

Discontinuity in the responses of ecosystem processes and multifunctionality to altered soil community composition

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Ecosystem management policies increasingly emphasize provision of multiple, as opposed to single, ecosystem services. Management for such “multifunctionality” has stimulated research into the role that biodiversity plays in providing desired rates of multiple ecosystem processes. Positive effects of biodiversity on indices of multifunctionality are consistently found, primarily because species that are redundant for one ecosystem process under a given set of environmental conditions play a distinct role under different conditions or in the provision of another ecosystem process. Here we show that the positive effects of diversity (specifically community composition) on multifunctionality indices can also arise from a statistical fallacy analogous to Simpson’s paradox (where aggregating data obscures causal relationships). We manipulated soil faunal community composition in combination with nitrogen fertilization of model grassland ecosystems and repeatedly measured five ecosystem processes related to plant productivity, carbon storage, and nutrient turnover. We calculated three common multifunctionality indices based on these processes and found that the functional complexity of the soil communities had a consistent positive effect on the indices. However, only two of the five ecosystem processes also responded positively to increasing complexity, whereas the other three responded neutrally or negatively. Furthermore, none of the individual processes responded to both the complexity and the nitrogen manipulations in a manner consistent with the indices. Our data show that multifunctionality indices can obscure relationships that exist between communities and key ecosystem processes, leading us to question their use in advancing theoretical understanding—and in management decisions—about how biodiversity is related to the provision of multiple ecosystem services.

aboveground–belowground interactions | ecosystem functioning | plant–soil feedbacks | soil biodiversity | soil fauna

Biodiversity contributes to the functioning of ecosystems by controlling both the rate and the variance of ecosystem processes, making understanding the consequences of biodiversity loss crucial to ecosystem management (1–3). Elucidating the likely impacts of belowground biodiversity loss is particularly important, as soil taxa play key roles in nearly every biogeochemical process that makes Earth an inhabitable planet (4, 5). However, the general relationship between soil biodiversity and ecosystem functioning remains largely unknown because positive, negative, and neutral effects of soil diversity on ecosystem processes are reported (6–10). Similarly idiosyncratic responses of individual ecosystem processes to loss of plant diversity prompted consideration of how biodiversity loss simultaneously affects multiple ecosystem processes, termed “ecosystem multifunctionality” (11). Recent studies on multifunctionality appear to suggest strong and consistent

negative effects of plant diversity loss on ecosystem functioning because species that do not contribute to one ecosystem process may play an important role in a separate process and/or under different conditions (11, 12). Similar assessments for soil biodiversity are in their infancy, but appear to lend support to the idea that the study of single ecosystem processes underestimates the importance of species and functional diversity for ecosystem functioning (13–16).

Soils contain a huge diversity of cryptic organisms living in an opaque environment, complicating direct assessment of the taxonomic, phylogenetic, and functional diversity of soil taxa and, in turn, the consequences of diversity loss in soil. Body size, however, presents a trait by which to manipulate the functional complexity of soil communities that is experimentally tractable as well as relevant to community change (6, 16, 17). First, larger body size is positively associated with susceptibility to mortality from human activities such as forest conversion to cropland and soil tillage (18, 19). Second, body size strongly affects ecosystem processes (2) because (*i*) it correlates with metabolic rate, generation time,

Significance

Ecosystem functioning is more strongly affected by biodiversity loss when multiple functions are considered because different species affect different functions. To quantify these biodiversity–functioning relationships, the emerging multifunctionality framework advocates calculation of indices that aggregate responses of individual functions. Data aggregation, however, is notorious for providing misleading information by obscuring true relationships between explanatory and response variables. We test the ability of common multifunctionality indices to reveal effects on key ecosystem functions of changes in soil communities. The multifunctionality indices all decrease with soil animal loss, but the responses of individual functions diverge markedly from these aggregated metrics. Application of the multifunctionality framework for landscape provision of multiple ecosystem services should therefore emphasize understanding relationships between communities and individual functions.

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population density, and food size (20); (ii) the physical structure of the soil habitat constrains access to resources for certain body sizes and hence modulates interactions between organisms (21); and (iii) the relationship of fauna with microflora shifts from predation to mutualism with increasing body size (22). As such, body size is a trait expected to be strongly related to ecosystem functioning (23).

We simulated the loss of soil functional complexity by excluding larger-bodied soil fauna from model grassland ecosystems. We compared treatments with only microorganisms and microfauna (nematodes and protists; low functional complexity) to those that also contained common larger-bodied meso- and macrofauna, such as springtails, mites, and earthworms (high functional complexity treatment). Taxonomic diversity therefore likely differed little among our experimental grasslands because we did not directly manipulate the microflora, which harbors the majority of soil taxonomic richness (5). We crossed the functional complexity treatment with inorganic nitrogen fertilizer, which is typically applied to more intensively managed grasslands and so is a management that typically accompanies the loss of larger-bodied fauna. Given that a central argument to the adoption of multifunctionality approaches (24, 25) is that different species maintain functions in different environmental contexts, the nitrogen fertilizer treatment allowed us to additionally ask how the effects of the functional complexity treatments were influenced by context. We assessed the responses of five ecosystem processes related to plant productivity, decomposition, ecosystem carbon storage, and respiration.

