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Simple measures of climate, soil properties and plant traits predict national-scale grassland soil carbon stocks

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Summary

1. Soil carbon (C) storage is a key ecosystem service. Soil C stocks play a vital role in soil fertility and climate regulation, but the factors that control these stocks at regional and national scales are unknown, particularly when their composition and stability are considered. As a result, their mapping relies on either unreliable proxy measures or laborious direct measurements.

2. Using data from an extensive national survey of English grasslands, we show that surface soil (0–7 cm) C stocks in size fractions of varying stability can be predicted at both regional and national scales from plant traits and simple measures of soil and climatic conditions.

3. Soil C stocks in the largest pool, of intermediate particle size (50–250 μm), were best explained by mean annual temperature (MAT), soil pH and soil moisture content. The second largest C pool, highly stable physically and biochemically protected particles (0.45–50 μm), was explained by soil pH and the community abundance-weighted mean (CWM) leaf nitrogen (N) content, with the highest soil C stocks under N-rich vegetation. The C stock in the small active fraction (250–4000 μm) was explained by a wide range of variables: MAT, mean annual precipitation, mean growing season length, soil pH and CWM specific leaf area; stocks were higher under vegetation with thick and/or dense leaves.

4. Testing the models describing these fractions against data from an independent English region indicated moderately strong correlation between predicted and actual values and no systematic bias, with the exception of the active fraction, for which predictions were inaccurate.

5. *Synthesis and applications.* Validation indicates that readily available climate, soils and plant survey data can be effective in making local- to landscape-scale (1–100 000 km^2) soil C stock predictions. Such predictions are a crucial component of effective management strategies to protect C stocks and enhance soil C sequestration.

Key-words: carbon sequestration, carbon storage, community weighted mean, particle size fractions, pH, soil carbon, soil organic matter

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Introduction

Soil carbon (C) stocks exceed those in both vegetation and the atmosphere by 2–3 times and play a vital role in

climate regulation and the maintenance of soil fertility (Trumper *et al.* 2009), but these stocks vary by orders of magnitude over regional and national scales, even within a single ecosystem type (Bellamy *et al.* 2005; Carey *et al.* 2008). Presently, their mapping relies upon either proxy measures that are often poor estimates of actual soil C stocks, particularly at local scales (Jones *et al.* 2005; Eigenbrod *et al.* 2010; Stevens *et al.* 2013), or direct measurements, which are expensive and laborious (Bellamy *et al.* 2005; Carey *et al.* 2008). Models are also used to predict soil C, but these are typically used to make large-scale or scenario-based projections and not fine-scale, extensive soil C stock mapping (Schimel *et al.* 1994; Smith *et al.* 2005).

Improved predictions of soil C stocks should be possible if the factors determining national, regional and local distributions of soil C are better understood. It has long been known that soil C is controlled by a wide range of factors (Jenny 1941; Schimel *et al.* 1994) that can be viewed as forming a 'hierarchy of controls' (Díaz *et al.* 2007; De Vries *et al.* 2012), which impact the basic processes of plant growth and organic matter (OM) decomposition and stabilization. At the apex of the hierarchy is climate, which controls the metabolism of plants, fauna and microbes (Burke *et al.* 1989; Schimel *et al.* 1994; Conant *et al.* 2011) and determines rates of rock weathering (White 2005), thus influencing soil properties. The next level in the hierarchy are soil abiotic properties, such as texture and pH, which are largely controlled by underlying geology and processes of weathering (Jenny 1941; White 2005), and which in turn influence soil C storage by affecting plant growth and microbial activity (Pietri & Brookes 2008; Schmidt *et al.* 2011). At a local level, soil C storage is also strongly affected by land-use type and intensity, which has an array of impacts on soil C cycling (Conant, Paustian & Elliot 2001; Smith 2014). Moreover, climate, soil properties and management all influence the composition and growth of the vegetation, which in turn affects the amount and chemistry of plant inputs, and the turnover of soil organic matter (SOM; Cornwell *et al.* 2008; De Deyn, Cornelissen & Bardgett 2008; De Vries *et al.* 2012).

Although it has long been acknowledged that the above factors are the primary regulators of soil C storage, their interdependence makes estimating their *relative* importance challenging. While some studies emphasize the importance of the physical and chemical properties of soil (Christensen 2001; Schmidt *et al.* 2011), there is also evidence that plant community composition plays a significant role (De Deyn, Cornelissen & Bardgett 2008). While the importance of vegetation properties has long been recognized, and is represented in C models (Parton *et al.* 1993; Smith *et al.* 2005), they have typically been viewed only from a tissue chemistry perspective, and the importance of other plant traits has rarely been considered. This may be an oversight as plant species vary along a 'fast-slow' traits axis, which distinguishes between fast-growing

species with rapidly decomposing litter and fast tissue turnover times and their opposite (Reich 2014). Accordingly, if species effects on ecosystem function are proportional to their biomass (Grime 1998), then community abundance-weighted means (CWMs) of species-level traits may explain variation in soil C storage and sequestration (Garnier *et al.* 2004). In line with this prediction, recent work shows that CWM trait measures can explain ecosystem-level variation in plant production, decomposition, photosynthesis, respiration and soil C concentration, and microbial community composition (Garnier *et al.* 2004; Díaz *et al.* 2007; De Vries *et al.* 2012; Grigulis *et al.* 2013; Everwand *et al.* 2014). While such studies point to the tractability of scaling up from traits of individual plants to ecosystem properties, the capacity of this approach to predict soil C at spatial scales large enough to matter to C stock management is unknown.

