doi.org/10.1126/science.1247355 (2014).

45. Muscina, L. & Rutherford, M. C. The vegetation of South Africa, Lesotho and Swaziland. (South African National Biodiversity Institute, 2010).

46. Ward, D., Hoffmann, M. T. & Colloccott, S. A. A century of woody plant encroachment in the dry Kimberley savanna of South Africa. African Journal of Range and Forage Science. 31, 107–121, https://doi.org/10.2989/12202119.2014.914974 (2014).

47. Grellier, S. et al. The indirect impact of encroaching trees on gully extension: A 64 year study in a sub-humid grassland of South Africa. Catena 98, 110–119, https://doi.org/10.1016/j.catena.2012.07.002 (2012).

48. Britz, M. L. & Ward, D. Dynamics of woody vegetation in a semi-arid savanna, with a focus on bush encroachment. African Journal of Range and Forage Science 24, 131–140, https://doi.org/10.2989/1472373X.2008.22926 (2007).

49. Mureva, A. & Ward, D. Spatial patterns of encroaching shrub species under different grazing regimes in a semi-arid savanna (eastern Karoo, South Africa). African Journal of Range and Forage Science 33, 77–89, https://doi.org/10.2989/12202119.2016.1148775 (2016).

50. Liu, F. et al. Woody plant encroachment into grasslands: spatial patterns of functional group distribution and community development. PLoS ONE 8, e68436, https://doi.org/10.1371/journal.pone.0084364 (2013).

51. Burt, R. Soil survey laboratory methods manual. Soil survey investigations. (United States Department of Agriculture, 2004).

52. Rundel, P. W., Ehleringer, J. R. & Nagy, K. A. Stable isotopes in ecological research. (Springer, 1989).

53. Evans, D. R. Physiological mechanisms influencing plant nitrogen isotope composition. Trends in Plant Science 6, 121–126, https://doi.org/10.1016/S1360-1385(01)01889-1 (2001).

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Author Contributions

Admore Mureva designed, collected and analysed data, wrote and edited the manuscript. David Ward designed, wrote and edited the manuscript. Tiffany Pillay designed, wrote and edited the manuscript. Pauline Chivenge designed, wrote and edited the manuscript. Michael Cramer designed, wrote and edited the manuscript.
Additional Information

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