Loss of soil taxa has divergent effects on individual ecosystem processes, often within the same experimental system (10). Some of these idiosyncrasies may arise because direct effects of the loss of taxa are counterbalanced through indirect effects of soil organisms on plant community composition and diversity (6, 7, 14). Earlier work, for example, in the experimental systems studied here showed that reductions in soil functional complexity decreased litter decomposition, but increased legume abundances (6). The N_2 -fixing ability of legumes might then have compensated for reductions in nitrogen resupply via organic matter breakdown and so sustained similar plant productivity rates in the low- and high-complexity treatments. We therefore hypothesized that two phenomena might diminish the usefulness of multifunctionality indices for understanding soil community relationships with ecosystem functioning: (i) contrasting responses of individual processes to functional complexity are common and arise through a combination of direct and indirect (via the plant community) effects of soil organisms (6, 14); and (ii) the combination of direct and indirect effects alters the soil environment, and hence relationships between complexity and ecosystem processes shift as the context changes in space and time (24, 25). For example, differences in legume abundance between low- and high-complexity treatments might mean that nitrogen fertilization stimulates plant productivity only where legume abundances are low. Such phenomena would mean that amalgamating individual process data to yield aggregated multifunctionality indices could alter, and potentially even reverse, our inferences about causal relationships between community change and ecosystem functioning (26).

Results and Discussion

Multifunctionality. We first examined the relationship between soil functional complexity and ecosystem multifunctionality. We measured rates of ecosystem processes that are either directly or indirectly mediated by soil biota through their effects on nutrient cycling and turnover of organic matter. The processes that we measured were net primary productivity (NPP), net ecosystem productivity (NEP), ecosystem respiration, and litter decomposition. Our approach reflects that of recent research into multifunctionality where investigators focus on the rates of a defined set of biogeochemical processes that together influence the quality

of services that an ecosystem provides (27). We calculated multifunctionality using three distinct methods (28), given that a standard approach has not yet been adopted and all of the proposed metrics have pros and cons (28). Regardless of the approach used, we find that the loss of functional complexity in soil significantly decreases ecosystem multifunctionality (Fig. 1 and Table 1).

The reduction in multifunctionality with the loss of functional complexity is shown most clearly by the averaging approach (15), where the mean index of average function is approximately five times less in the low- vs. high-complexity treatment (Table 1 and Fig. 1A). The effects of functional complexity loss on average multifunctionality were not influenced by nitrogen fertilization (Table 1). It is possible, however, that the averaging approach might give multifunctionality values that are driven by the response of a single ecosystem process (28). A threshold approach overcomes this limitation by scoring how many functions exceed a specified threshold of functioning (12). We explored two thresholds representing high and low functioning—80 and 20% of maximum observed functioning—and found that the soil community manipulations affected multifunctionality significantly only at higher levels of functioning (i.e., the 80% threshold; Table 1). That is, loss of functional complexity impaired multifunctionality only when higher threshold levels of functioning were considered.

One limitation of the single-threshold approach is that it does not necessarily quantify the extent to which functioning is affected, and so a multiple threshold approach has been proposed (28). We used this approach to simultaneously assess the relationship between functional complexity and multifunctionality for all thresholds between 5 and 99%. The effect (i.e., slope) of the loss of functional complexity increased with the threshold value, confirming that the consequences of the soil community manipulations were significant only when considering high levels of functioning (Fig. 1B and C). Specifically, loss of larger-bodied soil fauna began to significantly affect multifunctionality at a threshold value of 72%, peaked at 93%, and remained significantly lower in the low-complexity treatment at the 99% threshold value (Fig. 1C). Thus, loss of functional complexity from the soil communities reduced multifunctionality to about three-fourths of that observed in the high-complexity treatment (i.e., given that effects were evident at thresholds $\geq 72\%$).

Individual Functions. We hypothesized that contrasting responses of individual processes to loss of functional complexity, mediated in part by indirect effects of functional complexity on environmental context, would obscure true relationships between community composition and functioning when aggregated metrics of ecosystem multifunctionality were determined. Despite consistent effects of the functional complexity treatments on the three multifunctionality indices that we calculated (Fig. 1 and Table 1), only two of the individual processes showed significant positive relationships with functional complexity (Table 1). Furthermore, the magnitude and/or direction of the community effect typically strongly contrasted between multifunctionality and all of the individual processes (Fig. 2). We note that relationships between consumer and decomposer diversity on ecosystem multifunctionality, in both aquatic and terrestrial systems, were also consistently positive, but that the underlying process responses were not (16, 27). Our data, therefore, show that whereas functional community complexity has a positive effect on multifunctionality indices, the act of aggregating data to estimate these indices can reverse the direction and/or alter the magnitude of the relationship between community composition and individual process rates (as shown in Fig. 2). As such, multifunctionality indices may obscure insights into the mechanistic relationships required to understand and manage the influence of community change on ecosystem service provision.