Another drawback of existing methods of soil C stock prediction is that they typically predict only the total amount of soil C and not its composition (Jones *et al.* 2005; Stevens *et al.* 2013). Soil C is diverse in its chemistry and interactions with soil particles, and as a result, soil C particles vary greatly in their turnover rates (Trumbore 2000; Schmidt *et al.* 2011). Accordingly, soil C storage and sequestration is determined not just by the total soil C pool, but also by the half-lives of soil C components, which can be categorized into pools of varying stability (Schimel *et al.* 1994; Trumbore 2000). Such pools are arbitrarily defined but are represented in modern soil C models. Measuring them is inherently difficult, so we lack reliable baseline data on the amount of C in these pools for most of the Earth's land surface. While isotopic techniques (^{13}C and ^{14}C) can be employed (e.g. Trumbore 2000; Marschner *et al.* 2008), their use is impractical in large-scale surveys given their high cost and requirement for specialist equipment and personnel. An alternative approach is to use more readily measurable size and density fractions, which broadly correspond to C turnover times (Christensen 2001; Marschner *et al.* 2008). Fresh C inputs are predominantly found in large particles that constitute the active fraction, which turns over within months to a few years, making it the source of most soil C fluxes. In contrast, C found in particles of intermediate size is typically humified OM that turns over on decadal time-scales, while small and dense soil particles of physically and chemically protected soil comprise the stable C fraction. This typically turns over on the scale of centuries to millennia and is crucial to soil C sequestration (Schimel *et al.* 1994; Trumbore 2000; Christensen 2001). While relationships between many of the aforementioned drivers and *total* soil carbon over large scales have been quantified previously (e.g. Burke *et al.* 1989), their relationship with different C size fractions has received very little attention (Evans, Burke & Lauenroth 2011). The relative importance of the aforementioned drivers in determining stable soil C may differ from those controlling rapid turnover fractions, and this could explain discrepancies

between studies of soil C drivers, which typically focus upon total soil C.

In this study, we set out to identify which factors best explain national-scale patterns of different C fractions in the surface soil (0–7 cm) of grassland. This was done for two reasons: (i) to identify the potential abiotic and biotic (i.e. plant traits) determinants of these fractions at large spatial scales and (ii) to assess the potential for using a combination of simple plant trait and abiotic measures that are readily available to surveyors to predict these soil C stocks, that is to identify potential variables to be used in pedotransfer functions and/or ecological production functions for these fractions. To do this, we generated linear mixed-effects statistical models describing national-scale patterns of surface soil C in different size fractions across a wide spectrum of the soil and climatic conditions found across England, and a broad range of grassland types (including calcareous, mesotrophic, wet and acid, Rodwell 1992). These quantified the relative importance and predictive capacity of several abiotic factors and various CWM plant traits with strong hypothetical or known links with soil C cycling (De Deyn, Cornelissen & Bardgett 2008). We hypothesized that stocks of the active soil C fraction are best predicted by the drivers of plant inputs to soil and the decomposability of these inputs (e.g.