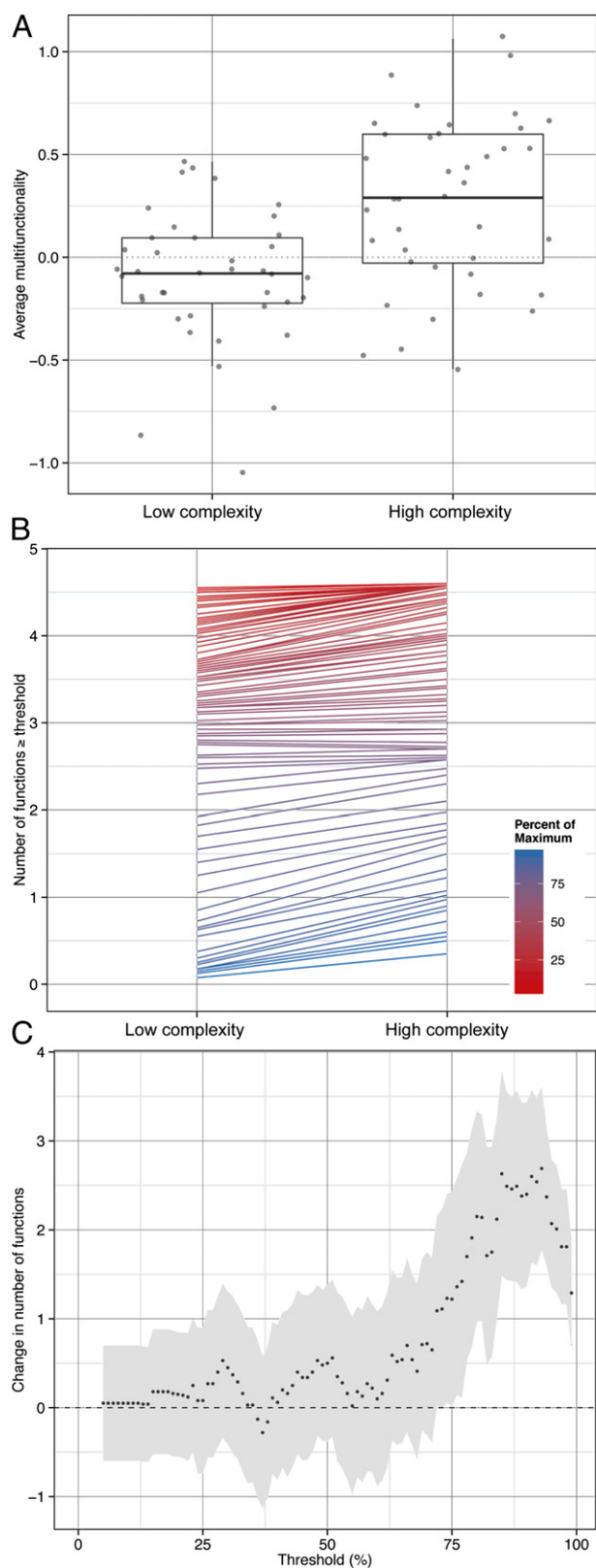


Fig. 1. Loss of soil community functional complexity decreases multifunctionality when measured as (A) average multifunctionality or (B) the number of functions exceeding threshold levels of maximum process rates. Data points in A are jittered to visualize vertical spread, and represent average multifunctionality of each replicate at each of the four time points at which measures were taken. The horizontal line shows the median, the box the 25th and 75th percentiles of the data, and the extent of the whiskers 1.5 times the

Net primary production strongly decreased with loss of functional complexity and increased with nitrogen fertilization (Fig. 2 and Fig. S1). Increased NPP with fertilization is to be anticipated, but the positive effect of larger-bodied fauna on NPP was surprising. Macro- and mesofauna do commonly accelerate processes such as litter decomposition and nutrient cycling (17, 29, 30). However, these effects do not seem to propagate to higher-order processes such as NPP and ecosystem respiration in species-rich plant communities, leading to the expectation that these processes are robust to the loss of larger-bodied fauna (6, 16, 31). Our experiment differs, however, from previous work in the complexity of the systems combined with the long timescale of the experiment. For example, our experimental microcosms took 7 mo to construct, as soil horizons were reconstructed, hundreds of individuals of 10 co-occurring plant species were planted, and soil microflora and fauna were systematically added over many events. The community treatments were then allowed 225 d to develop before the fertilizer treatment was applied and data were collected over a further 280 d (*Materials and Methods*). There was therefore likely enough time and ecological complexity within our experimental grasslands for feedbacks between belowground and plant community composition to develop and translate to alterations in an important ecosystem process (14). These aboveground–belowground interactions influenced all of the ecosystem processes and modified their response to fertilization; we discuss these interactions in turn below.

The marginally significant complexity by fertilization interaction effect on NPP arose because NPP responded strongly only to fertilization in the high-complexity treatment (Fig. 2 and Fig. S1), suggesting stronger plant nitrogen limitation in the absence of larger-bodied fauna. In support of this inference, legume biomass was >30% of NPP in the low-complexity treatment and <1% in the high-complexity treatment (Table S1). In addition, plant-available soil nitrogen was lower on average under high complexity (although this difference was not statistically significant) and was significantly increased by fertilization (Table S1). Furthermore, greater nitrogen limitation is consistent with the fact that the grass biomass in the high-complexity treatment had significantly lower foliar nitrogen contents than in the low-complexity treatment and responded most strongly (although not significantly) to fertilization (Table S1). Our results suggest that, from a management perspective, loss of soil functional complexity might not translate to decreased grassland performance because losses in aboveground yields trade off with increased forage quality (i.e., changes in the plant community through higher grass nitrogen content and higher legume abundance).

The low abundance of legumes and lower nitrogen content of grasses in the high-complexity treatment highlights the ability of belowground community composition to influence aboveground community composition and stoichiometry (7). Although it can be difficult to interpret such single “snapshot” metrics, our process data also support the importance of belowground–aboveground interactions in shaping system functioning. Specifically, the presence of larger-bodied fauna increased, as expected (29, 32, 33), decomposition rates of a standard litter substrate (Table 1, Fig. 2, and Fig. S1). However, the decomposition of the aboveground

interquartile range. (B) Lines represent the slope between soil fauna loss and the number of functions greater than or equal to a threshold value. Separate statistical models were fit at each threshold ranging from 5 to 99% of maximum functioning, with blue lines indicating high percentages of maximum functioning and red lines indicating low percentages of maximum functioning. The curve in C indicates the change in the slope of the relationship described in B across all thresholds with the gray area showing SE. When the error does not cross the zero line (starting at 72%), the loss of soil community functional complexity is associated with a statistically significant decrease in ecosystem multifunctionality.

Table 1. Soil community functional complexity and nitrogen fertilization effects on three different estimates of ecosystem multifunctionality and the five underlying ecosystem processes

Factor	Multifunctionality			Single functions				
	Average multifunctionality	Functioning $\geq 80\%$ threshold	Functioning $\geq 20\%$ threshold	NPP	NEP	Respiration	Standard litter decomposition	Returned litter decomposition
Complexity	0.37*** (0.18, 0.57)	0.40* (−0.02, 0.83)	0.15 (−0.13, 0.42)	42.50*** (28.97, 55.36)	0.35 (−0.25, 0.97)	−0.29*** (−0.42, −0.15)	0.08*** (0.05, 0.11)	0.01 (−0.02, 0.05)
Nitrogen (N)	−0.11 (−0.27, 0.05)	−0.37 [†] (−0.75, 0.01)	−0.25* (−0.45, −0.04)	13.03* (1.71, 24.41)	−1.10*** (−1.63, −0.55)	0.13* (0.00, 0.24)	−0.02 (−0.05, 0.01)	−0.02 (−0.05, 0.01)
Moisture	0.26** (0.09, 0.41)	0.57** (0.18, 0.96)	−0.25* (−0.45, −0.05)	30.43*** (19.52, 42.07)	−0.89** (−1.44, −0.35)	0.37*** (0.26, 0.49)	0.00 (−0.02, 0.03)	0.01 (−0.02, 0.04)
Time	−0.03 (−0.18, 0.13)	−0.33 [†] (−0.70, 0.07)	0.25* (0.05, 0.45)	−6.76 (−17.89, 4.47)	1.24*** (0.70, 1.78)	−0.20** (−0.31, −0.08)	−0.02 (−0.04, 0.01)	−0.01 (−0.04, 0.02)
Complexity: N	0.14 (−0.18, 0.43)	0.50 (−0.24, 1.27)	−0.50* (−0.90, 0.10)	21.63 [†] (−0.43, 43.58)	−0.72 (−1.76, 0.38)	0.24* (0.00, 0.46)	−0.00 (−0.5, 0.05)	0.00 (−0.06, 0.06)
Intercept	0.08 [†] (−0.01, 0.18)	1.90*** (1.68, 2.10)	4.50*** (4.36, 4.64)	87.58*** (81.04, 94.10)	0.11 (−0.20, 0.42)	1.71*** (1.64, 1.77)	0.75*** (0.73, 0.76)	0.67*** (0.66, 0.69)

Values are coefficients with statistical significance from the linear models with the lower and upper bounds shown in parentheses. Coefficients are centered to permit comparison of community and fertilization main effects that are also part of a significant community \times nitrogen interaction. Moisture and time are included in the analysis only to ensure that they do not contribute to the community complexity and fertilization effects (*Materials and Methods*). [†] $P < 0.1$, * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$.

litter returned to the same experimental replicate was unaffected by the community treatments (Table 1, Fig. 2, and Fig. S1). Presumably, the lower foliage litter quality in the high-functional-complexity treatment, due to lower legume abundances and low grass nitrogen contents, retarded decomposition. Such findings highlight the importance of decomposer community composition as an arbiter of how changes in plant communities affect ecosystem processes (14).

Respiration rates were measured at the level of the whole community and so include both autotrophic and heterotrophic ecosystem respiration. Given the greater plant biomass in the high-complexity treatment, both aboveground (i.e., NPP) and belowground (i.e., roots) (Table S1), it is surprising that this greater biomass did not translate to greater respiration rates. In fact, respiration was $\sim 25\%$ lower in the high-complexity treatment (Fig. 2 and Fig. S1). We therefore investigated whether the dominant source of heterotrophic respiration (i.e., soil microbes) was suppressed by the presence of larger-bodied fauna. Soil microbial biomass, however, was unaffected by the complexity and fertilization treatments being 3.6 ± 0.29 mg microbial C g soil^{−1} (mean \pm SE) across all replicates. We did not measure other properties of the microbial community, such as its structure or physiology, which might have helped explain the respiration responses (34). However, a plausible explanation for the higher respiration in the low-complexity treatment and the stronger fertilization effect on respiration in the high-complexity treatment (explaining the complexity \times fertilization interaction; Table 1), is that the responses are plant-mediated. Specifically, N₂ fixation by legumes is energetically demanding (35), and nitrogen contents of foliage correlate with the abundance of the photosynthetic enzyme, rubisco, which incurs a high respiratory cost for maintenance (36, 37). Therefore, we suggest that the low abundance of legumes and the lower nitrogen contents in the grasses in the high-complexity treatment reduced autotrophic respiration, which was then stimulated by nitrogen fertilization because it increased NPP and/or grass nitrogen content, despite reducing root biomass (Table S1 and Fig. 2).

Increases in aboveground NPP under the high-complexity and nitrogen fertilization treatments did not translate to greater net carbon capture (i.e., NEP) because carbon gains through higher plant productivity were generally offset by increased respiratory losses (Fig. 2 and Fig. S1). Indeed, the respiration response to the treatments was always opposite to the NEP response (compare

the sign of the coefficients in Table 1). These opposite responses are best represented by the effects of nitrogen fertilization, which increased aboveground NPP but also stimulated respiration, resulting in a significant and negative effect on NEP (Table S1, Fig. 2, and Fig. S1). Such dynamics represent how interconnected individual biogeochemical processes are in an ecosystem context, resulting in both positive and negative relationships that could aggregate to a common diversity–multifunctionality relationship with different underlying causes.