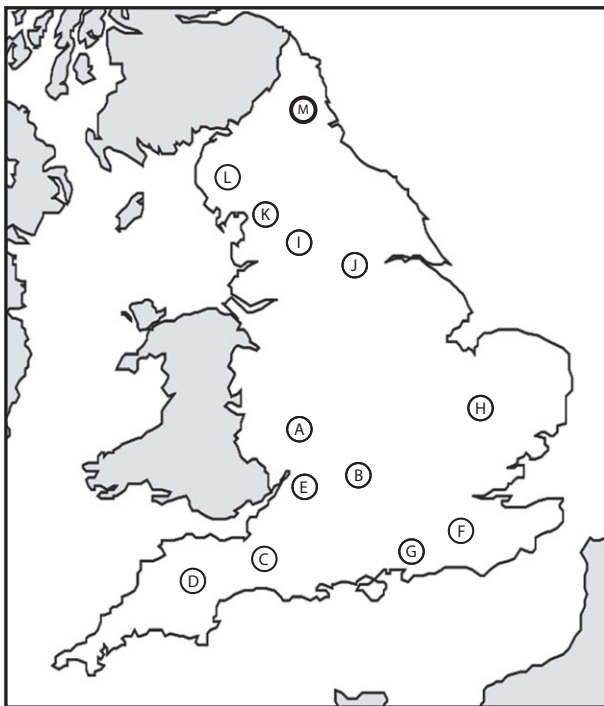


Fig. 1. Sampling regions within England. In each region, five farms were selected, and in each of these, three fields were sampled, one unimproved grassland, one semi-improved and one improved. Regions are (a) Worcester, (b) Upper Thames, (c) Somerset, (d) Devon, (e) Cotswolds, (f) High Weald, (g) South Downs, (h) Breckland, (i) Dales Meadows, (j) Yorkshire Ings, (k) Yorkshire Dales/South Lake District, (l) Lake District. In the validation region (m) Northumberland, 20 fields were sampled.

climate and plant traits), while the stable fraction is better explained by physical and chemical properties of soil (e.g. soil texture and pH). We focussed on grasslands because they cover *c.* 30% of the Earth's land surface and store *c.* 23% of the global terrestrial ecosystem C stock (Trumper *et al.* 2009). Moreover, in the United Kingdom (UK), where our study was performed, they cover 36% of the land surface and contain an estimated *c.* 32% of national soil C stocks (Ostle *et al.* 2009).

Materials and methods

GRASSLAND SURVEY

This work was conducted as part of a broader investigation aimed at quantifying relationships between agricultural intensification, botanical composition and soil properties, including microbial community composition, in temperate grasslands (De Vries *et al.* 2012). We sampled from 12 English regions during June and July 2005 (see Fig. 1). Within each region, there were five sites, each containing three fields, but subject to three broad management regimes: unimproved (U) and often designated as Site of Special Scientific Interest, semi-improved (SI) or improved (I) grassland, resulting in a total of 180 fields (Fig. S1, Supporting information). The survey represented the broad habitat classifications of acid (33 fields), calcicolous (42 fields), mesotrophic (81 fields) and wet grasslands (24 fields), the main grassland types in the UK (Rodwell 1992), and fields were allocated to land management intensity categories based on consultation with farmers and land managers, and expert opinion. This process also ensured that adjacent fields were of similar soil type and topography. Typically, unimproved grasslands receive $<25 \text{ kg N ha}^{-1} \text{ year}^{-1}$ and are lightly grazed by livestock and cut annually for hay, whereas SI and improved grasslands receive 25–50 and $>100 \text{ kg N ha}^{-1} \text{ year}^{-1}$, respectively, and are subject to higher grazing pressures and more frequent cutting for silage (Critchley, Fowbert & Wright 2007). This broad classification of grasslands has been used widely (e.g. De Vries *et al.* 2012; Grigulis *et al.* 2013) and reflects the typical grassland farming systems that are found across the UK and other parts of Europe (Rodwell 1992).

There were many different plant community types present in the more botanically diverse unimproved grasslands, but the improved categories were mainly the *Lolium perenne* (L.) dominated MG6 and MG7 communities of the UK's National Vegetation Classification (Rodwell 1992). Within each field, percentage cover of each plant species was visually estimated from three random 1-m² quadrats within a 25 × 25 m plot of homogeneous vegetation. These three cover values were averaged to obtain field-level abundance estimates. Within each quadrat, five random 2-cm-diameter, 7-cm-deep soil cores were taken and pooled. The use of 7-cm-deep cores follows the UK's Department of Environment, Food and Rural Affairs (DEFRA) recommended sampling depth for assessment of soil abiotic properties in permanent grassland (DEFRA 2010), and was selected to capture the zone of soil most influenced by plant traits, and of greatest C content relative to subsurface soil. We recognize that significant soil C stocks are found at depth in grasslands (Jobbagy & Jackson 2000), but sampling the whole soil profile was beyond the scope of this study, especially given the comprehensive suite of vegetation and soil properties measured.

SOIL ANALYSIS

Soil samples were sieved (4 mm), homogenized and air-dried, and analysed for moisture content, total C and pH, using standard methods (see Allen 1989 and Appendix S1 for methods). Standardized wet sieving (De Deyn *et al.* 2011) was then used to separate the soil particles and the C within them into the active (250–4000 μm), intermediate (50–250 μm) and stable fractions (0.45–50 μm) (see Appendix S1 for details). To calculate soil C stocks on a per-area basis, bulk density was calculated from core volume and dry soil weight after removing all stones and roots >3 mm diameter. It is possible that black C (charcoal) and inorganic C were present in our samples, though the proportion of these is likely to be small (see Appendix S1). Soil texture was classified by expert judgment and transformed into clay–silt–sand percentages using the central point of each category of the triangular classification developed by the Soil Survey of England and Wales (Hodgson 1997).

CLIMATE DATA

Both long-term climate data from Met Office UKCP09 data bases (Jenkins, Perry & Prior 2009) and the grassland survey data were assigned to 5 × 5 km grid cells. Mean annual temperature (MAT) and mean annual precipitation (MAP) were calculated from monthly data from 1981 to 2006. Mean growing season length (MGSL) values were taken from the UKCP09 data base containing monthly values from 1961 to 2003 and calculated as the number of days bounded by a daily temperature mean >5 and <5 °C after 1st July for more than five consecutive days. Mean growing degree-days (MGDD) was the day-by-day sum of the mean number of degrees by which air temperature exceeded 5.5 °C. It was calculated using the mean of values from 1961 to 2006. The differences in time periods between these measures reflect data availability in the UKCP09 data base.

PLANT TRAIT DATA

Plant species composition data were combined with data base values of plant traits to give field-level CWMs for plant traits with hypothetical links to soil processes (Garnier *et al.* 2004; Díaz *et al.* 2007; De Deyn, Cornelissen & Bardgett 2008; De Vries *et al.* 2012). To do this, trait values were assigned to all plant species occurring in the 180 fields sampled, and plant cover was used as the CWM weighting measure. Values for leaf dry matter content (LDMC) were taken from a published account of plant species in northern England (Grime, Hodgson & Hunt 2007). Values for specific leaf area (SLA), relative growth rate (RGR) and leaf nitrogen content (leaf N) were obtained from the TRY data base (Kattge *et al.* 2011, Appendix S2), which contains trait data from a wide range of authors and environments. See Appendix S1 for details of trait measurement and justification of trait choice.

STATISTICAL MODELLING

The grassland survey, climate and trait data were combined to form a single data set (see Table S1 to see the range of conditions covered) that was used to parameterize and test the likelihood of competing mixed-effects statistical models according to a model selection procedure (Pinheiro & Bates 2000). A separate model

was created to describe each soil C fraction as well as total C. Our model selection approach involved adding explanatory variables in fixed sequential order according to our hypothesized ‘hierarchy of controls’ (Appendix S1, Díaz *et al.* 2007). The process started with variables describing climatic conditions (MAP, MAT, MGSL, MGDD) and then added physical and chemical properties of soil that are driven mainly by underlying geology and local hydrology (soil pH, sand silt and clay content and soil moisture). The third set of terms was linked to management. Here, contrasts were made between three competing management terms, which either had three levels of U, SI and I or two, with either SI and U or SI and I merged. Finally, we added trait CWMs to estimate plant functional trait effects. CWMs for RGR, SLA, LDMC and leaf N were placed in the model in all combinations of one and two traits. Although microbial data were available (De Vries *et al.* 2012), they were not used to predict C stocks as they are not readily measurable by most surveyors. All models were linear mixed-effects models with a random effect for site to account for the spatial clustering of triplicate fields. Mixed models were fitted using maximum likelihood and the lme function of the statistical software R version 2.11.1 (Pinheiro & Bates 2000). Throughout the modelling process, quadratic terms were used when the optimum of biological activity occurs at intermediate levels (i.e. for temperature, pH and moisture). See Appendix S1 and Table S2 for details.

The explained variance (EV) of the final model was calculated as the r^2 when fitting a linear regression to the actual data, with the predicted values of the model as the explanatory variable. To estimate the variance explained by the fixed effects, we used the method of Nakagawa & Schielzeth (2013), which partitions the EV by comparing the fit of model predictions to the data when these terms are absent from the model to calculate a ‘marginal R^2 ’ ($R^2\text{M}$). We also used this technique to estimate the proportion of unique (total) variance explained by each class of variable in the model (soil, climate, plant traits). The importance of each variable in the model was also estimated by observing Akaike Information Criterion (AIC) change (Δi) on deletion.

MODEL VALIDATION

To validate the fitted models, we collected new data for all the variables retained in the models (Table 1) in 20 fields in the county of Northumberland, England, in summer 2012. This is a separate region to the north-east of the original 12 (Fig. 1). Methodology was identical to before with the exception of site selection. In this case, we intentionally chose sites covering a wide range of the predictor variables found in the original data set, but excluded sites from outside these ranges to avoid extrapolation (Table S1), because our models were not mechanistic. To validate the fitted models, predictor variable values for the Northumberland sites were fed into the fitted models to produce estimated values. These were then compared to actual values using Pearson’s correlation and paired t -tests.

Results

Total soil C stock to 7-cm depth was best described (Tables 1 and S3; EV = 74.2%, $R^2\text{M}$ = 26.9%) by a quadratic relationship for MAT (Fig. 2a), with C stocks being lowest at intermediate temperatures of *c.* 8.5 °C.

This temperature effect accounted for 13.7% of unique variance. Variation in total soil C stock was also related to soil pH, with stocks being lowest at intermediate pH values of *c.* 6 (Fig. 2a) (quadratic relationship). Finally, soil C stocks were related to soil moisture and maximal at moisture levels of *c.* 100%, on a dry soil weight basis. Together, these soil terms accounted for 15.2% of unique variance.