Implications for Management. The concept of ecosystem multifunctionality is being used to guide management of systems to provide multiple services (38–40), with biodiversity recommended as an ecosystem property that can be managed to increase or sustain multifunctionality (15). Our data show that three different multifunctionality metrics—all applied recently to investigate the consequences of diversity loss from various communities, including plants, soil biota, and salt marsh consumers (16, 27, 28)—show a consistent negative response to loss of soil functional complexity. These consistent responses seem to suggest that multifunctionality indices could help to provide a quantitative basis for improving ecosystem service provision in relation to managing communities. The mismatch between our community and fertilization effects on multifunctionality and the individual processes, however, cautions against using the framework as a predictive tool for achieving desired levels of functioning for multiple, specified ecosystem services. For the framework to be effective as a predictive tool requires that desired individual processes respond to community change in a positive, correlated fashion. In contrast, our data reveal that manipulations of belowground communities can elicit responses of individual processes that are widely divergent and that these divergent responses seem to arise through belowground effects on aboveground communities that feed back to affect ecosystem functioning.

Materials and Methods

Experimental Setup and Design. Ten terrestrial microcosms were established across 7 mo and then maintained in the Ecotron-controlled environment facility (41) for 505 d. The work presented here represents research conducted between experiment day 225 and 505 (280 d total), and research earlier in the experiment showed the potential for the functional complexity manipulations to affect plant community structure and carbon cycle dynamics (6, 29, 42). Each microcosm was 1 m² and housed within a 2 \times 2 \times 2-m walk-in chamber. Full details of the construction and conditions of the microcosms are provided in

with our design] to permit comparison with previous work and because each metric provides unique information (28).

Average multifunctionality determines the average level of a suite of functions by standardizing each function to a common scale and taking their mean (15, 44–47). Inclusion of strongly positively correlated (i.e., $r > 0.5$) individual processes is not recommended (46) in the calculation of multifunctionality indices, but our strongest correlation was 0.37 between the two litter-decomposition processes. We standardized the processes using a z-score transformation, which has advantages over other standardization procedures for the linear model-based statistics that we use (15). Specifically, a standardized function is subtracted by its mean and divided by its SD. The second index, the single-threshold approach, determines if multiple functions are simultaneously maintained above certain levels, determined as whether they exceed a specified threshold percentage of maximum functioning (12, 47, 48). This approach calculates the maximum value of each process across all observations and counts the number of functions—at each observational unit—that exceed a threshold established by the researcher. To define maximum functioning, we took the mean of the five highest values

for each function across all treatment types. We defined single-thresholds of 20 and 80%, representing stronger and weaker community effects on functioning, respectively. The multiple-thresholds approach does not require the investigator to choose a threshold value and instead investigates a continuous gradient of thresholds (28). Specifically, we calculated in 1% increments the values for 5–99% of maximum functioning, in addition to the number of functions per plot exceeding each threshold level. Threshold-based analysis was performed using the *multifunc* package in R (28). Further information on these metrics, as well as the linear mixed models used to assess relationships between the treatments and multifunctionality or single functions, are included in *SI Materials and Methods*.