Models explaining the three component fractions differed greatly in the variables they contained, indicating that each is controlled by different factors. The active fraction (4000–250 µm) (Fig. 3, Tables 1 and S4) accounted for 11.1% of total C stocks across grasslands, and the model describing it accounted for 41.0% of its variation ($R^2M = 37.6\%$) and contained five variables, each strongly linked to plant productivity and litter decomposition. The most important of these were quadratic relationships with MAT (Fig. 2b) and MAP; stocks of this C fraction were highest in cold, wet conditions. This pool was also positively associated with MGSL, presumably via higher net primary productivity, and resulting inputs of C to soil (Table 1). Together, these climate factors accounted for 22.0% of unique variance. Soil pH accounted for 8.7% of unique variance and also displayed a quadratic relationship with the active C fraction, being greatest in acidic soils. Finally, we found that the active C fraction was predicted by the CWM of SLA; stocks were

higher under vegetation with thick and/or dense leaves. This trait measure accounted for 4.3% of unique variance.

The intermediate fraction (50–250 µm) represented 54.7% of total soil C stocks to 7-cm depth across grasslands (Fig. 3) and was described by a model that was very similar to that describing total C stocks (EV = 78.4%, $R^2M = 19.9\%$, Tables 1 and S5); the retained terms described quadratic relationships with MAT (Fig. 2c), soil moisture and pH (Fig. 2c). Stocks of this C fraction were lowest in soils of neutral grassland and at intermediate climates (MAT *c.* 9 °C, pH *c.* 6). Of the retained variables, climate terms were marginally more important (11.8% unique variance) than soil terms (9.6% unique variance).

The stable soil C fraction (0.45–50 µm), which comprised 32.4% of the total C stocks across grasslands (Fig. 3), was not explained by climate or management variables. The model (EV = 74.2%, $R^2M = 17.53\%$, Tables 1 and S6) indicated a strong and quadratic relationship with soil pH, with the highest stocks being found in neutral and alkaline grassland soils (Fig. 2d). C stocks in this fraction also increased subtly with increasing CWM leaf N content. This trait effect accounted for far less variance (1.9% unique variance) than pH (14.16% unique variance).

Comparison of predicted and observed values of soil C stocks demonstrated that the fitted models made reason-

Table 1. Selected models for total soil carbon to 7-cm depth and soil carbon fractions of a range of size classes (kg C m⁻²)

Variable	Total soil carbon			Active fraction (4000–250 µm)			Intermediate fraction (50–250 µm)			Stable fraction (0.45–50 µm)		
	Param. est.	AIC change (Δi)*	P-value*	Param. est.	AIC change (Δi)*	P-value*	Param. est.	AIC change (Δi)*	P-value*	Param. est.	AIC change (Δi)*	P-value*
Intercept	64.35		<0.0001	12.57		<0.0001	39.21		0.0001	4.22		0.12
MAP (mm)			0.0001	–6.4e-4	14.21	0.0001						
MAP (mm) ²			0.0001	5e-6	18.71	0.0009						
MAT (°C)	–10.72	14.64	0.025	–2.28	17.99	<0.0001	–6.91	10.37	0.0008			
MAT (°C) ²	0.62	13.23	0.0067	0.11	8.97	<0.0001	0.40	10.81	0.0003			
Soil moisture (% dry weight)	0.035	–3.36	0.0002				0.021	2.05	0.049			
Soil moisture (% dry weight) ²	–1.8e-4	–5.36	<0.0001				–1.1e-4	4.02	0.014			
MGSL (days)				0.010	4.17	0.013						
Soil pH	–5.86	18.89		–0.99	5.51	0.0086	–2.95	9.38	0.0012	–1.59	17.57	<0.0001
Soil pH ²	0.52	11.64		0.077	4.3	0.012	0.26	6.69	0.0032	0.16	2.35	0.037
CWM SLA (mm ² mg ⁻¹)				–0.024	9.09	0.0009						
CWM leaf N content (mg N g ⁻¹)										0.039	4.13	0.01

AIC, Akaike Information Criterion; MAP, mean annual precipitation; MAT, mean annual temperature; CWM, community weighted mean; MGSL, mean growing season length; SLA, specific leaf area.

*Assessed with a likelihood ratio deletion test. This was a single d.f. test for most terms but two for the main effects of variables with a quadratic function. In these cases, both the main effect and the quadratic term were removed.

ably reliable predictions of observed stocks of total C and the intermediate and stable fractions, but not the active fraction (Fig. S2). Correlations between predicted and observed values were $r = 0.57\text{--}0.64$, and there was no significant difference between them (paired t -tests $P > 0.05$, $t = <2$, d.f. = 19), with the exception of the active fraction ($r = 0.03$, $P < 0.05$) (Table S7, Fig. S1).

Discussion

Our results indicate that regional and national patterns of C fractions in the surface soils of grasslands can be predicted using fairly simple measures of the abiotic environment and community-level plant traits. Reasonably accurate prediction of soil C stocks across broad gradients of soil and ecosystem types has been achieved previously using dynamic models (e.g. Parton *et al.* 1993) and proxy measures (Paruelo *et al.* 1998; Jones *et al.* 2005). However, it has not, to our knowledge, been achieved for different size fractions of soil C within a single land-use type, as shown here. The relationships identified here may not always be mechanistically causa-

tive because climate, management and underlying geology all directly affect soil C stocks while also selecting for different plant trait syndromes (De Vries *et al.* 2012), making trait measures an integrated measure of the environment. Nevertheless, all terms in the models accounted for unique variation, meaning that these relationships strongly indicate the primary regulators of these soil C fractions. Importantly, we found that several factors that influence soil C stocks at small scales, such as agricultural management (Conant, Paustian & Elliot 2001) and soil texture (Christensen 2001), do not explain national patterns in C stocks at these shallow depths. In contrast, plant traits did explain the C stocks of certain fractions. The surprising lack of soil texture effects on soil C pools may be because soil C is controlled by physical properties of soil that were not captured by our field assessment, for example mineral surface charges (Schmidt *et al.* 2011), and secondary and tertiary aspects of soil structure that determine the availability of C to decomposers, for example compaction and microaggregates (Christensen 2001). Alternatively, the lack of soil texture effects may be due to low data resolution or the rarity