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- Cardinale BJ, et al. (2011) The functional role of producer diversity in ecosystems. *Am J Bot* 98(3):572–592.
- Hooper DU, et al. (2005) Effects of biodiversity on ecosystem functioning: A consensus of current knowledge. *Ecol Monogr* 75(1):3–35.
- Naeem S, Duffy JE, Zavaleta E (2012) The functions of biological diversity in an age of extinction. *Science* 336(6087):1401–1406.
- Falkowski PG, Fenchel T, Delong EF (2008) The microbial engines that drive Earth's biogeochemical cycles. *Science* 320(5879):1034–1039.
- Wall DH, et al., eds (2012) *Soil Ecology and Ecosystem Services* (Oxford Univ Press, Oxford).
- Bradford MA, et al. (2002) Impacts of soil faunal community composition on model grassland ecosystems. *Science* 298(5593):615–618.
- De Deyn GB, et al. (2003) Soil invertebrate fauna enhances grassland succession and diversity. *Nature* 422(6933):711–713.
- Setälä H, Marshall VG, Trofymow JA (1996) Influence of body size of soil fauna on litter decomposition and ^{15}N uptake by poplar in a pot trial. *Soil Biol Biochem* 28(12):1661–1675.
- Heemsbergen DA, et al. (2004) Biodiversity effects on soil processes explained by interspecific functional dissimilarity. *Science* 306(5698):1019–1020.
- Nielsen UN, Ayres E, Wall DH, Bardgett RD (2011) Soil biodiversity and carbon cycling: A review and synthesis of studies examining diversity–function relationships. *Eur J Soil Sci* 62:105–116.
- Hector A, Bagchi R (2007) Biodiversity and ecosystem multifunctionality. *Nature* 448(7150):188–190.
- Gamfeldt L, Hillebrand H, Jonsson PR (2008) Multiple functions increase the importance of biodiversity for overall ecosystem functioning. *Ecology* 89(5):1223–1231.
- Bowker MA, Maestre FT, Mau RL (2013) Diversity and patch-size distributions of biological soil crusts regulate dryland ecosystem multifunctionality. *Ecosystems* 16:923–933.
- Eisenhauer N, Reich PB, Isbell F (2012) Decomposer diversity and identity influence plant diversity effects on ecosystem functioning. *Ecology* 93(10):2227–2240.
- Maestre FT, et al. (2012) Plant species richness and ecosystem multifunctionality in global drylands. *Science* 335(6065):214–218.
- Wagg C, Bender SF, Widmer F, van der Heijden MGA (2014) Soil biodiversity and soil community composition determine ecosystem multifunctionality. *Proc Natl Acad Sci USA* 111(14):5266–5270.
- Handa IT, et al. (2014) Consequences of biodiversity loss for litter decomposition across biomes. *Nature* 509(7499):218–221.
- Wardle DA (2002) *Communities and Ecosystems* (Princeton Univ Press, Princeton).
- de Vries FT, et al. (2013) Soil food web properties explain ecosystem services across European land use systems. *Proc Natl Acad Sci USA* 110(35):14296–14301.
- Peters RH (1983) *The Ecological Implications of Body Size* (Cambridge Univ Press, Cambridge, UK).
- Brussaard L, et al. (1997) Biodiversity and ecosystem functioning in soil. *Ambio* 26(8):563–570.
- Lavelle P (1997) Faunal activities and soil processes: Adaptive strategies that determine ecosystem function. *Adv Ecol Res* 27:93–132.
- Naeem S, Wright JP (2003) Disentangling biodiversity effects on ecosystem functioning: Deriving solutions to a seemingly insurmountable problem. *Ecol Lett* 6(6):567–579.
- Isbell F, et al. (2011) High plant diversity is needed to maintain ecosystem services. *Nature* 477(7363):199–202.
- Hillebrand H, Matthiessen B (2009) Biodiversity in a complex world: Consolidation and progress in functional biodiversity research. *Ecol Lett* 12(12):1405–1419.
- Gelman A, Shor B, Bafumi J, Park D (2007) Rich state, poor state, red state, blue state: What's the matter with Connecticut? *Quart J Poli Sci* 2:345–367.
- Hensel MJS, Silliman BR (2013) Consumer diversity across kingdoms supports multiple functions in a coastal ecosystem. *Proc Natl Acad Sci USA* 110(51):20621–20626.
- Byrnes JEK, et al. (2014) Investigating the relationship between biodiversity and ecosystem multifunctionality: Challenges and solutions. *Methods Ecol Evol* 5(2):111–124.
- Bradford MA, Tordoff GM, Eggers T, Jones TH, Newington JE (2002) Microbiota, fauna, and mesh size interactions in litter decomposition. *Oikos* 99(2):317–323.
- García-Palacios P, Maestre FT, Kattge J, Wall DH (2013) Climate and litter quality differently modulate the effects of soil fauna on litter decomposition across biomes. *Ecol Lett* 16(8):1045–1053.
- Hunt HW, Wall DH (2002) Modelling the effects of loss of soil biodiversity on ecosystem function. *Glob Change Biol* 8(1):33–50.
- Swift MJ, Heal OW, Anderson JM (1979) *Decomposition in Terrestrial Ecosystems. Studies in Ecology* (Blackwell Scientific, Oxford), Vol 5.
- Wall DH, et al. (2008) Global decomposition experiment shows soil animal impacts on decomposition are climate-dependent. *Glob Change Biol* 14(11):2661–2677.
- Allison SD, Wallenstein MD, Bradford MA (2010) Soil-carbon response to warming dependent on microbial physiology. *Nat Geosci* 3:336–340.
- Vance CP, Heichel GH (1991) Carbon in N_2 fixation: Limitation or exquisite adaptation. *Annu Rev Plant Physiol Plant Mol Biol* 42:373–392.
- Shipley B, Lechowicz MJ, Wright I, Reich PB (2006) Fundamental trade-offs generating the worldwide leaf economics spectrum. *Ecology* 87(3):535–541.
- Tjoelker MG, Oleksyn J, Lorenc-Plucinska G, Reich PB (2009) Acclimation of respiratory temperature responses in northern and southern populations of *Pinus banksiana*. *New Phytol* 181(1):218–229.
- Milder JC, Hart AK, Dobie P, Minai J, Zaleski C (2014) Integrated landscape initiatives for African agriculture, development, and conservation: A region-wide assessment. *World Dev* 54:68–80.
- Lovell ST, Taylor JR (2013) Supplying urban ecosystem services through multifunctional green infrastructure in the United States. *Landscape Ecol* 28(8):1447–1463.
- Schindler S, et al. (2014) Multifunctionality of floodplain landscapes: Relating management options to ecosystem services. *Landscape Ecol* 29(2):229–244.
- Lawton JH (1996) The Ecotron facility at Silwood Park: The value of “big bottle” experiments. *Ecology* 77(3):665–669.
- Bradford MA, et al. (2007) Carbon dynamics in a model grassland with functionally different soil communities. *Funct Ecol* 21(4):690–697.
- Bradford MA, Fierer N, Reynolds JF (2008) Soil carbon stocks in experimental mesocosms are dependent on the rate of labile carbon, nitrogen and phosphorus inputs to soils. *Funct Ecol* 22(6):964–974.
- Hooper DU, Vitousek PM (1998) Effects of plant composition and diversity on nutrient cycling. *Ecol Monogr* 68(1):121–149.
- Maestre FT, Castillo-Monroy AP, Bowker MA, Ochoa-Hueso R (2012) Species richness effects on ecosystem multifunctionality depend on evenness, composition and spatial pattern. *J Ecol* 100(2):317–330.
- Mouillot D, Villéger S, Scherer-Lorenzen M, Mason NWH (2011) Functional structure of biological communities predicts ecosystem multifunctionality. *PLoS ONE* 6(3):e17476.
- Pasari JR, Levi T, Zavaleta ES, Tilman D (2013) Several scales of biodiversity affect ecosystem multifunctionality. *Proc Natl Acad Sci USA* 110(25):10219–10222.
- Zavaleta ES, Pasari JR, Hulvey KB, Tilman GD (2010) Sustaining multiple ecosystem functions in grassland communities requires higher biodiversity. *Proc Natl Acad Sci USA* 107(4):1443–1446.

Supporting Information

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SI Materials and Methods

We used three complementary approaches to measure multifunctionality: weighted average (1, 2), single threshold (3), and multiple threshold (4). To calculate the weighted average, values for each separate process were first standardized by their variance so all processes were comparable in magnitude. The average of all of the processes is then the weighted average metric. For the single-threshold approach, we first calculated the maximum observed value for each process by taking the average of the $n + 1$ highest observed values, where n is the smallest sample size of a single treatment (e.g., low complexity, control). We then selected thresholds of this maximum value at which to assess the effect of the treatments. For each single threshold (20 and 80%), we calculated the number of processes that had values greater than or equal to the chosen threshold of the maximum value. For example, with five processes with maximum values of 20, 10, 30, 70, and 100, our response variable at the 20% threshold would be the number of processes that, for a given replicate microcosm, exceed 4, 2, 6, 14, and 20, respectively. If the values for a replicate were then 5, 7, 5, 13, and 11 for the five processes, respectively, our response variable would equal 2 (of 5 maximum). Given that the determination of these thresholds is not standardized and can be arbitrary (3), we also used a multiple thresholds approach. For this approach, we calculated all of the thresholds between 5 and 99% at 1% intervals (4). We then modeled the relationship between the complexity treatment and the multifunctional response for a continuous range of thresholds. This multiple-threshold approach allowed us to evaluate the thresholds at which the treatments began to have a significant impact on the response (4).

To model the relationship between functional complexity, fertilization, and their interaction on multifunctionality, we used linear mixed models (LMMs). We also used LMMs to assess the

treatment responses of the five ecosystem processes used to calculate multifunctionality. The sign of net ecosystem productivity was reversed so that positive values reflected greater carbon storage. We then considered greater values of all variables to be optimal, as they would indicate greatest rates of plant production, carbon storage, and nutrient turnover. All response variables fit a Gaussian error distribution. Plot was used as a random effect to account for the repeated measurement (four times) of each ecosystem process following fertilization (5). We included moisture (wetter or drier) and measurement time (cut 1–4) as main effects in the models because we recognized that they could explain variance in our data that might otherwise be included in the community complexity and nitrogen effects. Moisture and time effects were not, however, explored beyond model inclusion because they were not the main focus of the study and we did not have enough degrees of freedom to test how they interacted with complexity or fertilization. Our models contained a mix of categorical and continuous factors, which have different variances, and so we centered categorical data by subtracting the mean and standardized continuous variables by subtracting the mean and dividing by two SDs (6). An advantage of this analytic approach is that centering our independent variables makes main effects interpretable even when involved in interactions (7, 8).

The LMMs were fit using the lme4 package for the R statistical program (9). The F-statistic is not considered accurate for the lme4 package (10), so we used a Markov Chain Monte Carlo approach to estimate coefficients and P values. All reported P values are quasi-Bayesian, rather than the classical frequentist P values, and, like the coefficients, retain the same interpretation. We considered coefficients with $P < 0.05$ significant and coefficients with $P < 0.10$ marginally significant (11). Supporting metadata (e.g., root densities) were also analyzed using LMMs but without time because these metrics were assessed only at the end of the experiment.

1. Maestre FT, et al. (2012) Plant species richness and ecosystem multifunctionality in global drylands. *Science* 335(6065):214–218.
2. Wagg C, Bender SF, Widmer F, van der Heijden MGA (2014) Soil biodiversity and soil community composition determine ecosystem multifunctionality. *Proc Natl Acad Sci USA* 111(14):5266–5270.
3. Zavaleta ES, Pasari JR, Hulvey KB, Tilman GD (2010) Sustaining multiple ecosystem functions in grassland communities requires higher biodiversity. *Proc Natl Acad Sci USA* 107(4):1443–1446.
4. Byrnes JEK, et al. (2014) Investigating the relationship between biodiversity and ecosystem multifunctionality: Challenges and solutions. *Methods Ecol Evol* 5(2):111–124.
5. Bolker BM, et al. (2009) Generalized linear mixed models: A practical guide for ecology and evolution. *Trends Ecol Evol* 24(3):127–135.
6. Gelman A (2008) Scaling regression inputs by dividing by two standard deviations. *Stat Med* 27(15):2865–2873.
7. Schielzeth H (2010) Simple means to improve the interpretability of regression coefficients. *Methods Ecol Evol* 1(2):103–113.
8. Schielzeth H, Nakagawa S (2013) Nested by design: Model fitting and interpretation in a mixed model era. *Methods Ecol Evol* 4(1):14–24.
9. R Core Team (2012) *R: A Language and Environment for Statistical Computing* (R Foundation for Statistical Computing, Vienna).
10. Baayen RH, Davidson DJ, Bates DM (2008) Mixed-effects modeling with crossed random effects for subjects and items. *J Mem Lang* 59:390–412.
11. Hurlbert SH, Lomabardi CM (2009) Final collapse of the Neyman-Pearson decision theoretic framework and rise of the neoFisherian. *Ann Zool Fenn* 46(5):311–349.