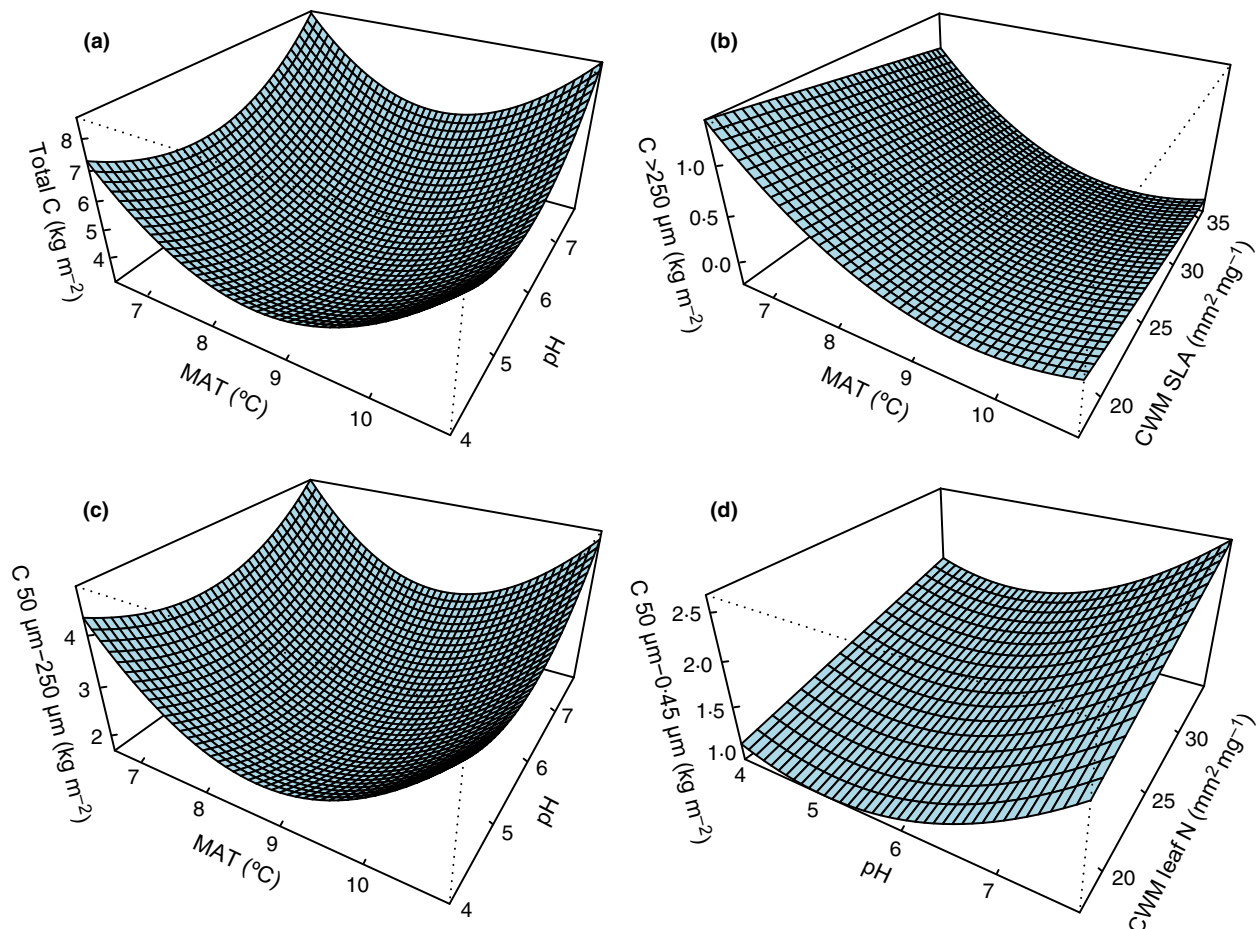


Fig. 2. Fitted relationships between abiotic and plant community trait variables and grassland soil carbon stocks. In all figures, the other variables in the models (Table 1) were held constant at their mean in the data set (Table S1). Relationships are between (a) mean annual temperature (MAT) and pH with total soil carbon stocks; (b) MAT and mean annual precipitation (MAP) with carbon in the active fraction; (c) soil pH and MAT with carbon in the intermediate fraction; and (d) soil pH and the community abundance-weighted mean (CWM) of leaf nitrogen content and carbon in the stable fraction. Stocks are for the top 7 cm of the soil.

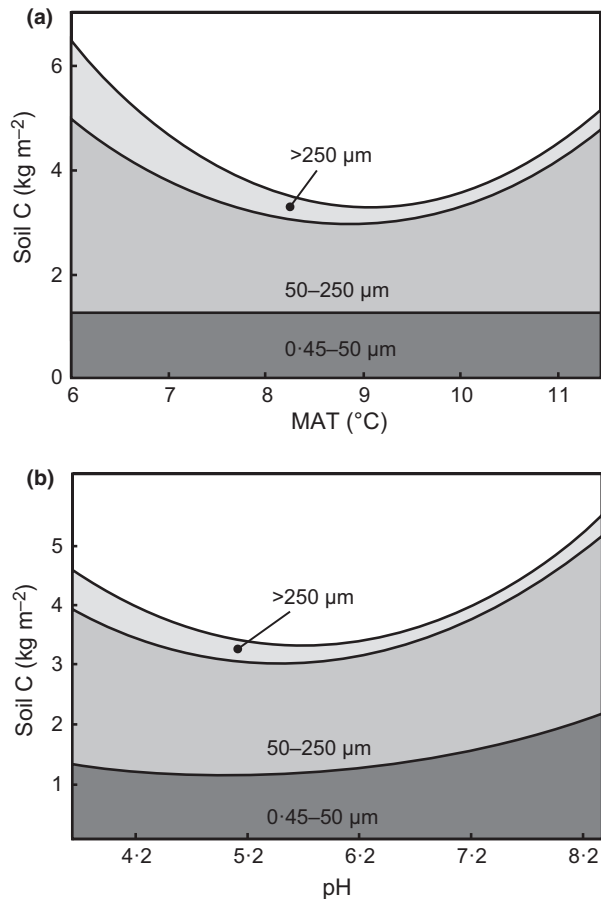


Fig. 3. Changes in grassland soil carbon stocks across (a) temperature and (b) soil pH gradients. MAT, mean annual temperature. The stocks shown are the three size fractions predicted by the fitted models when all other variables are held constant at their mean in the data set (Table S1).

of clay-rich soils sampled (Table S1). We also highlight that although plant traits explained a small proportion of variance, their real importance may be greater than in our models, given their correlation with many of the other descriptor variables and their basal position in our hierarchy of controls modelling procedure.

Looking at each model in turn provides insight into the factors driving each pool and emphasizes the need to view soil C as a heterogeneous material when attempting to understand its dynamics and meaningfully quantify C stocks. The active fraction model demonstrates that stocks in this fraction are highest where plant growth is high (high MAT and MGSL), but decomposition is possibly slow (low pH and high MAP) (Cornwell *et al.* 2008; Pietri & Brookes 2008). There was also, as hypothesized, a strong relationship with the CWM of SLA. Where vegetation possessed leaves that were thin and/or low density and lacked more slowly decomposing structural materials (Reich 2014), stocks of this fraction were lower (Fig. 2b), a finding that is consistent with previous studies linking SLA to litter decomposition rates (e.g. Garnier *et al.* 2004). The poor predictive capacity of our

active fraction model may be due to the dynamic nature of this pool, which is highly variable seasonally (Christensen 2001). It may be better predicted by models in which plant production and decomposition are more explicitly defined.