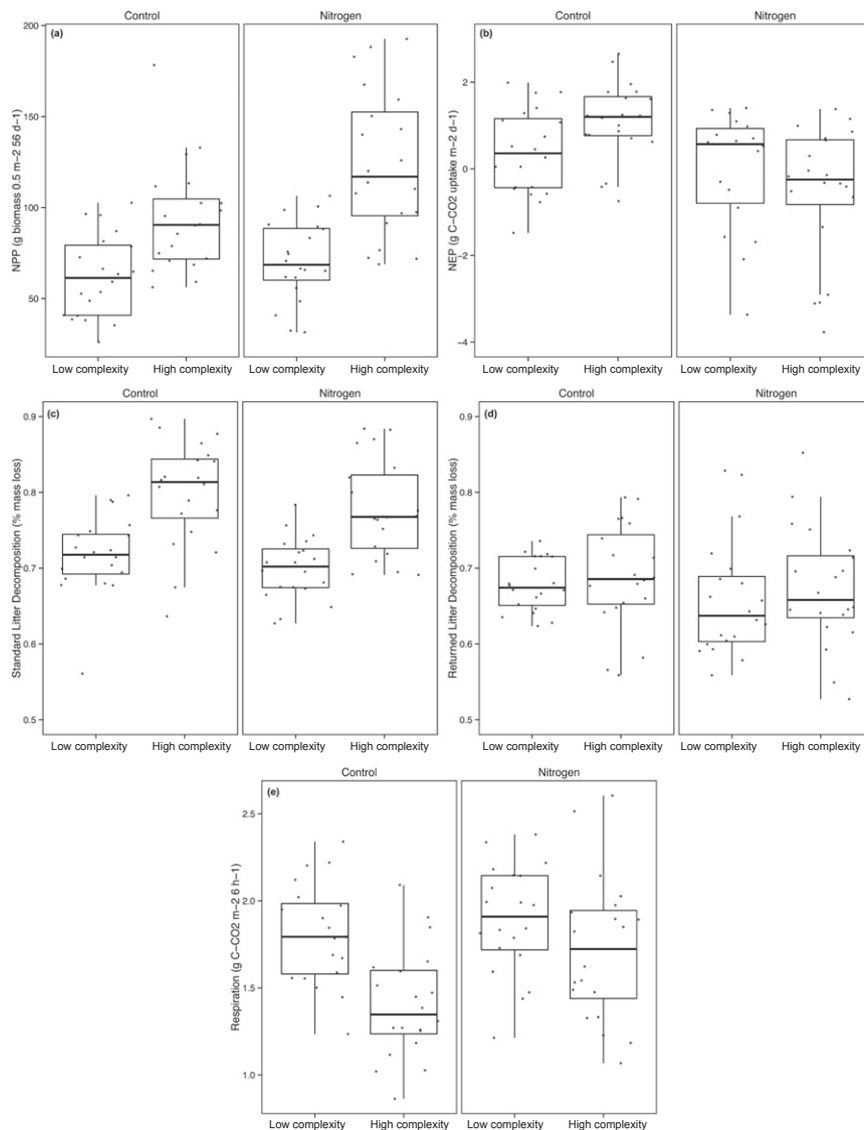


Fig. S1. Effects of soil community functional complexity loss and nitrogen fertilization on the rates of the five biogeochemical processes evaluated: (A) net primary productivity (NPP); (B) net ecosystem productivity (NEP); (C) decomposition of a standard litter; (D) decomposition of litter returned within each replicate community; and (E) total community respiration. The horizontal line in each box plot shows the median, the box indicates the 25th and 75th percentiles of the data, and the extent of the whiskers 1.5 times the interquartile range. Data points are jittered to visualize vertical spread and represent replicate values for each of the four time points that measures were taken. The statistical significance of the community complexity by nitrogen treatments is given in Table 1.

Table S1. Soil community functional complexity and nitrogen fertilization effects on ecosystem and community properties of the experimental grasslands

Variables	Control		Nitrogen added		Coefficients and significance	
	Low complexity	High complexity	Low complexity	High complexity	Complexity	Nitrogen
Legume biomass (% community)	38.7 ± 10.02	0.1 ± 0.05	33.6 ± 10.23	0.1 ± 0.05	-36*** (51, -20)	NS
Grass nitrogen (% mass)	2.46 ± 0.21	1.98 ± 0.16	2.47 ± 0.24	2.22 ± 0.15	-0.36 [†] (-0.81, 0.05)	NS
Roots—organic (g·m ⁻³)	4,679 ± 972	8,282 ± 2196	3,873 ± 1019	3,520 ± 326	NS	-2,467 [†] (-5241, 253)
Roots—mineral (g·m ⁻³)	1,058 ± 209	1,828 ± 265	1,103 ± 134	1,257 ± 255	0.34* (0.01, 0.68)	NS
Soil NH ₄ ⁺ (μg N g·soil ⁻¹)	17.5 ± 1.45	15.2 ± 3.04	19.4 ± 2.44	28.0 ± 6.38	NS	7.96* (0.58, 16.42)
Soil NO ₃ ⁻ (μg N g·soil ⁻¹)	5.0 ± 0.64	4.0 ± 0.96	10.1 ± 2.75	8.6 ± 1.96	NS	0.61** (0.21, 1.01)

Shown are attributes of the aboveground plant community (legume and grass), root densities, and estimates of plant-available soil nitrogen. Values are means ± SE ($n = 5$) at the end of the experiment, and the final two columns show the coefficients with statistical significance from the linear models with the lower and upper bounds shown in parentheses. Interactions between the community complexity and nitrogen fertilization effects are not shown because none were significant. [†] $P < 0.1$, * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$; NS, not significant.