Unlike the other C fractions, the stable pool was not explained by climate, possibly because much of this C would have entered this pool and become stabilized in different climatic conditions to those experienced today. This finding is consistent with some large-scale gradient studies, which show stable soil C stocks to be largely insensitive to temperature (Conant *et al.* 2011), although in other regions (e.g. Inner Mongolia) mineral-associated C is the largest C pool and shows a strong relationship with climate (Evans, Burke & Lauenroth 2011). In contrast, but in line with our hypotheses, stable C was strongly influenced by soil pH. Higher stocks in more neutral and alkaline soils may reflect greater microbial processing of SOM in higher pH soils, resulting in greater transfer of C to chemically protected pools (Fornara *et al.* 2011). There was also a relatively small and unexpected effect of CWM leaf N content, which might be explained by N-rich plant material reducing the need for ‘microbial mining’, a process where soil microbes decompose SOM to acquire N (Craine, Morrow & Fierer 2007). Given that CWM leaf N is higher in improved, fertilized grasslands (De Vries *et al.* 2012), it might also reflect higher inorganic N availability in a more statistically parsimonious way than the deleted management term. Management was not retained in any of our models, and this may reflect the very broad categories used, which cover a range of fertilizer and mowing regimes, and grazing intensities. Gathering detailed and accurate data for such factors requires considerable effort, and plant traits that respond to all these factors may act as a good proxy substitute for them.

Models describing the total C stocks and the intermediate fraction were extremely similar, which is unsurprising given that most soil C was in the intermediate fraction. The decline of soil C stocks at intermediate pH is likely caused by the balance of two contrasting processes: reduced decomposer activity and the accumulation of plant inputs in low pH conditions (Pietri & Brookes 2008), and greater transfer of C to the stable C fraction in more neutral and calcareous soils (Fornara *et al.* 2011). The moisture term in the total C model demonstrates that stocks peaked at soil moisture levels typical of waterlogged, or wet grasslands where soil microbial activity is low. The lack of plant trait terms in these models may reflect the fact that most older soil C has either undergone chemical and/or physical transformation into more complex forms, or because current plant community composition does not reflect its origin.

Previous studies have shown that regression models can predict soil C variation using climate and soil texture data at very large scales and within several continents (>100 000 km²) (Burke *et al.* 1989; Paruelo *et al.* 1998; but see Evans, Burke & Lauenroth 2011). Our

findings indicate that a combination of plant trait data and simple climate and soil measures can also help to predict regional- and national-scale soil C stocks (1–100 000 km²) in the surface soil, in a range of C pools of varying stability. It is possible that this approach could greatly improve regional- and national-level predictions of surface soil C stocks compared to current land cover proxy methods (Eigenbrod *et al.* 2010). Climate data are available for many parts of the world, soil pH can be measured quickly and with little equipment, and many countries produce regular national surveys of plant community composition (e.g. Carey *et al.* 2008). Furthermore, large international trait data bases now exist (Kattge *et al.* 2011), and some traits, such as leaf N, may be predictable from remote sensing (Dahlin, Asner & Field 2013). Our approach may also complement current soil C mapping approaches, which use a combination of dynamic models such as CEN-TURY (Parton *et al.* 1993) and RothC (Smith *et al.* 2005), direct measurements (Bellamy *et al.* 2005; Carey *et al.* 2008) and proxy measures (Jones *et al.* 2005; Eigenbrod *et al.* 2010), in two ways. First, it could be used to parameterize the starting conditions for soil C pools in models, and secondly, it could provide more extensive and fine-scale coverage than might be possible from direct measurement, for example for cases in which landowners seek to map soil C.

The large amount of variation captured by the random effects in our models is likely to represent site differences in geology and legacies of landscape history (e.g. land use and glaciation), which may already be captured in coarse-scale soil surveys. The measures here could help refine these coarse maps using local-scale information about soils, climate and plant communities. Similarly, this approach could help refine existing models that use proxy measures with extensive geographic coverage (e.g. land cover and climate) (e.g. Paruelo *et al.* 1998; Jones *et al.* 2005), by improving the characterization of existing relationships and including trait-based vegetation measures that are general, more detailed and mechanistically informative. Such an approach requires extension to a wider range of soil depths, environmental conditions and ecosystem types before it can be widely applied. Nevertheless, our results show that it has great potential, especially given the urgent need for large-scale, cost-effective and accurate soil C stock characterization. Such information is a precursor for the inclusion of soil C into C-trading schemes and improved ecosystem service management.

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Data accessibility

All data used in this article are available in Dryad Digital Repository, doi: 10.5061/dryad.s7867 (Manning *et al.* 2015).

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Supporting Information

Additional Supporting Information may be found in the online version of this article.

Appendix S1. Details of soil C measurement and statistical modelling procedures.

Fig. S1. Demonstration of sampling strategy.

Fig. S2. Comparison of soil carbon stocks predicted by the statistical models and actual stocks.

Table S1. The mean and range of data used in model fitting and validation.

Table S2. Parameter combinations fitted in the statistical modelling procedure.

Table S3–S6. Model statistics for total soil carbon, and the active, intermediate and stable fractions.

Table S7. Model validation comparison of predicted and actual values.

Appendix S2. Additional references for trait data sources and appendix S1.